When is an ecological network complex? Connectance drives degree distribution and emerging network properties

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- 4 Connectance and degree distributions are important components of the structure of eco-
- logical networks. In this contribution, we use a statistical argument and simple network
- generating models to show that properties of the degree distribution are driven by net-
- work connectance. We discuss the consequences of this finding for (1) the generation of
- random networks in null-model analyses, and (2) the interpretation of network structure
- and ecosystem properties in relationship with degree distribution.

10 1 Introduction

Ecologists developed a strong interest for network theory, as it allowed to make sense of some of the 11 complexity of ecological communities. In contrast to early approaches on "community modules" 12 (groups of a few species within a large community, Holt, 1997) a network level approach allows 13 one to account for the whole community scale (J. A. Dunne, 2006), thus integrating all direct and 14 indirect interactions (Berlow et al., 2009). Ecological networks have often been called "complex" 15 (R. J. Williams & Martinez, 2000), on account of the fact that they represent objects (ecological communities) with complex dynamics (i.-e. non-linear, sensitive to indirect interactions). Because networks are multi-faceted objects with a rich range of structure, ecologists have been looking for emerging properties that can be easily measured and analyzed, and that relate to ecological properties and processes. Early in the ecological network literature, connectance, i.e. the proportion of realized ecological 21 interactions among the potential ones (most often the squared species richness), has been recognized 22 as a central network property (May, 1972; Yodzis, 1980; N. D. Martinez, 1992). In part, this success can 23 be attributed to the relationship between connectance and early definitions of network complexity 24 (Pimm, 1982), and to the fact that connectance predicts reasonably well key dynamical properties of 25 ecological networks (J. A. Dunne, Williams, & Martinez, 2002; 2002) including their stability (May, 26 1972). More recently, attention shifted from connectance, a community-averaged property, to the 27 degree distribution, that is the statistical properties of the distribution of number of interactions per species. Variation of degree distribution among networks has often been taken as evidence that assembly or interaction mechanisms differ (Vázquez, 2005; Williams, 2011), and increasingly refined 30 methods to estimate degree distribution have been devised (Williams, 2009). Some authors proposed that degree distribution, rather than connectance, is driving higher level network properties such as nestedness or modularity, which are important drivers of network dynamics (Miguel Angel Fortuna et al., 2010).

However, it is worth asking if we were not too quick in focusing most of our research effort on degree

distribution, in detriment to more fondamental work on connectance and its effects. A network, ecological or otherwise, can be viewed as a physical space that edges (interactions) occupy. The size 37 of this space is limited by the number of nodes. This means that there are physical constraints on the 38 filling of a network, due to the fact that placing the first edge will limit the number of ways to place 39 the remaining edges, and so on. For example, there is only one way to have a fully connected network, and there are a limited number of ways to have a network with the lowest possible connectance. For this reason, and given the rising importance of degree distribution in the literature, it is important that we clearly understand how constrained this distribution actually is in relation to connectance. In this contribution, using an argument from combinatorial statistics and simulations of pseudo-random networks under two different models, we present strong evidence that degree distribution, along with 45 other emerging network properties, are constrained (and can be predicted to a certain extent) by connectance. We discuss the consequences of our results for the comparison of different ecological networks, and for the generation of random networks in null-model analyses. 48

2 Statistical argument

Assuming an ecological network made of n species, and assuming undirected interactions with no self-edges (e.g. no species can interact with itself), there can be at most M = n(n-1)/2 interactions 51 in this network, in which case it is a complete graph (the results presented below hold qualitatively for 52 both directed graphs, and graphs in which self-edges are allowed). We note this maximal number of 53 links M_n . With this information in hand, it is possible to know the total number of possible networks 54 given a number l of interactions. Ecologists are often more familiar with networks being represented 55 as their adjacency matrix, i.e. (with minimal simplifications) a matrix with as many rows and columns 56 as there are species, and a 1 at the intersection of two species that interact. In an undirected network, 57 the existence of an edge between species A and B imply two interactions (i.e. $A \rightarrow B$ and $B \leftarrow A$), and so assuming no self-edges, the total number of ones in the adjacency matrix of a complete, undirected graph, is n(n-1). Throughout this paper, we represent networks as graphs, and not as adjacency matrices. We consider only the situation of unipartite networks, *i.e.* that can be represented by a square matrix. While the shape of the matrix (*i.e.* the ratio of columns numbers over rows number) will have an impact on the results, we conducted preliminary simulations showing that the results hold qualitatively in bipartite networks of varying shapes.

If we term S_n the set of all possible M_n edges in a n-node network, then the number $G_{n,l}$ of possible networks with l links is the number of l-combinations of S_n , i.e. how many possibilities are there to pick l edges among M_n . Formally, this is expressed as $G_{n,l} = C_l^{M_n}$, (where C_x^y is the binomial coefficient, i.e. the number of possible ways to pick x elements among a set of y elements) or

$$G_{n,l} = \frac{M_n!}{l!(M_n - l)!}$$

Note that this number of possible networks include some graphs in which nodes have a degree of o, and that in most ecological studies, such nodes will be discarded. We therefore have to evaluate how 70 many of such networks will be found withing $G_{n,l}$. In addition, in a null-model context (Bascompte et al., 2003; Miguel A. Fortuna & Bascompte, 2006), having unconnected nodes in random replicates will change the richness of the community, thus possibly biasing the value of randomized emerging 73 properties. As some measures of network structure covary with species richness, if one is to generate 74 a randomly expected distribution of the values of these properties, then it is important to hold 75 species richness constant. Finding out the number of networks in which a given node has a degree 76 of o is similar to finding out how many networks exist with l links between the n-1 nodes. If 77 one node is removed from the network, there are C_{n-1}^n possible combinations of nodes (which is 78 (n)!/((n-1)!(n-(n-1))!), which further simplifies to n). For each of these, there are $G_{n-1,l}$ possible 79 networks configurations. Note that these networks will also include situations in which more than 80 one species has a degree of o, so that by recurrence, evaluating $G_{n-2,l}$ and so forth is not necessary 81 (all networks with more than one node of null degree are within the set of the networks with at least 82 one node of null degree). Calling $R_{n,l}$ the number of networks with n nodes and l edges in which all nodes have at least one edge attached, we can write that the number of networks with all nodes

having at least one edge is the total number of networks minus the number of networks having at least one node of null degree (evaluated for each node), or

$$R_{n,l} = G_{n,l} - C_{n-1}^n \times G_{n-1,l}$$

We call the quantities R and G, respectively, the *realized* and *total* network space. They measure how many networks of n nodes and l edges exists, either allowing or preventing the existence of nodes with no interactions. Based on this reasoning, we can make two predictions.

Prediction 1: Because $C_x^y = C_{y-x}^y$, it comes that the total network space is largest when $l = M_n/2$. As in this context the maximal number of edges is M_n , we define effective connectance as $Co = l/M_n$, so $\max(G_{n,l})$ is reached at Co = 1/2. The algebraic expression of the maximum value of $R_{n,l}$ is hard to find, but simulations show that it also occurs around Co = 1/2. In other words, regardless of the number of nodes in a network, the "degrees of freedom" of network structure, as indicated by the size of the realized and total network spaces, is maximized at intermediate connectance.

Prediction 2: $R_{n,l}$ will become asymptotically closer to $G_{n,l}$ when l is close to M_n . In other words, there is only one way to fill a network of n nodes with M_n interactions, and in this situation there is no possibility to have nodes with a degree of n. In the situation in which n0 which n0 with n0 given that n0 which n0 with n0 w

We now illustrate these predictions using networks of 10 nodes, with a number of edges varying from 104 10 to M_{10} (*i.e.* 45 edges). As illustrated in Fig. 1, the size of the network space has a hump-shaped relationship with connectance, and the size of the realized network space becomes closer to the size of the total network space when connectance increases.

In Fig. 2, we show that regardless of the network size, the relative size of the realized network space increases with connectance. The rate at which it occurs increases with network size. However, in all

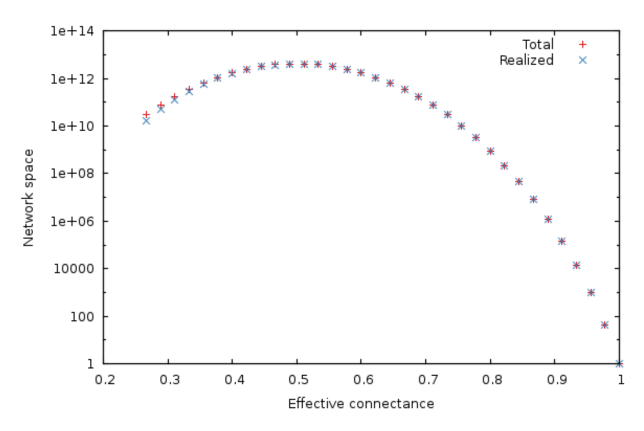


Figure 1: Size of the total and realized network space for n = 10. As predicted in the main text, (1) the size of network spaces peaks at Co = 1/2, and (2) the size of the realized network space becomes asymptotically closer to the size of the total network space when connectance increases.

cases, when connectance is low, there are only a very small proportion of the total network space in which all nodes have at least one edge. This suggest that the structure of extremely sparse networks is also strongly constrained. This is congruent with historical findings by Erdos & Rényi (1959), namely that the probability of each node being connected to the graph largest connected component (*i.e.* any set of vertices of which any two are connected by at least one path) increases with average degree (thus for high connectances, all nodes are likely to be connected to the giant component, hence no node has a degree of o). In the context of ecology, in which most networks have a low connectance, it implies that generating random networks to test null hypotheses can be a computationally intensive task, as the realized network space is (proportionally) small.

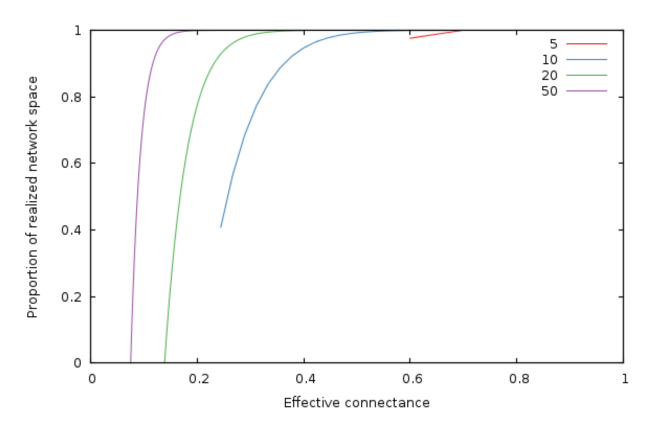


Figure 2: Relative size of the realized network space compared to the total network space when connectance increases, for four different network sizes.

3 Simulations

In the previous part, we show mathematically that connectance (the number of realized *vs.* possible interactions), relative to the network size, determines the size of the *network space*, *i.e.* how many possible network combinations exist. Based on this, we can therefore expect that the degree distribution will be contingent upon network connectance. Specifically, we expect that the variance of the degree distribution, which is often related to ecosystem properties and other network structures (Miguel Angel Fortuna et al., 2010), will display a hump-shaped relationship with connectance. The mean, kurtosis, and skewness of the degree distribution should all vary in a monotonous way with connectance.

In the simulations below, we use networks of 30 nodes, filled with 35 to M_{30} interactions, by steps of 127 10. We use two different routines to generate random networks, that are contrasted in the way they 128 distribute edges among nodes. First, we generate Erdős-Rényi (ER, undirected) graphs, meaning 129 that every potential interaction has the same probability of being realized (Erdos & Rényi, 1959). We 130 use an algorithm inspired by Knuth (1997), allowing to fix the number of edges in the graph rather 131 than the probability of an edge occurring, although the generated graphs have the same properties as 132 the original ER model. A total of 19000 networks are generated this way. Second, we use the niche 133 model of food webs (R. J. Williams & Martinez, 2000), which generates (directed) networks under 134 rules representing hypothesized mechanisms of prey-selection in empirical ecosystems (Gravel et al., 135 2013). This particular model assumes that the existence of interactions is constrained by the position 136 of species along a "niche" axis, for example body size. Other randomization methods for food webs 137 exists, but given that Stouffer et al. (2005) showed that they yield similar degree distributions to the 138 niche model, we will not use them here. A total of 500 replicates for each value of l are generated. All 139 networks generated with the two models are free of self-edges and nodes with a null degree.

For each replicate, we measure the degree distribution and report its variance, coefficient of variation, kurtosis, and skewness. In addition, for each network, we fit a power-law distribution on the sorted degree distribution using the least-squares method; we report the power-law exponent.

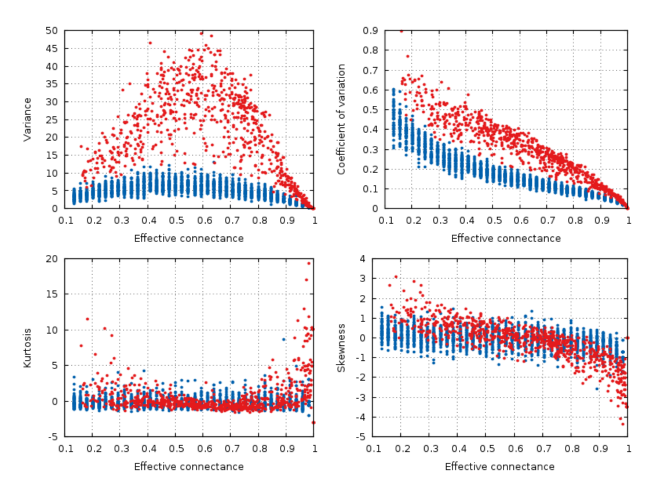


Figure 3: Statistical descriptors of the degree distribution of randomized networks, n = 30, increasing connectance. These results show that central properties of the degree distribution are contingent upon connectance, at a given network size, and under a given network generation model. ER networks are in blue, niche-model networks are in red. Each point represent a single generated network.

Qualitatively, the random graphs and the niche networks behave exactly the same. With the exception 144 of the kurtosis, all statistical descriptors of the degree distribution were influenced by the effective 145 connectance (Fig. 3). As predicted in the previous part, variance of the degree distribution is hump-146 shaped with regard to connectance, which implies that as average degree increases with connectance, 147 the coefficient of variation of the degree distribution decreases at high connectances. Note also that 148 the range of variances in the degree distribution is higher at intermediate connectances, but lower at the extreme. Due to the fact that the Erdős-Rényi graphs we simulate are essentially Poisson random 150 graphs, it is expected that the variance of their degree distribution would be lower than for the niche 151 model, which in contrast forces strong difference in the degree of species according to their niche 152 position. 153

To quantify the impact of connectance on the different network properties, we measured the pro-154 portion of variance explained by the linear regression of a given property against connectance (in 155 such cases as had a linear relationship between the two, i.e. all measures but variance). Kurtosis is 156 independent of connectance ($R_{\text{niche}}^2 = 0.04$, $R_{\text{ER}}^2 = 0.06$), while skewness decreases with connectance, 157 although more markedly so in the niche model ($R_{\text{niche}}^2 = 0.66$, $R_{\text{ER}}^2 = 0.26$). This result is expected. 158 Positively skewed distribution have longer or fatter right tails, indicating mostly low values (low 159 degree): unconnected networks are made mostly of species with a weak generality (Schoener, 1989). 160 On the other hand, negative skewness indicate that most of the values in the distribution are high. 161 Ecologically, it means that most species are wide-range generalists, which happens in densely con-162 nected networks. This bears important ecological consequences, as it indicates that due to physical 163 constraints acting on the filling of interactions within the graphs, networks with intermediate con-164 nectances are expected to have species with both low and high generality (Schoener, 1989). The 165 coefficientof variation of the degree distribution is extremely well predicted by connectance alone 166 $(R_{\text{niche}}^2 = 0.91, R_{\text{ER}}^2 = 0.87).$ 167

The estimate of the power-law exponent increases when connectance increases (Fig. 4, $R_{\text{niche}}^2 = 0.91$, $R_{\text{ER}}^2 = 0.70$). This indicates that the degree distribution flattens when connectance increases. Taken with the elements presented above, we show that all of the estimators of the degree distribution vary

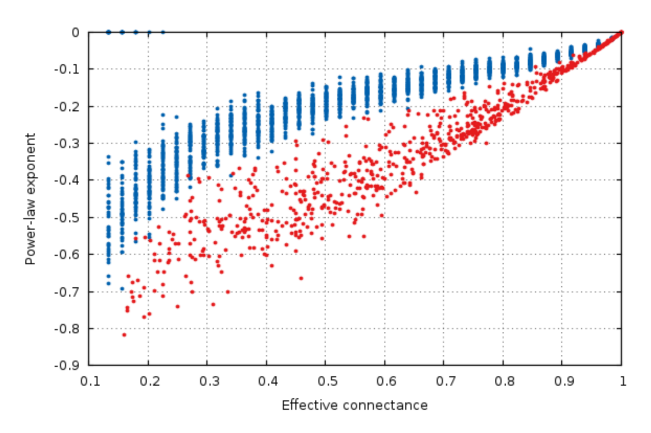


Figure 4: The estimate of the power-law exponent increases with connectance, reaching a flat distribution for complete graphs.

strongly with connectance of the network. Although power-laws should be truncated as soon as the probability of a species having a degree of $2 \times n$ (or n in undirected networks) is not negligible, and as such the fitting of power-laws should not be done on highly connected networks for practical purposes, this result emphasizes the key role of connectance in driving central network structure properties.

176 4 Practical consequences

Randomized null models are often used to estimate how much a given network property deviates from 177 its random expectation (Flores et al., 2011). Our results show two things. First, except for extremely 178 high or low connectance, the proportion of the network space that will be explored using 103 or 104 179 replicates (typical values in null models analyses) is orders of magnitude smaller than the realized 180 network space. Although this is somewhat compensated by the fact that a part of these networks are 181 isomorphic, the risk of inferring deviation from the random expectation based on a drastically small sampling of the network space is real, and un-addressed; at the very least, it seems that that intensity of the replication should be dependent upon the connectance of the network one tries to replicate. On 184 the other hand, when connectance is high, the number of unique network combinations decreases, and there is a risk to generate a number of replicates that is larger than the realized network space, 186 thus decreasing the information content of the randomizations. To the best of our knowledge, these 187 issues have seldom be addressed in the literature on ecological network randomization. 188

Second, generating null models with a low connectance is a computationally intensive task. When connectance decreases, the *realized* network space decreases faster than the *total* network space, meaning that the probability of picking a network with no o degree nodes (which is simply $R_{n,l}/G_{n,l}$) goes toward zero. For this reason, classical rejection sampling (accept the random network if no nodes have no edges, reject it if not) is bound to take an unreasonable amount of time in networks with low connectance. In addition, there is a risk of selecting some particular types of networks. It makes intuitive sense that networks with extremely skewed degree distributions have less chance of

being generated this way, as when a few nodes collect most of the edges, the probability than the remaining nodes each have at least one edge decreases. To the best of our knowledge, this source of bias has not received important attention in the literature. For this reason, using a purely random matrix shuffling as a starting point, then swapping interactions until no free nodes remain, seems to be a promising way to address this problem. Given the important of null-model approaches in network analysis, the generation of efficient and unbiased algorithms is a fruitful research avenue.

5 Conclusions

Connectance is an extremely intuitive property of network, expressing how much of the potential 203 interactions are realized. Through statistical reasoning and simple simulations using models of 204 random networks, we show that for a given number of species, connectance drives (i) how many 205 different networks exist, and (ii) some key elements of the degree distribution. We observed both 206 among and between model quantitative changes in degree distribution along a connectance gradient. 207 The niche model is a particularly striking example of this, with the variance in the degree distribution increasing 50-fold when connectance moves from 0.1 to 0.5. This result has practical implications 209 for network comparisons. As descriptors of degree distribution vary with connectance, connectance 210 should be factored out from all analyses. So as to avoid colinearity issues, this can be done by either 211 working on the residuals of the degree distributions' property of interest. To some extent, the impact 212 of connectance is lesser in the 0.05-0.3 range where most empirical food webs lies (although bipartite 213 networks can have much higher connectances), but the effect is high enough that it should not be 214 ignored: at equal number of species, networks with different connectances are expected to have 215 different degree distributions. 216 Finally, this analysis raises interesting ecological questions. Early analyses focusing on degree distribu-217 tion argued that ecological mechanisms were responsible for the shape of the distribution (Vázquez, 218 2005; Miguel Angel Fortuna et al., 2010; Williams, 2011). In this contribution, we show that con-219 nectance will impose a lower and higher limit for the shape of the degree distribution. Given this 220

information, it is time to bring the debate full-circle: is connectance the cause of observed network properties, or an emergent property of pairwise species interactions? As the later seems far more 222 likely, it now makes sense to focus on why some networks deviate, or not, from the expected degree 223 distribution knowing their connectance. As the density of interaction plays such a central role in 224 May's criteria for stability (May, 1972), clarifying how connectance is shaped by mechanisms regulat-225 ing pairwise species interactions offers the opportunity of integrating the effects of these mechanisms up to their impact on emergent, community-wide properties. Okuyama & Holland (2008) showed 227 that in mutualistic systems, resilience (to perturbation) is affected both by degree network size and interaction strength, but also by the degree distribution and connectance of the network; we show 229 here that degree distribution and connectance are tightly linked, and alternative approaches to the 230 question of resilience can focus on the deviation of degree distribution knowing the connectance, 231 rather than the "raw" degree distribution. In keeping with the results we present in the first part of 232 the paper, Rozdilsky & Stone (2001) report that there is a inverted hump-shaped relationship between 233 connectance, and the proportion of "feasible and stable" systems, with the lowest proportion of such 234 systems being found at $Co \approx 0.5$. This strongly suggests that the same mechanisms that limit the 235 realized network space may affect ecological properties, thus emphasizing the need not to discard 236 connectance in profit of more emerging properties in future ecological network analyses. 237

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240 6 References

Bascompte, J., Jordano, P., Melián, C. J., & Olesen, J. M. 2003. *The nested assembly of plant-animal mutualistic networks*. Proceedings of the National Academy of Sciences of the United States of America 100(16):9383–9387.

Berlow, E. L., Dunne, J. a, Martinez, N. D., Stark, P. B., Williams, R. J., & Brose, U. 2009. *Simple prediction of interaction strengths in complex food webs.* Proceedings of the National Academy of

- Sciences of the United States of America 106(1):187–91.
- Dunne, J. A. 2006. The Network Structure of Food Webs. In J. A. Dunne & M. Pascual (Eds.), Ecological
- networks: Linking structure and dynamics (pp. 27–86). Oxford University Press.
- Dunne, J. A., Williams, R. J., & Martinez, N. D. 2002. Food-web structure and network theory: the
- 250 role of connectance and size. Proceedings of the National Academy of Sciences of the United States of
- 251 America 99(20):12917.
- Dunne, J. A., Williams, R. J., & Martinez, N. D. 2002. Network structure and biodiversity loss in food
- webs: robustness increases with connectance. Ecology Letters 5(4):558-567.
- Erdos, P., & Rényi, A. 1959. On random graphs. Publicationes Mathematicae Debrecen 6:290-297.
- ²⁵⁵ Flores, C. O., Meyer, J. R., Valverde, S., Farr, L., & Weitz, J. S. 2011. Statistical structure of host-phage
- 256 interactions. Proceedings of the National Academy of Sciences of the United States of America
- 257 108(28):E288-297.
- Fortuna, M. A., & Bascompte, J. 2006. Habitat loss and the structure of plant-animal mutualistic
- 259 *networks*. Ecology Letters 9(3):281–286.
- Fortuna, M. A., Stouffer, D. B., Olesen, J. M., Jordano, P., Mouillot, D., Krasnov, B. R., Poulin, R., &
- Bascompte, J. 2010. Nestedness versus modularity in ecological networks: two sides of the same coin?.
- Journal of Animal Ecology 78:811–817.
- ²⁶³ Gravel, D., Poisot, T., Albouy, C., Velez, L., & Mouillot, D. 2013. *Inferring food web structure from*
- 264 predator-prey body size relationships. Methods in Ecology and Evolution.
- Holt, R. D. 1997. Community modules. In A. C. Gange & V. K. Brown (Eds.), Multitrophic Interactions
- in Terrestrial Ecosystems (pp. 333–349). Blackwell Science.
- Knuth, D. E. 1997. *Volume 2: Seminumerical Algorithms*. The Art of Computer Programming 192.
- ²⁶⁸ Martinez, N. D. 1992. Constant connectance in community food webs. The American Naturalist
- 269 139(6):1208-1218.
- 270 May, R. M. 1972. Will a large complex system be stable?. Nature 238:413–414.

- Okuyama, T., & Holland, J. N. 2008. Network structural properties mediate the stability of mutualistic
- 272 communities. Ecology Letters 11(3):208–216.
- Pimm, S. L. 1982. Food Webs. London: Chapman & Hall.
- 274 Rozdilsky, I. D., & Stone, L. 2001. Complexity can enhance stability in competitive systems. Ecology
- 275 Letters 4(5):397-400.
- ²⁷⁶ Schoener, T. W. 1989. Food webs from the small to the large. Ecology 70(6):1559–1589.
- Stouffer, D. B., Camacho, J., Guimera, R., Ng, C. A., & Amaral, L. A. N. 2005. Quantitative patterns in
- 278 the structure of model and empirical food webs. Ecology 86(5):1301–1311.
- ²⁷⁹ Vázquez, D. P. 2005. Degree distribution in plant-animal mutualistic networks: forbidden links or
- 280 random interactions?. Oikos 108(2):421-426.
- ²⁸¹ Williams, R. J. 2009. Simple MaxEnt models explain food web degree distributions. Theoretical Ecology
- 282 3(1):45-52.
- ²⁸³ Williams, R. J. 2011. Biology, Methodology or Chance? The Degree Distributions of Bipartite Ecological
- Networks. (J. Langowski, Ed.)PLoS One 6(3):e17645.
- ²⁸⁵ Williams, R. J., & Martinez, N. D. 2000. Simple rules yield complex food webs. Nature 404(March):180-
- 286 183.
- Yodzis, P. 1980. The connectance of real ecosystems. Nature 284:544-545.