Keystoneness, centrality, and the structural controllability of ecological networks

- ¹ E. Fernando Cagua¹, Kate L. Wootton^{1,2}, Daniel B. Stouffer¹
- ² Centre for Integrative Ecology, School of Biological Sciences, University of Canterbury, Private Bag
- ³ 4800, Christchurch 8041, New Zealand
- ⁴ Current address: Department of Ecology, Swedish University of Agricultural Sciences, Box 7044, SE-750
- 5 07 Uppsala, Sweden
- 6 Author for correspondence: Daniel B. Stouffer (daniel.stouffer@canterbury.ac.nz) +64 3 364 2729 -
- ⁷ Centre for Integrative Ecology, School of Biological Sciences, University of Canterbury, Private Bag 4800,
- 8 Christchurch 8140, New Zealand

Abstract

16

17

18

19

20

21

23

25

26

27

28

- 1. An important dimension of a species' role is its ability to alter the state and maintain the diversity of its community. Centrality metrics have often been used to identify these species, which are sometimes 10 referred as "keystone" species. However, the relationship between centrality and keystoneness is 11 largely phenomenological and based mostly on our intuition regarding what constitutes an important 12 species. While centrality is useful when predicting which species' extinctions could cause the largest 13 change in a community, it says little about how these species could be used to attain or preserve a 14 particular community state. 15
 - 2. Here we introduce structural controllability, an approach that allows us to quantify the extent to which network topology can be harnessed to achieve a desired state. It also allows us to quantify a species' control capacity—its relative importance—and identify the set of species that , collectively, are critical in this context because they have the largest possible control capacity. We illustrate the application of structural controllability with ten pairs of uninvaded and invaded plant-pollinator communities.
- 3. We found that the controllability of a community is not dependent on its invasion status, but on 22 the asymmetric nature of its mutual dependences. While central species were also likely to have a large control capacity, centrality fails to identify species that, despite being less connected, were critical in their communities. Interestingly, this set of critical species was mostly composed of plants and included every invasive species in our dataset was part of it. We also found that species with high control capacity, and in particular critical species, contribute the most to the stable coexistence of their community. This result was true, even when controlling for its the species' degree, abundance/interaction strength, and the relative dependence of their partners. 29
- 4. Synthesis: Structural controllability is strongly related to the stability of a network and measures 30 the difficulty of managing an ecological community. It also identifies species that are critical to 31 sustain biodiversity and to change or maintain the state of their community and are therefore likely 32 to be very relevant for management and conservation. 33
- Keywords: Keystone species, Invasive species, keystone species, management interventions, mutualism, 34
- network control theory, plant population and community dynamics, species' importance, control capacity,
- structural stability , controllability

Introduction

A major goal in ecology is to understand the roles played by different species in the biotic environment. Within community ecology, a complex-systems approach has led to the development of a variety of analytical and simulation tools with which to compare and contrast the roles of species embedded in a network of interactions (J.-Bascompte & Stouffer, 2009; Coux, Rader, Bartomeus, & Tylianakis, 2016; Guimerà & Amaral, 2005; Stouffer, Sales-Pardo, Sirer, & Bascompte, 2012). A particularly relevant dimension of any species' role is its ability to alter the abundance of other species and the state of the 42 community—since changes of this nature can have knock-on effects on ecosystem function, diversity, processes, and services (Thompson et al., 2012; Tylianakis, Didham, Bascompte, & Wardle, 2008; Tylianakis, Laliberté, Nielsen, & Bascompte, 2010). This ability is sometimes referred to as a species' "keystoneness" (Mills & Doak, 1993). A significant proportion of the network tools used to estimate species' roles in this context rely on the calculation of a species' centrality—a relative ranking of its positional importance that originally stems from social-network research (Friedkin, 1991; Martín González, Dalsgaard, & Olesen, 2010). Generally speaking, central species tend to be better connected and consequently are more likely to participate in the network's "food chains". Because species that participate in more chains are more likely to affect the abundances of other species, centrality metrics have often been used to identify keystone species in the community (Jordán, Benedek, & Podani, 2007). Centrality metrics have been shown to be useful tools to rank species in regard to their potential to alter the abundances of other species, in particular 54 when estimating the probability of secondary extinctions that may follow the loss of a species (Dunne, 55 Williams, & Martinez, 2002; Kaiser-Bunbury, Muff, Memmott, Müller, & Caflisch, 2010). Despite being conceptually intuitive, the relationship between centrality and a species' presumed impact 57 on the state of the community is largely phenomenological. On the one hand, substantive changes in ecosystem functioning can also occur without complete removal of a species (Mouillot, Graham, Villéger, Mason, & Bellwood, 2013). On the other, we are often interested in a specific state of the community that might be desirable to attain (or preserve) because of its biodiversity, resilience, functioning, or the ecosystem services it provides. In these cases, it might be less useful to understand which species may 62 cause any change in the community. Instead, we are better served by understanding how the structure 63 of the network can be harnessed to achieve the desired state and which species may play the largest role in this targeted process. When the state of a community is underpinned by more than a single 65 species (often the case in real communities) and we move beyond single-species removals, we might expect the accuracy of centrality to diminish. As a result, community ecology could arguably benefit from an alternative, perhaps more mechanistically-grounded, approach to understand how species affect each other's abundance.

Species' abundances—and consequently the state of the community as a whole—are influenced both by the structure of their interactions and the dynamics of these interactions, including the mechanisms of self-regulation (Lever, van Nes, Scheffer, & Bascompte, 2014). However, community and population 72 dynamics can be modelled in innumerable ways, and empirical support for one versus another is often 73 still ambiguous (Holland, DeAngelis, & Bronstein, 2002). The alternative approach should, therefore, ideally acknowledge ecosystem dynamics, but without being overly dependent on the particular choices of 75 how they are characterised. Among the various possibilities structural controllability, a branch of control theory, appears to be a strong candidate (Isbell & Loreau, 2013). Control theory is a widely-studied branch 77 of engineering used to determine and supervise the behaviour of dynamical systems (A. E. Motter, 2015). It is inherently designed to deal with system feedbacks and its application has recently been expanded to 79 complex networks (Lin, 1974; Liu & Barabási, 2016). Consistent with long-standing ecological questions, 80 advances in structural controllability have established a clear link between the structure of the network 81 and the way nodes affect each other. Unlike centrality indices, however, this link is not based on a priori assumptions between network metrics and keystoneness but is instead based on well-established advances in both dynamical and complex-systems theory (A. E. Motter, 2015). At its fundamental level, structural controllability first determines whether a system is controllable or not; that is, it asks if a system could ever be driven to a desired state within a finite amount of time. Although the controllability of a network is a whole-system property, it has recently been shown that asking for the 87 controllability of a complex-system is equivalent to finding a particular set of relevant nodes: the set with which is possible to control the state of the whole network (Liu & Barabási, 2016). Importantly, this set of nodes is not always unique for a given network. This implies that an examination of the distinct sets provides a means to connect nodes with their general ability to modify the system to which they belong. Here, we apply methods from structural controllability to a particular ecological problem and show how it can be used to generate insight into the role of species in an ecological network. Specifically, we outline 93 the approach using a set of ten pairs of uninvaded and invaded plant-pollinator communities. We use 94 invaded communities because there is strong empirical evidence showing that invasive species play an 95 important role shaping the abundances of other species, something which is particularly true in these ten networks (Bartomeus, Vilà, & Santamaría, 2008; Lopezaraiza-Mikel, Hayes, Whalley, & Memmott, 2007). This choice thus offers us an opportunity to explicitly contrast our theoretical observations with empirical evidence. Moreover, empirical observations indicate that steering the state of some communities—for example during ecosystem restoration or invasive species removal—can be a very difficult task (Woodford et al., 2016). Therefore, we first ask whether there are differences between the controllability of invaded and uninvaded networks. We then expand existing methods from control theory to effectively link the 102 controllability (Table 1) of a network with the role of its constituent species. We ask—from a control-103

Table 1: Glossary

network control

A network is said to be controllable if it is possible to steer it from an initial to an arbitrary final state within finite time.

controllability

The intrinsic difficulty of controlling an ecological community. It is measured by the relative size of the minimum driver-node set, n_D . It also indicates the extent to which network structure can be harnessed for network control.

minimum driver-node set

One of the sets of species whose abundances need to be directly managed in order to achieve full control of the community. The minimum driver-node sets can be obtained by finding all maximum matchings in a network.

maximum matching

A matching is a set of links that do not share any common start or end nodes; the largest possible matching is called a maximum matching.

control configuration

One of the species combinations with which is possible to achieve network control. Optimal control configurations are given by the minimum driver-node sets.

control capacity

The relative frequency ϕ which with a species is part of the optimal control configurations of a network.

critical species

A species with a maximal control capacity $\phi = 1$.

superior node

A species is a superior node if it can internally affect the abundance of other species in the network. Superior nodes make up the chains that propagate the control signals through the network.

- theoretic perspective—whether there are key differences between species in the role they play at driving
- the state of the community and explore the ecological factors related to these differences. This allows
- us to identify species that might be critical for network control and show that they have a larger than
- expected impact on the stable coexistence of the community. Finally, we compare the proposed approach
- to current methods based on species centrality and show how these methods are indeed valuable but
- ultimately paint a limited picture in regard to the "keystoneness" of a species.

Materials and methods

- 110 We used ten paired pollination communities to apply the control-theoretic approach. Each community
- pair was composed of a community invaded by a plant and a community free of the invasive species.

Four pairs correspond to natural or semi-natural vegetation communities in the city of Bristol, UK 112 (Lopezaraiza–Mikel et al., 2007). These communities are comprised of comprised 19–87 species (mean 55), 113 and non-invaded plots were obtained by experimentally removing all the flowers of the invasive species 114 Impatients grandulifera. The other six pairs were obtained from lower diversity Mediterranean shrublands in Cap de Creus National Park, Spain (Bartomeus et al., 2008). These communities are comprised of comprised 30-57 species (mean 38); in contrast to the above, uninvaded communities were obtained from 117 plots that had not yet been colonised by either of the invasive species Carpobrotus affine acinaciformis or 118 Opuntia stricta. The structure of all these communities was defined by the pollinator visitation frequency, 119 which has been shown to be an appropriate surrogate for interspecific effects in pollination networks (J. 120 Bascompte, Jordano, & Olesen, 2006; Diego P. Vázquez, Morris, & Jordano, 2005). Full details about the 121 empirical networks can be found in the Supporting Information Section S1. 122 The first step in applying methods of control theory is to construct a directed network that is able to 123 provide an indication of the extent to which species affect each other's abundance. In some ecological 124 networks, establishing the directionality can be relatively straightforward, for example when links represent biomass transfer or energy flow (Isbell & Loreau, 2013). In pollination networks, however, this directionality 126 is less obvious as both species can, in theory, benefit from the interaction. We overcome that obstacle by 127 noting that the extent to which species i affects species j relative to the extent to which j affects i can be 128 summarised by their interaction asymmetry (J. Bascompte et al., 2006). This asymmetry is given by

$$a(i,j) = a(j,i) = \frac{|d_{ij} - d_{ji}|}{max(d_{ij}, d_{ji})},$$

129

where the dependence of plant i on pollinator j, d_{ij} , is the proportion of the visits from pollinator j 130 compared to all pollinator visits to plant i. Previous research has shown that mutualistic interactions are 131 often highly asymmetric in natural communities; in other words, if a plant species is largely dependent 132 on a pollinator species, that pollinator tends to depend rather weakly on the plant (and vice versa). 133 We therefore create a directed link from species i to species j when $d_{ij} - d_{ji} \ge 0$ to establish the most likely direction of control between a species pair (Figure 1a). Sometimes Sometimes (2.4% of the observed interactions in our datasets) there is no observed asymmetry between species pairs $(d_{ij} = d_{ji})$, and we cannot infer a dominant direction of control. When this occurs, we deem both species to be equally likely 137 to affect each other and leave a reciprocal interaction between them (a link from i to j and another from j138 to i). By basing the direction of the links on the asymmetry of their dependence, we are able to generate 139 a network that is consistent with the dynamics of the community while satisfying the requirements of 140 structural controllability. This allows us to calculate the controllability of the networks and investigate 141 whether there are differences between invaded and uninvaded communities. 142

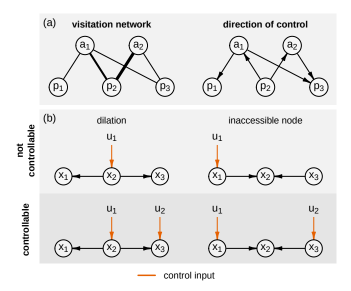


Figure 1: The direction of control and controllability conditions. (a) To establish the direction of control, we start with a weighted visitation network (on the left). In this network, the width of the links corresponds to the frequency of visitation between animals a_i and plants p_i , with wider links indicating more visits. Plant p_1 is visited exclusively by a_1 but p_1 represents only a small fraction of the floral resources exploited by a_1 . Therefore, the population of p_1 is more likely to be affected by a_1 than vice versa. We represent this with a directed link from a_1 to p_1 in the control network (on the right). The direction of control between all other species pairs can be similarly determined by inspecting the difference between their relative dependences. (b) Once we have established the directions of control, we can determine whether the network is controllable or not. Any system defined by a directed network (with state nodes x_i ; species' populations in an ecological context) and external control inputs (nodes u_i , orange links) is structurally controllable if it satisfies two conditions: it has no dilations (expansions in the network) and no inaccessible nodes. The system on the top left is not controllable because there is a dilation since node x_2 is being used to control two nodes simultaneously; in other words, there are fewer superiors (x_2) than subordinates $(x_1 \text{ and } x_3)$. The network on the top right is not controllable because node x_3 is inaccessible for the only input node u_1 in the system. Both systems can be made controllable by adding an extra input node (u_2 in both bottom networks).

Controllability

A system is said to be controllable if it is possible to steer it from an initial to an arbitrary final state within finite time (Kalman, 1963). A simple version of such a system can be described by $\frac{dx}{dt} = Ax + Bu(t)$, 144 where the change of its state over time $\left(\frac{dx}{dt}\right)$ depends on its current state x (for example the species) 145 abundances), an external time-varying input u(t) (the control signal), and two matrices A and B, which encode information about the network structure and how species respond to external inputs, respectively. In classic control theory, determining whether this system is controllable can be achieved by checking that 148 its controllability matrix $R = \begin{bmatrix} B & AB & A^2B & \dots & A^{n-1}B \end{bmatrix}$ has full rank. In complex systems, however, 149 employing this rank condition, or numerical approximations of it is infeasible because it is hard to fully 150 parameterise A and B (either because the weight of the links changes over time or because they are 151 difficult to measure). Here, we use an approach based on the structural controllability theorem (Lin, 152 1974), which assumes that we are confident about which elements of A and B have either non-zero or 153 zero values (there is an interaction or not), but that we are less sure about the precise magnitude of the 154 non-zero values. Using this structural approach, we can find out the controllability of a system for every 155 non-zero realisation of the parameters. We are often able to estimate A in ecological networks, as this matrix represents the interactions between 157 species. Part of the control problem thus resides in estimating a supportable estimation of B, which 158 represents the links between external inputs and species. Naively, any ecological community (and any 159 system for that matter) could be controlled if we control the state of every species independently, but 160 such an approach is typically impractical. Here, we are interested in finding a minimum driver-node 161 set (effectively finding B) with which to make the system controllable. The brute-force search for this 162 minimum driver-node set is computationally prohibitive for most networks as it involves the evaluation of 163 2^N different controllability matrices where N is the number of species in the community. We therefore instead employ a recently-developed approach that shows that the control problem of finding the minimum 165

Maximum matching is a widely studied topic in graph theory and is commonly used in multiple applications, ranging from dating apps and wireless communications to organ transplant allocation and peer-to-peer file sharing. A matching in an unweighted directed graph is defined as a set of links that do not share common start or end nodes; the largest possible matching is called a maximum matching. For example, in a network composed of jobs and job applicants, a matching is any pairing between applicants and positions that satisfies one basic constraint: an applicant can be assigned to at most one position and vice versa. Consequently, a maximum matching is an optimal pairing, one that maximises the number of applicants with jobs and the number of positions filled. Admittedly, the link between matchings and

driver-node set can be mapped into a graph-theoretic problem: maximum matching (Liu & Barabási,

2016; Liu, Slotine, & Barabási, 2011).

167

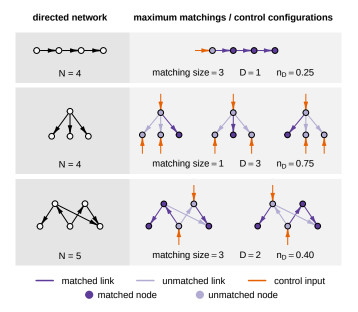


Figure 2: Maximum matchings and control configurations. In directed networks, a maximum matching is the largest possible set of links that do not share start or end nodes (dark purple). Maximum matchings are not necessarily unique; instead, each of them is related to a possible minimum driver-node set in the network (the nodes to which an external control input, in orange, should be applied in order to ensure controllability). The size of the minimum driver-node set D corresponds exactly to the number of unmatched nodes (the number of nodes in the network N minus the matching size). To account for network size, we use the size of the minimum driver-node set relative to the total number of nodes $n_D = D/N$ as a measure of the extent to which the network structure can be harnessed to control the system.

176 structural controllability may appear far from straightforward.

This link becomes apparent after examining the graphical interpretation of structural controllability: from 177 a topological perspective, a network is structurally controllable if there are no inaccessible nodes—that is, 178 nodes without incoming links—or dilations—expansions of the network (Figure 1b; Supporting Information 179 Section S2). The key is to note that these two fundamental conditions of structural controllability imply 180 that there is a one-to-one relationship between superior and subordinate nodes just like the one-to-one relationship between jobs and applicants (Figure 1b, bottom left). We thus use the maximum-matching 182 algorithm to find an optimal pairing of superior (those that can control another node) and subordinate 183 nodes (those that can be controlled by another node) in a manner consistent with the controllability 184 conditions (Supporting Information Section S3.1). Given the result, we can further decompose the 185 matching into a set of paths that reveal how a control signal can flow across the links in a network to 186 reach every node within it. As recently shown (Liu et al., 2011), the minimum driver-node set—those to 187 which an external control input should be applied to make the system controllable—corresponds exactly 188 to the unmatched nodes in the network (Figure 2).

Differences between invaded and uninvaded networks

Our first ecological Our first objective is to investigate whether the controllability of a community is

associated with invasion status or not. Finding out exactly how difficult it is to control a network depends 191 strongly on the particularities of the desired control trajectory (i.e. the path to the desired final state) 192 as well as the dynamical relationship between nodes. However, we are interested in understanding the controllability of a network in a more general sense, such that it can be applied even when the precise control scenario is known only incompletely. To this end, we chose an indicator that follows directly from our approach: the size of the minimum driver-node set. This simple metric provides a general indication 196 of how difficult controlling a network might be, as systems that require a large number of external inputs 197 to be fully controlled are intuitively more difficult or costly to manage. For instance, achieving full control 198 in a "network" in which species do not interact at all is relatively more difficult as we would require an 199 intervention for every single species. Conversely, the structure of a linear trophic chain can be harnessed 200 to achieve full control using just one intervention targeted at the top species; a suitable control signal 201 could then cascade through the trophic levels and reach other species in the community. Specifically, 202 drawing from the structural-controllability literature, we use the size of the minimum driver-node set relative to the total number of species $n_D = \frac{D}{N}$ as a measure of the controllability of a network—the extent to which the network structure can be harnessed to control the community. The lower n_D the more 205 controllable the community. In an ecological context, external inputs can be thought of as management interventions that modify the abundance of a particular species. 207

After finding the minimum driver-node set in each of our networks, we wanted to test whether invasion 208 status or other predictors are correlated to controllability. We do this using a set of generalised linear 209 models with binomial error structure Gaussian errors and a logit link function. The response variable was 210 the relative size of the minimum driver-node set n_D of the twenty empirical networks (ten invaded and 211 ten uninvaded), and we included invasion status as a predictor. As predictors, we also include the network connectance, the network nestedness (NODF), the number of species (since one might naively expect to see a negative relationship between richness and controllability; Menge, 1995), the network asymmetry (an indication of the balance between plant and pollinator diversity), and the interaction strength asymmetry 215 (the asymmetry on the dependences between trophic levels; N. Blüthgen, Menzel, Hovestadt, Fiala, & 216 Blüthgen, 2007). We compared models using the Akaike Information Criterion for small sample sizes 217 (AICc). 218

In addition, we also explored whether real networks differ in their architecture from random ones in a concerted way that could impact these results. Specifically, we used two null models each with 99 randomisations per network. In the first, we followed $\frac{\text{Diego P}}{\text{P}}$ Vázquez et al. (2007) and maintained the connectance of the network but randomised the visits across species such that the relative probabilities of interactions were maintained. We then re-estimated the direction of control and the corresponding size of the minimum driver-node set, n_D . For the second null model, we used the empirical directed network described above and randomly shuffled the direction of control between a species pair prior to re-estimating the size of the minimum driver-node set.

Species' roles

Our second objective is related to how species differ in their ability to drive the population dynamics of
the community. We in turn examine whether these differences are also reflected in the role species play at
supporting the stable coexistence of other species in the community. Ecologically, these differences are
relevant because resources and data are limited, and therefore full control is infeasible. While calculating
the size of the minimum drive-node set can measure the controllability of an ecological community, it
does not provide information about the roles that particular species play.

To answer this question, we harness the fact there may be multiple maximum matchings for a given network, and each of these maximum matchings indicates a unique combination of species with which it is possible to control the network. Moreover, some species belong to these combinations more often than do others. We call this property a species, "control capacity," ϕ . The higher a species, control capacity, the greater the likelihood that they it would need to be directly managed to change (or maintain) the ecological state of their community. Therefore, a species, control capacity provides an estimation of their its relative importance at driving the state of the community (Jia & Barabási, 2013).

To calculate a species' control capacity ϕ , we must first enumerate all possible maximum matchings (Supporting Information Section S3.2). Unfortunately, enumerating all maximum matchings is extremely 241 expensive from a computational perspective—a network with a couple dozen species has several hundred million unique maximum matchings. To solve this problem, we employ a recently-developed algorithm that reveals the control correlations between the nodes in the graph while requiring considerably less computational resources (Zhang, Lv, & Pu, 2016). Using this algorithm, we are able to identify species that are possible control inputs—those that belong to the minimum driver-node set in at least one of the 246 possible control configurations. Here, we extend this algorithm such that it is possible to calculate a highly 247 accurate approximation of the control capacity ϕ of every species in the network (Supporting Information 248 Section S3.3). In the networks that contained reciprocal links (because there was no asymmetry in the dependences of a species pair), we averaged a species' control capacity ϕ across every possible 250 "non-reciprocal" version of the network (Supporting Information Section S3.4). 251

We then examined how species-level properties were related to control capacity using a set of candidate generalised linear models with binomial error structure. These models included five predictor variables that mirror the network-level predictors. First, the species' contribution to nestedness, which has been proposed as a key feature that promotes stability and robustness in mutualistic networks (S.-Saavedra,

Stouffer, Uzzi, & Bascompte, 2011). Second, the species the species' strength (the sum of a species' 256 visits), which quantifies the strength of a species' associations and is indirectly related to its abundance 257 (Poisot, Canard, Mouquet, & Hochberg, 2012). Third, the direction of asymmetry which quantifies the 258 net balance in dependencies; that is, it indicates if a species affects other species more than what they 259 affect it or not (Diego P-Vázquez et al., 2007). Fourth, the species' degree in order to account for the intrinsic centrality of a species. Finally, we included a categorical variable for the species' trophic level 261 (plant or pollinator) and an interaction term between trophic level and the previous four variables. To 262 facilitate comparison between predictors, degree and visitation strength were log-transformed and all 263 four continuous variables were scaled to have a mean of zero and a standard deviation of one. In these 264 models, species from all networks were analysed together. We initially included random effects to account 265 for possible variation across communities. Specifically, we tested structures that allowed for a random 266 intercept for the network, site, and the study it comes from. However, we found that in all cases the 267 among-group variance was effectively zero, and therefore we did not include any random effect in further analyses. We then generated all possible candidate models across the space of models with all, some, and none of the predictor variables. To identify the models that were best supported by the data, we first 270 determined the most parsimonious random structure using the AICc. The relative importance of variables was then assessed by looking at their effect sizes in the top-ranked models and the cumulative weight of 272 the models in which they are present. 273

In addition, we wanted to understand how a species' control capacity ϕ described above relates to metrics
of keystoneness based on centrality. Specifically, in each network, we calculated the species' degree,
betweenness, closeness centrality (Martín González et al., 2010), page rank (McDonald-Madden et al.,
2016), and eigen centrality (Jordano, Bascompte, & Olesen, 2006). We then calculated the spearman
correlation coefficient between control capacity and each of these centrality metrics.

Our analysis revealed that some species have a control capacity $\phi = 1$. These species are critical to 279 controlling their community because they are part of the minimum driver-node set in every control 280 scenario. In other words, it is theoretically impossible to drive the state of the community to a desired 281 state without directly managing the abundance of these species. We thus anticipate that these species have 282 a disproportionally large impact on the community dynamics. To test this hypothesis, we identified these 283 critical species in each of the networks and investigated whether they have a larger than average impact 284 on the stable coexistence of species in the community. Within mutualistic networks, one useful measure of 285 stable coexistence is called structural stability (R. P. Rohr, Saavedra, & Bascompte, 2014). Mathematically, the structural stability of a network represents the size of the parameter space (i.e., growth rates, carrying capacities, etc.) under which all species can sustain positive abundances (S. Saavedra, Rohr, Olesen, & Bascompte, 2016). The contribution of any given species i to stable coexistence can be estimated 289

by calculating the structural stability of the community when the focal species i is removed. To allow comparison across communities, the structural stability values were scaled within each network to have a mean of zero and a standard deviation of one. Given these species-specific estimates of structural stability, we then used a t-test to compare the contribution to stable coexistence of critical and non-critical species.

More details about the calculation of structural stability can be found in the Supporting Information Section S4.

Testing assumptions

Just like the centrality metrics, the information obtained by applying structural controllability depends on 296 the ability of the network to accurately represent the ecological community. We thus tested the sensitivity 297 of our approach to two fundamental assumptions. First, we tested that visitation is an appropriate proxy 298 to infer interspecific effects by comparing the results obtained using visitation to two alternative metrics in a separate dataset that lacked invasive species (Ballantyne, Baldock, & Willmer, 2015). Specifically, we also calculated the controllability (the size of the minimum driver node-set) and the control capacity of networks constructed using pollinator efficiency (which measures the pollen deposition of an interaction) 302 and pollinator importance (which accounts for both pollen deposition and visitation and hence is regarded 303 as a more accurate estimation of the pollination service received by plants; Ne'eman, Jürgens, Newstrom-304 Lloyd, Potts, & Dafni, 2009). More details in the 2010). See Supporting Information Section S5 for more 305 details. 306 Second, because interspecific dependencies themselves depend on the network topology and consequently 307 on the accurate sampling of interactions, we tested the robustness of structural controllability to the 308 uncertainty involved with the sampling of interactions. Here, we compared the results obtained when using the full network and when randomly removing interactions from the weakest links in the network. This effectively removed the rare interactions from the networks (more details in the Supporting Information 311 Section S6). 312

Results

Controllability

The size of the minimum driver-node set relative to the number of species in each network n_D ranged between $n_D = 0.58$ and $n_D = 0.88$ (median 0.74).

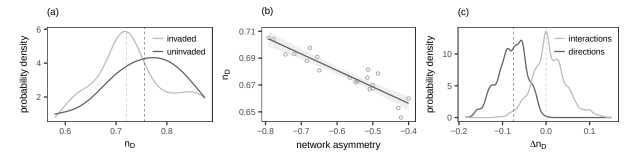


Figure 3: Drivers of network controllability. (a) Probability density of the relative size of the minimum driver-node set n_D in the invaded (light) and uninvaded (dark) empirical networks. (b) Relationship between the asymmetry plant/pollinator richness and n_D . (c) Probability density of the difference between the relative size of the minimum driver-node set of random networks and that of empirical networks. We randomised either the species visitation patterns (light line) or randomised the direction of control between a species pair (dark line). The vertical dashed lines in (a) and (c) indicate the median values of the distributions.

Differences between invaded and uninvaded networks

We found that the relative size of the minimum driver-node set of invaded communities was not significantly different from that of communities that have not been invaded (Figure 3a). In contrast, there was a large negative relationship between n_D and the network asymmetry (Figure 3b). Furthermore, there were also negative, albeit weaker, relationships between n_D and connectance, nestedness and species richness (Table S3). The relative size of the minimum driver-node set n_D of empirical networks did not differ from that of a null model that roughly preserved the degree distribution and fully preserved the network connectance (p = 0.66; Figure 3c). However, empirical networks had a larger n_D than null models that preserved the interactions but shuffled the direction of control of the empirical network ($p = 2.4 \times 10^{-7}$).

Species' roles

Species varied widely in their control capacity (Figure 4). Pollinators had, in average, larger control capacities than plants. That said, almost no pollinator was critical for network control, (where a species is critical for control if it has control capacity $\phi_i = 1$). Plants had a multimodal distribution of control 325 capacity with maxima at both extremes of the distribution (Figure 4a). Intriguingly, every invasive species 326 was critical for network control in each of their communities. The species-level models identified a positive 327 relationship between control capacity ϕ and a species' contribution to nestedness, visitation strength, and 328 the asymmetry of its dependences (Table 2; Figure 5; Table S4). Comparatively, species' degree was only 329 weakly associated with control capacity (Table S5). In fact, many species with a low degree, especially 330 pollinators, exhibited a large control capacity in their communities (Figure S10a). 331

Species' control capacity ϕ was only weakly correlated with commonly-used centrality metrics. The Spearman correlation between these ranged between -0.14 (with betweeness-centrality) and 0.42 (with

Table 2: Selection table of the binomial generalised linear models of species control capacity, ϕ . Only models with a weight larger or equal to 0.01 are shown.

model terms												
int.	k	l	a	n	s	k: l	l: a	l: n	l: s	d.f.	$\Delta {\rm AICc}$	weight
-1.20		+	0.80	0.15	0.29		+	+		7	0.00	0.48
-1.19		+	0.76	0.13	0.35		+	+	+	8	1.52	0.22
-1.26	-1.24	+	1.44	0.39	1.07	+	+		+	9	4.09	0.06
-1.37	-0.66	+	1.03		1.06	+	+		+	8	4.39	0.05
-1.27	-1.15	+	1.37	0.33	1.07	+	+	+	+	10	4.92	0.04
-1.37	-0.10	+	0.90		0.43	+	+			7	6.36	0.02
-1.25	-0.28	+	1.24	0.40		+	+			7	6.47	0.02
-1.24	-0.62	+	1.29	0.38	0.40	+	+			8	6.50	0.02
-1.39	0.30	+	0.83			+	+			6	6.72	0.02
-1.28	-0.17	+	1.16	0.32		+	+	+		8	7.03	0.01
-1.26	-0.53	+	1.23	0.32	0.39	+	+	+		9	7.42	0.01
-1.02		+	0.69	0.30	0.31		+			6	7.48	0.01

Terms: intercept (int), degree (k), trophic level (l), asymmetry (a), contribution to nestedness (n), visitation strength (s).

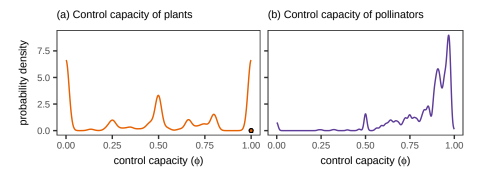


Figure 4: Probability density of the control capacity ϕ of (a) plants and (b) pollinators across all networks. The control capacity of all invasive species is $\phi = 1$ and is depicted with solid circles.

eigen-centrality), see Figure S11a. The correlation coefficient with degree was -0.13, however most species
with high degree also tended to attain a high control capacity (Figure S10a).

Finally, we found that critical species have a particularly large impact on species coexistence when compared to non-critical species. Indeed, the structural stability of the networks where critical species were removed was considerably lower to those in were non-critical species were removed ($p = 2 \times 10^{-15}$; Figure 6; Supporting Information S4).

Testing assumptions

We found that using visitation as a proxy for the strength of species' interactions leads to similar results
than those obtained using pollinator importance (regarded as an accurate measure of the pollination service
to plants; Supporting Information Section S5; Ne'eman et al., 20092010). Importantly, we also found that
structural stability is robust to incomplete sampling of interactions. Indeed, we found strong agreement
between results obtained using the complete empirical networks and those obtained by randomly removing
the weakest interactions (Supporting Information Section S6). Despite removing rare interactions and

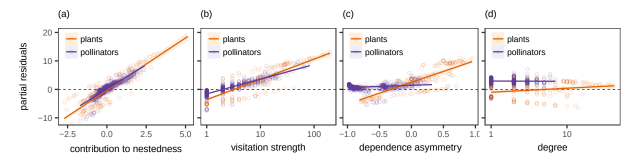


Figure 5: Partial-residual plots for the independent variables: (a) contribution to nestedness, (b) visitation strength, (c) asymmetry of dependences, and (d) degree. Partial-residual plots show the relationship between control capacity and each of the independent variables while accounting for all other remaining variables. Ploted values correpond to the predictions of the weighted average of the models shown in Table 2.

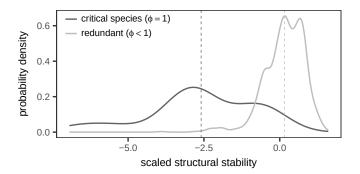


Figure 6: Probability density of the structural stability of the communities after a single focal species is removed. Mathematically, the structural stability of a network represents the size of the parameter space (i.e., growth rates, carrying capacities, etc.) under which all species can sustain positive abundances. The structural stability of communities in which critical species have been removed (darker line) is considerably smaller than that of communities in which non-critical species have been removed. This indicates that critical species contribute more to the stable coexistence of their communities. To allow comparison across communities, the structural stability values were scaled within each network to have a mean of zero and a standard deviation of one. Here, we assume values of the mutualistic trade-off and mean interspecific competition of $\delta = 0$ and $\rho = 0.01$ respectively (S.–Saavedra et al., 2016). However, the choice of these parameters does not affect the results (Supporting Information S4).

species, the relative size of the minimum driver-node set, the superior species, and the relative rankings of control capacity were generally maintained. Of particular note, we found that critical species in the full network were also critical in the vast majority of rarefied networks.

Discussion

Our main goal was to understand the role that species play at in both modifying the abundance of the 349 species they interact with and the state of the community as a whole. To achieve that goal we applied 350 structural controllability, a field at the intersection between control and complex theory that allow-allowed 351 us to obtain two key pieces of information: the controllability of a network and a species' control capacity 352 (Table 1). We found that the controllability of a network does not depend on its invasion status and that 353 the species that are critical to altering the state of the community are also the ones that most sustain the 354 stable coexistence of species in their communities. 355 Glossary network control — A network is said to be controllable if it is possible to steer it from an initial 356 to an arbitrary final state within finite time. controllability — The intrinsic difficulty of controlling 357 an ecological community. It is measured by the relative size of the minimum driver-node set, n_D . It 358 also indicates the extent to which network structure can be harnessed for network control. minimum 359 driver-node set — One of the sets of species whose abundances need to be directly managed in order 360 to achieve full control of the community. The minimum driver-node sets can be obtained by finding all 361 maximum matchings in a network. maximum matching — A matching is a set of links that do not 362 share any common start or end nodes; the largest possible matching is called a maximum matching. 363 control configuration — One of the species combinations with which is possible to achieve network 364 control. Optimal control configurations are given by the minimum driver-node sets. control capacity The relative frequency ϕ which with a species is part of the optimal control configurations of a network. 366 critical species — A species with a maximal control capacity $\phi = 1$ superior node — A species is a 367 superior node if it can internally affect the abundance of other species in the network. Superior nodes 368 make up the chains that propagate the control signals through the network. 369 Our results indicate that fully controlling ecological networks might currently be out of reach for all 370 but the smallest communities (A. E. Motter, 2015). Indeed, the median size of the relative minimum 371 driver-node set in our dataset was $n_D = 0.74$, a high value when compared to other complex systems 372 in which controllability has been investigated (the lower n_D the more controllable the community). For instance, only gene regulation networks appear to achieve similar levels of controllability while most social, power transmission, Internet, neuronal, and even metabolic networks seem to be "easier" to control

 $(0.1 < n_D < 0.35)$ (Liu et al., 2011). Structural controllability provides solid theoretical rationale for the 376 many difficulties encountered in the management and restoration of natural communities (Woodford et 377 al., 2016). Nevertheless, structural controllability might be helpful at identifying communities in which 378 changes in the ecological state are more likely to occur. 379 The differences between the controllability across networks are likely to arise from differences in their 380 structure rather than their invasion status. Specifically, when controlling for network structure, we found 381 no difference between the controllability of invaded and uninvaded networks. instead controllability 382 is almost completely constrained by the patterns of species richness at each trophic guild and their 383 degree distributions (N. Blüthgen et al., 2007; C. J. Melián & Bascompte, 2002). These two factors are 384 particularly relevant because they govern the asymmetric nature of mutual dependences, which themselves 385 provide the foundation of structure and stability in mutualistic networks (Astegiano, Massol, Vidal, Cheptou, & Guimarães, 2015; J.-Bascompte et al., 2006; J.-Memmott, Waser, & Price, 2004). 387 Accordingly, our results suggest that structural controllability is closely related to the dynamic persistence 388 of an ecological community based on two lines of evidence. First, we found a comparatively small 389 but thought-provoking negative relationship between the controllability of a network and its nestedness. 390 Previous studies indicate that nestedness promotes species coexistence and confers robustness to extinction 391 (Bastolla et al., 2009; J. Memmott et al., 2004) even at the expense of the dynamic stability of the 392 mutualistic community (S. Saavedra et al., 2016). These observations are in agreement with our results, 393 as we would expect the dynamic stability (the ability to return to equilibrium after a perturbation in 394 species abundances) of a community to be correlated to the difficulty to control it. Second, species' control capacity was strongly correlated to their contribution to nestedness and critical species had the largest impact to the stable coexistence of species in their communities. Therefore, species that play a key role at determining the state of the community might also be more key to "maintain the organization and diversity of their ecological communities", one of the hallmarks of keystone species (Mills & Doak, 1993). 399 When controlling for a species' strength species' visitation strength (the sum of a species' visits), which 400 is indirectly a proxy of its abundance, and the net balance of its dependencies, we found that control 401 capacity could not be easily predicted by species' degree or other metrics of centrality. For instance, some 402 species with a low degree achieved the maximum control capacity and were critical for control in their 403 communities. At first glance, our findings challenge numerous studies that highlight the role that central species play in the dynamics of their communities and their utility at predicting species extinctions (Jordan, 2009). However, further -inspection shows that our results do not contradict these findings; most 406 species with a large degree also have a large control capacity and all of them were classified as superior 407 nodes which corroborates the utility of classic centrality metrics. Putting these observations together, 408

our results therefore take previous findings one step further and suggest that centrality might paint an

incomplete picture of the relevance of species.

Other conceptual differences between structural controllability and centrality metrics provide three key 411 insights into the conservation of ecological networks. First, structural controllability emphasizes that the effect a species has on the abundance of other species is not independent of the effects of other other species in their that the other species have in the community. The rankings provided by centrality 414 metrics and other heuristics fail to account for the collective influence of several species at once. Second, 415 it demonstrates that to ensure the persistence of a community it is often necessary to consider the 416 abundances of more than a single species, even when full control is infeasible or undesired (for example 417 90% of our communities contained more than one critical species). Third, structural controllability 418 explicitly acknowledges the existence of multiple management strategies and some will be better than 419 others depending on the context. Approaches to prioritise species for conservation and reintroduction 420 based on traits or centrality are still useful and are likely to overlap with species control capacity' control 421 capacity (Devoto, Bailey, Craze, & Memmott, 2012; Pires, Marquitti, & Guimarães, 2017). Stepping back, our results also provide support to the idea that management decisions should not be based on a single technique but indicate that focusing on ecosystem processes and interactions may be more effective than 424 traditional ranking-based approaches (Harvey, Gounand, Ward, & Altermatt, 2017). As much potential 425 as any metric or metrics to summarise species' importance might appear to have, it's clear that we also 426 need more empirical studies in different types of networks in order to build intuition and ground truth 427 their usefulness. 428 Our choice of studying invaded/uninvaded networks was based on a desire to contrast the extensive empirical evidence of the role of invasive plants with our theoretical results. We found that invasive plants were always critical for network control and as such our results were in line with our expectations. Invasive plants have been previously found to exacerbate the asymmetries in their communities (Aizen, Morales, & Morales, 2008; Bartomeus et al., 2008; Henriksson, Wardle, Trygg, Diehl, & Englund, 2016) and to be 433 central and to attain high centrality in their communities (Palacio, Valderrama-Ardila, & Kattan, 2016; 434 Vila et al., 2009). We found that invasive plants are, however, unlikely to be inherently different from that 435 it is not that invasive plants have some different mechanism for influencing the community compared to 436 their native counterparts (Emer, Memmott, Vaughan, Montoya, & Tylianakis, 2016; Stouffer, Cirtwill, & 437 Bascompte, 2014). Just like any other mutualist in our dataset, Both native species and invasive plants 438 tended to attain a high control capacity proportional to the degree to which they contribute if they 439 were important to network persistence, are were abundant, and depended little on other species. Furthermore, our observation that changes in the abundance of invasive plants (and presumably all critical species) are crucial to modify the state of the community agrees with recent evidence showing that ecosystem restoration focused on the eradication of invasive plants can have transformative desirable 443

effects in plant-pollinator communities (Kaiser-Bunbury et al., 2017). However, our results also suggest 444 that removals must be exercised with caution. Not only it is hard to predict the direction in which the system will change, but we also show that critical species can underpin the coexistence of species and therefore some communities may be acutely vulnerable to their eradication (Albrecht, Padron, Bartomeus, & Traveset, 2014; Traveset et al., 2013). Structural controllability assumes that the networks can be approximated using linear functional responses 449 (Liu & Barabási, 2016). The ramifications of this assumption imply that, while structural controllability 450 is useful to identify species that are relevant for network control, it cannot be used to design the 451 exact interventions that should be applied to these species in order to achieve a desired state. In an 452 ideal scenario, we would completely incorporate the species' dynamics into the controllability analysis 453 (Cornelius, Kath, & Motter, 2013); the reality is that such information is rarely available in most ecological 454 scenarios. In contrast, structural controllability only requires a quantitative approximation of the network's interactions to gain valuable insight from the community. Furthermore, while the relationship between centrality and keystoneness is based on an intuitive understanding of what a keystone species is, the assumptions of structural controllability are explicit and the estimation of a species' importance arises 458 from a mechanistic understanding of the population dynamics between species. By accounting for network 459 dynamics (even if in a simple way), structural stability incorporates more ecological realism, especially in 460 the extreme scenario in which the state of a community is only marginally affected by the topology of 461 their interactions structure of interactions within the community only marginally affects the community's 462 $\underbrace{\text{state}}_{\cdot}$. 463

Conclusions

Here we show that structural controllability can be applied in an ecological setting to gain insight into the stability of a community and the role that species play at in modifying the abundance of other species and ultimately the state of the community. These characteristics make structural stability an ideal framework to evaluate the effects of invasions and other types of perturbations. Importantly, structural controllability 467 can be used to identify critical species in the community that promote biodiversity and underpin the 468 stable coexistence of species in their community. Collectively, critical species dominate the have the 469 ability to strongly influence the state of their community and therefore are likely to be highly relevant for 470 ecosystem management and conservation. While useful, centrality metrics, which metrics—which have 471 often been used as a proxy for keystoneness, fail keystoneness—fail to identify some of these species, 472 highlighting their limitations when we fully embrace the notion that ecological communities are dynamical 473 systems. Paine (1969) showed nearly 50 years ago that one a single species can dominate the state of its

- community sometimes fill this role on its own. Structural controllability suggests that this situation might
- be the exception rather than the rule. We see our study as a starting point to study the controllability of
- ecological and socio-ecological systems where many exciting questions lie ahead.

Acknowledgements

- 478 The authors thank Jane Memmott and co-authors, and everyone that has made their data available
- 479 to us, Takeuki Uno for the insight provided to find the set of all maximum matching algorithms, and
- 480 Jason Tylianakis, Bernat Bramon Mora, Guadalupe Peralta, Rogini Runghen, Michelle Marraffini, Mark
- 481 Herse, Warwick Allen, Matthew Hutchinson, and Marilia Gaiarsa for feedback and valuable discussions.
- 482 EFC acknowledges the support from the University of Canterbury Doctoral Scholarship, the University
- 483 of Canterbury Meadow Mushrooms Postgraduate Scholarship, a New Zealand International Doctoral
- Research Scholarship, and a travel grant from the European Space Agency. DBS acknowledges the support
- of a Marsden Fast-Start grant and a Rutherford Discovery Fellowship, administered by the Royal Society
- 486 Te Aparangi.

Author contributions

- DBS conceived the idea; all authors contributed to the development of the theoretical framework. EFC
- performed all analysis. EFC and DBS wrote the manuscript. All authors contributed to its revision.

Data accessibility

- 489 All data used in this manuscript have already been published by Lopezaraiza-Mikel et al. (2007),
- Bartomeus et al. (2008), and Ballantyne et al. (2015) The reader should refer to the original sources to
- access the data.

References

- ⁴⁹² Aizen, M. A., Morales, C. L., & Morales, J. M. (2008). Invasive Mutualists Erode Native Pollination
- ⁴⁹³ Webs. *PLoS Biology*, 6(2), e31. doi:10.1371/journal.pbio.0060031
- ⁴⁹⁴ Albrecht, M., Padron, B., Bartomeus, I., & Traveset, A. (2014). Consequences of plant invasions on
- compartmentalization and species' roles in plant-pollinator networks. Proceedings of the Royal Society B:
- ⁴⁹⁶ Biological Sciences, 281(1788), 20140773–20140773. doi:10.1098/rspb.2014.0773
- ⁴⁹⁷ Astegiano, J., Massol, F., Vidal, M. M., Cheptou, P.-O., & Guimarães, P. R. (2015). The Robustness
- of Plant-Pollinator Assemblages: Linking Plant Interaction Patterns and Sensitivity to Pollinator Loss.
- 499 PLOS ONE, 10(2), e0117243. doi:10.1371/journal.pone.0117243
- 500 Ballantyne, G., Baldock, K. C. R., & Willmer, P. G. (2015). Constructing more informative plantPollinator
- plant-pollinator networks: Visitation and pollen deposition networks in a heathland plant community.
- ₅₀₂ Proceedings of the Royal Society B: Biological Sciences, 282(1814), 20151130. doi:10.1098/rspb.2015.1130
- Bartomeus, I., Vilà, M., & Santamaría, L. (2008). Contrasting effects of invasive plants in plantPollinator
- plant-pollinator networks. Oecologia, 155(4), 761–770. doi:10.1007/s00442-007-0946-1
- ⁵⁰⁵ Bascompte, J., & Stouffer, D. B. (2009). The assembly and disassembly of ecological networks. *Philosophical*
- Transactions of the Royal Society B: Biological Sciences, 364(1524), 1781–1787. doi:10.1098/rstb.2008.0226
- Bascompte, J., Jordano, P., & Olesen, J. M. (2006). Asymmetric Coevolutionary Networks Facilitate
- 508 Biodiversity Maintenance. Science, 312(5772), 431–433. doi:10.1126/science.1123412
- Bastolla, U., Fortuna, M. A., Pascual-García, A., Ferrera, A., Luque, B., & Bascompte, J. (2009). The
- architecture of mutualistic networks minimizes competition and increases biodiversity. Nature, 458(7241),
- ⁵¹¹ 1018–1020. doi:10.1038/nature07950
- Blüthgen, N., Menzel, F., Hovestadt, T., Fiala, B., & Blüthgen, N. (2007). Specialization, Con-
- 513 straints, and Conflicting Interests in Mutualistic Networks. Current Biology, 17(4), 341–346.
- ⁵¹⁴ doi:10.1016/j.cub.2006.12.039
- ⁵¹⁵ Cornelius, S. P., Kath, W. L., & Motter, A. E. (2013). Realistic control of network dynamics. Nature
- 516 Communications, 4, 1942. doi:10.1038/ncomms2939
- ⁵¹⁷ Coux, C., Rader, R., Bartomeus, I., & Tylianakis, J. M. (2016). Linking species functional roles to their
- network roles. Ecology Letters, 19(7), 762–770. doi:10.1111/ele.12612
- 519 Devoto, M., Bailey, S., Craze, P., & Memmott, J. (2012). Understanding and planning ecological restora-
- tion of plant-pollinator networks: Understanding network restoration. Ecology Letters, 15(4), 319–328.

- doi:10.1111/j.1461-0248.2012.01740.x
- 522 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. doi:10.1046/j.1461-
- 524 0248.2002.00354.x
- Emer, C., Memmott, J., Vaughan, I. P., Montoya, D., & Tylianakis, J. M. (2016). Species roles in
- plant-pollinator communities are conserved across native and alien ranges. Diversity and Distributions,
- ⁵²⁷ 22(8), 841–852. doi:10.1111/ddi.12458
- Friedkin, N. E. (1991). Theoretical Foundations for Centrality Measures. American Journal of Sociology,
- 96(6), 1478–1504.
- ⁵³⁰ Guimerà, R., & Amaral, L. A. N. (2005). Cartography of complex networks: Modules and universal
- roles. Journal of Statistical Mechanics: Theory and Experiment, 2005(02), P02001. doi:10.1088/1742-
- 532 5468/2005/02/P02001
- Harvey, E., Gounand, I., Ward, C. L., & Altermatt, F. (2017). Bridging ecology and conservation: From
- ecological networks to ecosystem function. Journal of Applied Ecology, 54(2), 371–379. doi:10.1111/1365-
- 535 2664.12769
- Henriksson, A., Wardle, D. A., Trygg, J., Diehl, S., & Englund, G. (2016). Strong invaders are strong
- defenders implications for the resistance of invaded communities. Ecology Letters, 19(4), 487–494.
- 538 doi:10.1111/ele.12586
- Holland, J. N., DeAngelis, D. L., & Bronstein, J. L. (2002). Population Dynamics and Mutualism:
- Functional Responses of Benefits and Costs. The American Naturalist, 159(3), 231–244. doi:10.1086/338510
- Isbell, F., & Loreau, M. (2013). Human impacts on minimum subsets of species critical for maintaining
- ecosystem structure. Basic and Applied Ecology, 14(8), 623–629. doi:10.1016/j.baae.2013.09.001
- Jia, T., & Barabási, A.-L. (2013). Control Capacity and A Random Sampling Method in Exploring
- ⁵⁴⁴ Controllability of Complex Networks. Scientific Reports, 3(1). doi:10.1038/srep02354
- Jordan, F. (2009). Keystone species and food webs. Philosophical Transactions of the Royal Society B:
- 546 Biological Sciences, 364(1524), 1733-1741. doi:10.1098/rstb.2008.0335
- Jordano, P., Bascompte, J., & Olesen, J. M. (2006). The ecological consequences of complex topology and
- nested structure in pollination webs. In N. M. Waser & J. Ollerton (Eds.), Plant-Pollinator Interactions:
- From Specialization to Generalization (pp. 173–199). University of Chicago Press.
- Jordán, F., Benedek, Z., & Podani, J. (2007). Quantifying positional importance in food webs: A comparison

- of centrality indices. Ecological Modelling, 205(1-2), 270–275. doi:10.1016/j.ecolmodel.2007.02.032
- 552 Kaiser-Bunbury, C. N., Mougal, J., Whittington, A. E., Valentin, T., Gabriel, R., Olesen, J. M., &
- Blüthgen, N. (2017). Ecosystem restoration strengthens pollination network resilience and function.
- 554 Nature, 542(7640), 223–227. doi:10.1038/nature21071
- Kaiser-Bunbury, C. N., Muff, S., Memmott, J., Müller, C. B., & Caflisch, A. (2010). The robustness
- of pollination networks to the loss of species and interactions: A quantitative approach incorporating
- ₅₅₇ pollinator behaviour. Ecology Letters, 13(4), 442–452. doi:10.1111/j.1461-0248.2009.01437.x
- Kalman, R. E. (1963). Mathematical Description of Linear Dynamical Systems. Journal of the Society for
- 559 Industrial and Applied Mathematics Series A Control, 1(2), 152–192. doi:10.1137/0301010
- Lever, J. J., van Nes, E. H., Scheffer, M., & Bascompte, J. (2014). The sudden collapse of pollinator
- 561 communities. Ecology Letters, 17(3), 350–359. doi:10.1111/ele.12236
- Lin, C. T. (1974). Structural Controllability. IEEE Transactions on Automatic Control, 19(3), 201–208.
- ⁵⁶³ doi:10.1109/TAC.1974.1100557
- Liu, Y.-Y., & Barabási, A.-L. (2016). Control principles of complex systems. Reviews of Modern Physics,
- 565 88(3). doi:10.1103/RevModPhys.88.035006
- Liu, Y.-Y., Slotine, J.-J., & Barabási, A.-L. (2011). Controllability of complex networks. Nature, 473(7346),
- ⁵⁶⁷ 167–173. doi:10.1038/nature10011
- Lopezaraiza-Mikel, M. E., Hayes, R. B., Whalley, M. R., & Memmott, J. (2007). The impact of an alien
- plant on a native plantPollinator network: An experimental approach. Ecology Letters, 10(7), 539–550.
- oi:10.1111/j.1461-0248.2007.01055.x
- Martín González, A. M., Dalsgaard, B., & Olesen, J. M. (2010). Centrality measures and the
- importance of generalist species in pollination networks. Ecological Complexity, 7(1), 36–43.
- 573 doi:10.1016/j.ecocom.2009.03.008
- McDonald-Madden, E., Sabbadin, R., Game, E. T., Baxter, P. W. J., Chadès, I., & Possingham,
- H. P. (2016). Using food-web theory to conserve ecosystems. Nature Communications, 7, 10245.
- 576 doi:10.1038/ncomms10245
- Melián, C. J., & Bascompte, J. (2002). Complex networks: Two ways to be robust?: Complex networks:
- Two ways to be robust? *Ecology Letters*, 5(6), 705–708. doi:10.1046/j.1461-0248.2002.00386.x
- Memmott, J., Waser, N. M., & Price, M. V. (2004). Tolerance of pollination networks to species extinctions.
- Proceedings of the Royal Society B: Biological Sciences, 271(1557), 2605–2611. doi:10.1098/rspb.2004.2909
- Menge, B. A. (1995). Indirect Effects in Marine Rocky Intertidal Interaction Webs: Patterns and

- ⁵⁸² Importance. Ecological Monographs, 65(1), 21–74. doi:10.2307/2937158
- Mills, L. S., & Doak, D. F. (1993). The Keystone-Species Concept in Ecology and Conservation. BioScience,
- ⁵⁸⁴ 43(4), 219–224. doi:10.2307/1312122
- ⁵⁸⁵ Motter, A. E. (2015). Networkcontrology. Chaos, 25, 097621. doi:10.1063/1.4931570
- Mouillot, D., Graham, N. A., Villéger, S., Mason, N. W., & Bellwood, D. R. (2013). A functional
- ⁵⁸⁷ approach reveals community responses to disturbances. Trends in Ecology & Evolution, 28(3), 167–177.
- 588 doi:10.1016/j.tree.2012.10.004
- Ne'eman, G., Jürgens, A., Newstrom-Lloyd, L., Potts, S. G., & Dafni, A. (20092010). A framework for com-
- paring pollinator performance: Effectiveness and efficiency. Biological Reviews, no-no. doi:10.1111/j.1469-
- ⁵⁹¹ 185X.2009.00108.x
- ⁵⁹² Paine, R. T. (1969). A Note on Trophic Complexity and Community Stability. The American Naturalist,
- ⁵⁹³ 103(929), 91–93.
- ⁵⁹⁴ Palacio, R. D., Valderrama-Ardila, C., & Kattan, G. H. (2016). Generalist Species Have a Central Role
- In a Highly Diverse Plant-Frugivore Network. Biotropica, 48(3), 349–355. doi:10.1111/btp.12290
- ⁵⁹⁶ Pires, M. M., Marquitti, F. M., & Guimarães, P. R. (2017). The friendship paradox in species-rich
- ecological networks: Implications for conservation and monitoring. Biological Conservation, 209, 245–252.
- ⁵⁹⁸ doi:10.1016/j.biocon.2017.02.026
- Poisot, T., Canard, E., Mouquet, N., & Hochberg, M. E. (2012). A comparative study of ecological
- specialization estimators: Species-Level Specialization. Methods in Ecology and Evolution, 3(3), 537–544.
- 601 doi:10.1111/j.2041-210X.2011.00174.x
- Rohr, R. P., Saavedra, S., & Bascompte, J. (2014). On the structural stability of mutualistic systems.
- Science, 345(6195), 1253497-1253497. doi:10.1126/science.1253497
- 604 Saavedra, S., Rohr, R. P., Olesen, J. M., & Bascompte, J. (2016). Nested species interactions promote
- feasibility over stability during the assembly of a pollinator community. Ecology and Evolution, 6(4),
- 997-1007. doi:10.1002/ece3.1930
- 607 Saavedra, S., Stouffer, D. B., Uzzi, B., & Bascompte, J. (2011). Strong contributors to network persistence
- are the most vulnerable to extinction. Nature, 478(7368), 233–235. doi:10.1038/nature10433
- Stouffer, D. B., Cirtwill, A. R., & Bascompte, J. (2014). How exotic plants integrate into pollination
- ${}_{610}\quad \text{networks. } \textit{Journal of Ecology},\ 102(6),\ 1442-1450.\ doi: 10.1111/1365-2745.12310$
- 611 Stouffer, D. B., Sales-Pardo, M., Sirer, M. I., & Bascompte, J. (2012). Evolutionary Conservation of

- 612 Species' Roles in Food Webs. Science, 335(6075), 1489–1492. doi:10.1126/science.1216556
- Thompson, R. M., Brose, U., Dunne, J. A., Hall, R. O., Hladyz, S., Kitching, R. L., ... Tylianakis, J. M.
- 614 (2012). Food webs: Reconciling the structure and function of biodiversity. Trends in Ecology & Evolution,
- 615 27(12), 689–697. doi:10.1016/j.tree.2012.08.005
- Traveset, A., Heleno, R., Chamorro, S., Vargas, P., McMullen, C. K., Castro-Urgal, R., ... Olesen,
- ₆₁₇ J. M. (2013). Invaders of pollination networks in the Galapagos Islands: Emergence of novel com-
- munities. Proceedings of the Royal Society B: Biological Sciences, 280(1758), 20123040–20123040.
- doi:10.1098/rspb.2012.3040
- ⁶²⁰ Tylianakis, J. M., Didham, R. K., Bascompte, J., & Wardle, D. A. (2008). Global change and species inter-
- 621 actions in terrestrial ecosystems. Ecology Letters, 11(12), 1351–1363. doi:10.1111/j.1461-0248.2008.01250.x
- Tylianakis, J. M., Laliberté, E., Nielsen, A., & Bascompte, J. (2010). Conservation of species interaction
- networks. Biological Conservation, 143(10), 2270-2279. doi:10.1016/j.biocon.2009.12.004
- Vázquez, D. P., Melián, C. J., Williams, N. M., Blüthgen, N., Krasnov, B. R., & Poulin, R. (2007). Species
- 625 Abundance and Asymmetric Interaction Strength in Ecological Networks Author(s): Diego P. Vázquez,
- ⁶²⁶ Carlos J. Melián, Neal M. Williams, Nico Blüthgen, Boris R. Krasnov and Robert Poulin. Oikos, 116(7),
- 627 1120-1127. doi:10.1111/j.2007.0030-1299.15828.x
- Vázquez, D. P., Morris, W. F., & Jordano, P. (2005). Interaction frequency as a surrogate for the total
- effect of animal mutualists on plants: Total effect of animal mutualists on plants. Ecology Letters, 8(10),
- 630 1088–1094. doi:10.1111/j.1461-0248.2005.00810.x
- Vila, M., Bartomeus, I., Dietzsch, A. C., Petanidou, T., Steffan-Dewenter, I., Stout, J. C., & Tscheulin, T.
- 632 (2009). Invasive plant integration into native plant-pollinator networks across Europe. Proceedings of the
- 633 Royal Society B: Biological Sciences, 276(1674), 3887–3893. doi:10.1098/rspb.2009.1076
- Woodford, D. J., Richardson, D. M., MacIsaac, H. J., Mandrak, N. E., van Wilgen, B. W., Wilson, J. R.
- U., & Weyl, O. L. F. (2016). Confronting the wicked problem of managing biological invasions. NeoBiota,
- 636 31, 63–86. doi:10.3897/neobiota.31.10038
- ⁶⁵⁷ Zhang, X., Lv, T., & Pu, Y. (2016). Input graph: The hidden geometry in controlling complex networks.
- 638 Scientific Reports, 6(1). doi:10.1038/srep38209