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Rudolf P. Rohr *et al.*
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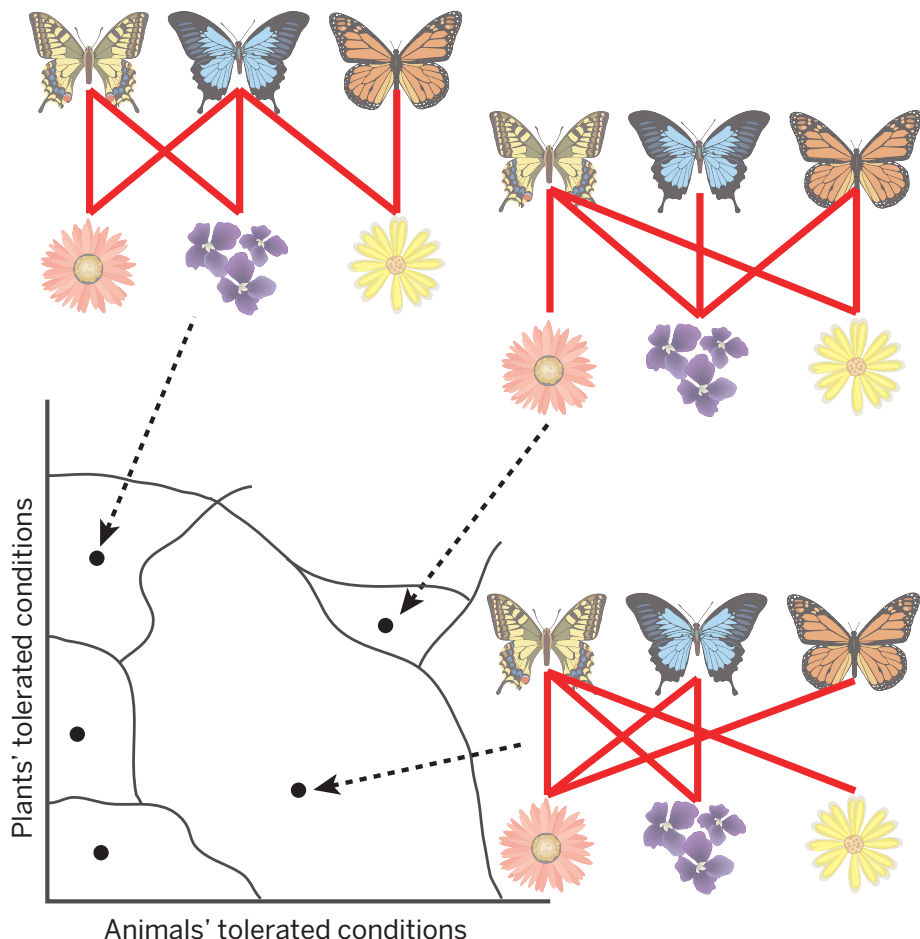
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RESEARCH ARTICLE SUMMARY

ECOLOGICAL NETWORKS

On the structural stability of mutualistic systems

Rudolf P. Rohr,^{1,2} Serguei Saavedra,¹ Jordi Bascompte^{1*}

The architecture of plant-animal mutualistic networks modulates the range of conditions leading to the stable coexistence of all species. The area of the different domains represents the structural stability of a model of mutualistic communities with a given network architecture. The nested networks observed in nature—illustrated here by the network at the bottom—lead to a maximum structural stability.

INTRODUCTION: Several major developments in theoretical ecology have relied on either dynamical stability or numerical simulations, but oftentimes, they have found contradictory results. This is partly a result of not rigorously checking either the assumption that a steady state is feasible—meaning, all species have constant and positive abundances—or the dependence of results to model parameterization. Here, we extend the concept of structural

stability to community ecology in order to account for these two problems. Specifically, we studied the set of conditions leading to the stable coexistence of all species within a community. This shifts the question from asking whether we can find a feasible equilibrium point for a fixed set of parameter values, to asking how large is the range of parameter values that are compatible with the stable coexistence of all species.

RATIONALE: We begin by disentangling the conditions of global stability from the conditions of feasibility of a steady state in ecological systems. To quantify the domain of stable coexistence, we first find its center (the structural vector of intrinsic growth rates). Next, we determine the boundaries of such a domain by quantifying the amount of variation from the structural vector tolerated before one species goes extinct. Through this two-step approach, we disentangle the effects of the size of the feasibility domain from how close a solution is to its boundary, which is at the heart of previous contradictory results. We illustrate our method by exploring how the observed architecture of mutualistic networks between plants and their pollinators or seed dispersers affects their domain of stable coexistence.

RESULTS: First, we determined the network architecture that maximizes the structural stability of mutualistic systems. This corresponds to networks with a maximal level of nestedness, a small trade-off between the number and intensity of interactions a species has, and a high level of mutualistic strength within the constraints of global stability. Second, we found that the large majority of observed mutualistic networks are close to this optimum network architecture, maximizing the range of parameters that are compatible with species coexistence.

CONCLUSION: Structural stability has played a major role in several fields such as evolutionary developmental biology, in which it has brought the view that some morphological structures are more common than others because they are compatible with a wider range of developmental conditions. In community ecology, structural stability is the sort of framework needed to study the consequences of global environmental change—by definition, large and directional—on species coexistence. Structural stability will serve to assess both the range of variability a given community can withstand and why some community patterns are more widespread than others. ■

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RESEARCH ARTICLE

ECOLOGICAL NETWORKS

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Rudolf P. Rohr,^{1,2*} Serguei Saavedra,^{1*} Jordi Bascompte^{1†}

In theoretical ecology, traditional studies based on dynamical stability and numerical simulations have not found a unified answer to the effect of network architecture on community persistence. Here, we introduce a mathematical framework based on the concept of structural stability to explain such a disparity of results. We investigated the range of conditions necessary for the stable coexistence of all species in mutualistic systems. We show that the apparently contradictory conclusions reached by previous studies arise as a consequence of overseeing either the necessary conditions for persistence or its dependence on model parameterization. We show that observed network architectures maximize the range of conditions for species coexistence. We discuss the applicability of structural stability to study other types of interspecific interactions.

A prevailing question in ecology [particularly since May's seminal work in the early 1970s (1)] is whether, given an observed number of species and their interactions, there are ways to organize those interactions that lead to more persistent communities. Conventionally, studies addressing this question have either looked into local stability or used numerical simulations (2–4). However, these studies have not yet found a unified answer (1, 5–12). Therefore, the current challenge is to develop a general framework in order to provide a unified assessment of the implications of the architectural patterns of the networks we observe in nature.

Main approaches in theoretical ecology

Dynamical stability and feasibility

Studies based on the mathematical notions of local stability, *D*-stability, and global stability have advanced our knowledge on what makes ecological communities stable. In particular, these studies explore how interaction strengths need to be distributed across species so that an assumed feasible equilibrium point can be stable (1–4, 13–17). By definition, a feasible equilibrium point is that in which all species have a constant positive abundance across time. A negative abundance makes no sense biologically, and an abundance of zero would correspond to an extinct species.

The dynamical stability of a feasible equilibrium point corresponds to the conditions under which the system returns to the equilibrium point

after a perturbation in species abundance. Local stability, for instance, looks at whether a system will return to an assumed feasible equilibrium after an infinitesimally small perturbation (1–3, 13). *D*-stability, in turn, looks at the local stability of any potential feasible equilibrium that the system may have (15–17). More generally, global stability looks at the stability of any potential feasible equilibrium point after a perturbation of any given amplitude (14–17). A technical definition of these different types of dynamical stability and their relationship is provided in (18).

In most of these stability studies, however, a feasible equilibrium point is always assumed without rigorously studying the set of conditions allowing its existence (5, 14, 15, 19). Yet in any given system, we can find examples in which we satisfy only one, both, or none of the feasibility and stability conditions (3, 16, 17, 19). This means that without a proper consideration of the feasibility conditions, any conclusion for studying the stable coexistence of species is based on a system that may or may not exist (3, 5, 19).

To illustrate this point, consider the following textbook example of a two-species competition system

$$\begin{cases} \frac{dN_1}{dt} = N_1(\alpha_1 - \beta_{11}N_1 - \beta_{12}N_2) \\ \frac{dN_2}{dt} = N_2(\alpha_2 - \beta_{21}N_1 - \beta_{22}N_2) \end{cases} \quad (1)$$

where N_1 and N_2 are the abundances of species 1 and 2; β_{11} and β_{22} are their intraspecific competition strengths; β_{12} and β_{21} are their interspecific competition strengths; and α_1 and α_2 are their intrinsic growth rates. An equilibrium point of the system is a pair of abundances N_1^* and N_2^* that makes the right side of the ordinary differential equation system equal to zero.

Although the only condition necessary to guarantee the global stability of any feasible equilibrium point in this system is that the interspecific

competition strengths are lower than the intraspecific ones ($\beta_{12}\beta_{21} < \beta_{11}\beta_{22}$), the feasibility conditions are given by $N_1^* = \frac{\beta_{22}\alpha_1 - \beta_{12}\alpha_2}{\beta_{11}\beta_{22} - \beta_{12}\beta_{21}} > 0$ and $N_2^* = \frac{\beta_{11}\alpha_2 - \beta_{21}\alpha_1}{\beta_{11}\beta_{22} - \beta_{12}\beta_{21}} > 0$ (3, 4, 19). This implies that if we set, for example, $\beta_{11} = \beta_{22} = 1$, $\beta_{12} = \beta_{21} = 0.5$, $\alpha_1 = 1$, and $\alpha_2 = 2$, we fulfill the stability condition but not the feasibility condition, whereas if we set $\beta_{11} = \beta_{22} = 0.5$, $\beta_{12} = \beta_{21} = 1$, and $\alpha_1 = \alpha_2 = 1$, we can satisfy the feasibility condition but not the stability one. To have a stable and feasible equilibrium point, we need to set, for instance, $\beta_{11} = \beta_{22} = 1$, $\beta_{12} = \beta_{21} = 0.5$, and $\alpha_1 = \alpha_2 = 1$ (a graphical illustration is provided in Fig. 1).

The example above confirms the importance of verifying both the stability and the feasibility conditions of the equilibrium point when analyzing the stable coexistence of species (3–5, 19). Of course, we can always fine-tune the parameter values of intrinsic growth rates so that the system is feasible (16, 17). This strategy, for example, has been used when studying the success probability of an invasive species (20). However, when fixing the parameter values of intrinsic growth rates, we are not anymore studying the overall effect of interspecific interactions on the stable coexistence of species. Rather, we are answering the question of how interspecific interactions increase the persistence of species for a given parameterization of intrinsic growth rates. As we will show below, this is also the core of the problem in studies that are based on arbitrary numerical simulations.

Numerical simulations

Numerical simulations have provided an alternative and useful tool with which to explore species coexistence in large ecological systems in which analytical solutions are precluded (3). With this approach, one has as a prerequisite to parameterize the dynamical model, or at least to have a good estimate of the statistical distribution from which these parameters should be sampled. However, if one chooses an arbitrary parameterization without an empirical justification, any study has a high chance of being inconclusive for real ecosystems because species persistence is strongly dependent on the chosen parameterization.

To illustrate this point, we simulated the dynamics of an ecological model (6) with three different parameterizations of intrinsic growth rates (21). Additionally, these simulations were performed over an observed mutualistic network of interactions between flowering plants and their pollinators located in Hickling Norfolk, UK (table S1), a randomized version of this observed network, and the observed network without mutualistic interactions (we assume that there is only competition among plants and among animals). As shown in Fig. 2, it is possible to find a set of intrinsic growth rates so that any network that we analyze is completely persistent and, at the same time, the alternative networks are less persistent.

This observation has two important implications. First, this means that by using different parameterizations for the same dynamical model

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and network of interactions, one can observe from all to a few of the species surviving. Second, this means that each network has a limited range of parameter values under which all species coexist. Thus, by studying a specific parameterization, for instance, one could wrongly conclude that a random network has a greater effect on community persistence than that of an observed network, or vice versa (10–12). This sensitivity to parameter values clearly illustrates that the conclusions that arise from studies that use arbitrary values in intrinsic growth rates are not about the effects of network architecture on species coexistence, but about which network architecture maximizes species persistence for that specific parameterization.

Traditional studies focusing on either local stability or numerical simulations can lead to apparently contradictory results. Therefore, we need a different conceptual framework to unify results and seek for appropriate generalizations.

Structural stability

Structural stability has been a general mathematical approach with which to study the behavior of dynamical systems. A system is considered to be structurally stable if any smooth change in the model itself or in the value of its parameters does not change its dynamical behavior (such as the existence of equilibrium points, limit cycles, or deterministic chaos) (22–25). In the context of ecology, an interesting behavior is the stable coexistence of species—the existence of an equilibrium point that is feasible and dynamically stable. For instance, in our previous two-species competition system there is a restricted area in the parameter space of intrinsic growth rates that leads to a globally stable and feasible solution as long as $\rho < 1$ (Fig. 3, white area). The higher the competition strength ρ , the larger the size of this restricted area (Fig. 3) (19, 26). Therefore, a relevant question here is not only whether or not the system is structurally stable, but how large is the domain in the parameter space leading to the stable coexistence of species.

To address the above question, we recast the mathematical definition of structural stability to that in which a system is more structurally stable, the greater the area of parameter values leading to both a dynamically stable and feasible equilibrium (27–29). This means that a highly structurally stable ecological system is more likely to be stable and feasible by handling a wider range of conditions before the first species becomes extinct. Previous studies have used this approach in low-dimensional ecological systems (3, 19). Yet because of its complexity, almost no study has fully developed this rigorous analysis for a system with an arbitrary number of species. An exception has been the use of structural stability to calculate an upper bound to the number of species that can coexist in a given community (6, 30).

Here, we introduce this extended concept of structural stability into community ecology in order to study the extent to which network architecture—strength and organization of inter-

specific interactions—modulates the range of conditions compatible with the stable coexistence of species. As an empirical application of our framework, we studied the structural stability of mutualistic systems and applied it on a data set of 23 quantitative mutualistic networks (table S1). We surmise that observed network architectures increase the structural stability and in turn the likelihood of species coexistence as a function of the possible set of conditions in an ecological system. We discuss the applicability of our framework to other types of interspecific interactions in complex ecological systems.

Structural stability of mutualistic systems

Mutualistic networks are formed by the mutually beneficial interactions between flowering plants

and their pollinators or seed dispersers (31). These mutualistic networks have been shown to share a nested architectural pattern (32). This nested architecture means that typically, the mutualistic interactions of specialist species are proper subsets of the interactions of more generalist species (32). Although it has been repeatedly shown that this nested architecture may arise from a combination of life history and complementarity constraints among species (32–35), the effect of this nested architecture on community persistence continues to be a matter of strong debate. On the one hand, it has been shown that a nested architecture can facilitate the maintenance of species coexistence (6), exhibit a flexible response to environmental disturbances (7, 8, 36), and maximize total abundance (12). On the other hand, it has also been suggested that this nested

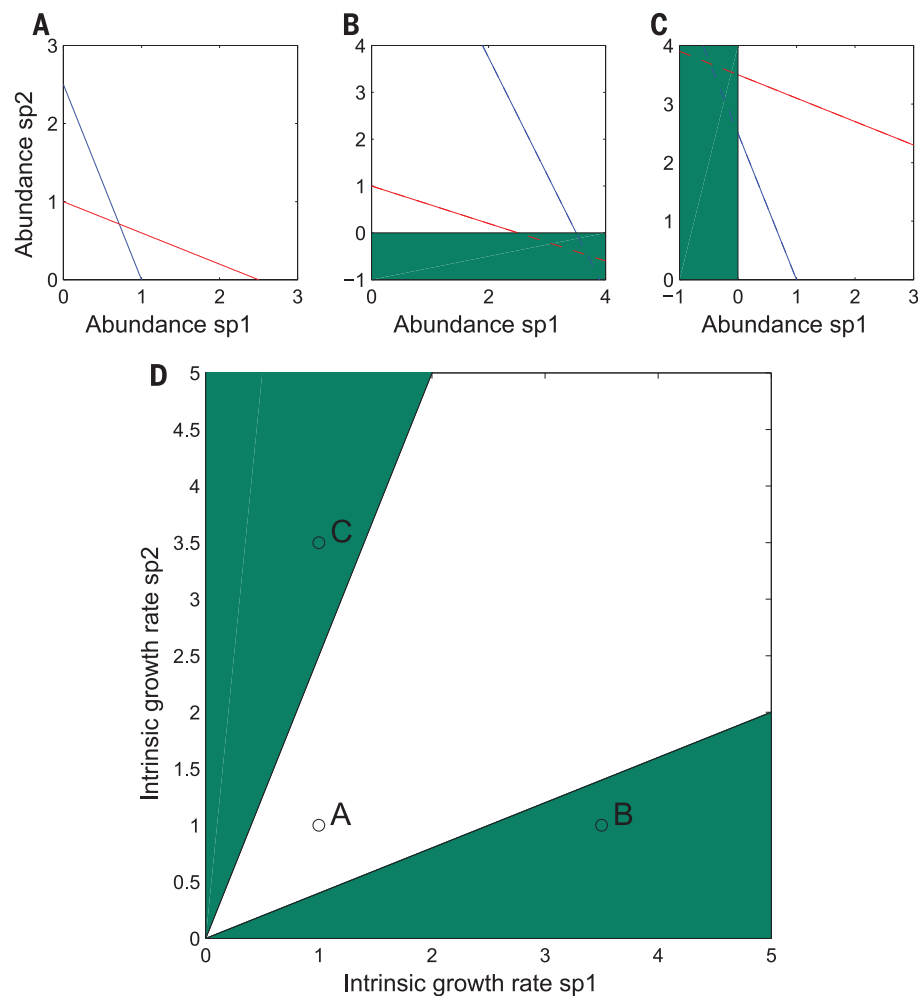


Fig. 1. Stability and feasibility of a two-species competition system. For the same parameters of competition strength (which grant the global stability of any feasible equilibrium), (A to C) represent the two isoclines of the system. Their intersection gives the equilibrium point of the system (3, 19). Scenario (A) leads to a feasible equilibrium (both species have positive abundances at equilibrium), whereas in scenarios (B) and (C), the equilibrium is not feasible (one species has a negative abundance at equilibrium). (D) represents the area of feasibility in the parameter space of intrinsic growth rates, under the condition of global stability. This means that when the intrinsic growth rates of species are chosen within the white area, the equilibrium point is globally stable and feasible. In contrast, when the intrinsic growth rates of species are chosen within the green area, the equilibrium point is not feasible. Points "A," "B," and "C" indicate the parameter values corresponding to (A) to (C), respectively.

architecture can minimize local stability (9), have a negative effect on community persistence (10), and have a low resilience to perturbations (12). Not surprisingly, the majority of these studies have been based on either local stability or numerical simulations with arbitrary parameterizations [but, see (6)].

Model of mutualism

To study the structural stability and explain the apparently contradictory results found in studies

of mutualistic networks, we first need to introduce an appropriate model describing the dynamics between and within plants and animals. We use the same set of differential equations as in (6). We chose these dynamics because they are simple enough to provide analytical insights and yet complex enough to incorporate key elements—such as saturating, functional responses (37, 38) and interspecific competition within a guild (6)—recently adduced as necessary ingredients for a reasonable theoretical exploration of mutualistic

interactions. Specifically, the dynamical model has the following form

$$\begin{cases} \frac{dP_i}{dt} = P_i \left(\alpha_i^{(P)} - \sum_j \beta_{ij}^{(P)} P_j + \frac{\sum_j \gamma_{ij}^{(P)} A_j}{1 + h \sum_j \gamma_{ij}^{(P)} A_j} \right) \\ \frac{dA_i}{dt} = A_i \left(\alpha_i^{(A)} - \sum_j \beta_{ij}^{(A)} A_j + \frac{\sum_j \gamma_{ij}^{(A)} P_j}{1 + h \sum_j \gamma_{ij}^{(A)} P_j} \right) \end{cases} \quad (2)$$

where the variables P_i and A_i denote the abundance of plant and animal species i , respectively.

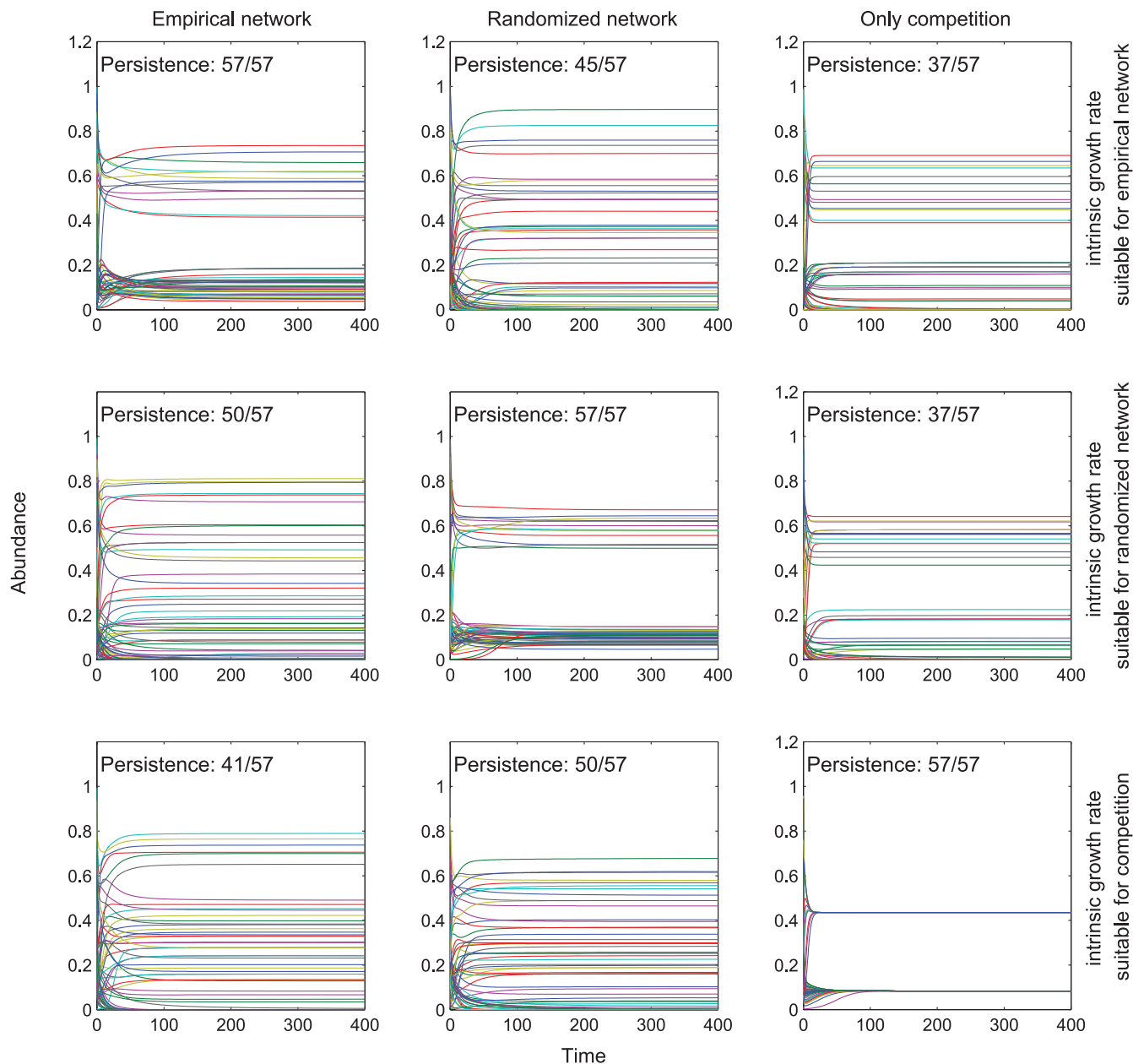


Fig. 2. Numerical analysis of species persistence as a function of model parameterization. This figure shows the simulated dynamics of species abundance and the fraction of surviving species (positive abundance at the end of the simulation) using the mutualistic model of (6). Simulations are performed by using an empirical network located in Hickling, Norfolk, UK (table S1), a randomized version of this network using the probabilistic model

of (32), and the network without mutualism (only competition). Each row corresponds to a different set of growth rate values. It is always possible to choose the intrinsic growth rates so that all species are persistent in each of the three scenarios, and at the same time, the community persistence defined as the fraction of surviving species is lower in the alternative scenarios.

The parameters of this mutualistic system correspond to the values describing intrinsic growth rates (α_i), intra-guild competition (β_{ij}), the benefit received via mutualistic interactions (γ_{ij}), and the saturating constant of the beneficial effect of mutualism (h), commonly known as the handling time. Because our main focus is on mutualistic interactions, we keep as simple as possible the competitive interactions for the sake of analytical tractability. In the absence of empirical information about interspecific competition, we use a mean field approximation for the competition parameters (6), where we set $\beta_{ii}^{(P)} = \beta_{ii}^{(A)} = 1$ and $\beta_{ij}^{(P)} = \beta_{ij}^{(A)} = \rho < 1$ ($i \neq j$).

Following (39), the mutualistic benefit can be further disentangled by $\gamma_{ij} = (\gamma_0 y_{ij})/(k_i^0)$, where $y_{ij} = 1$ if species i and j interact and zero otherwise, k_i is the number of interactions of species i , γ_0 represents the level of mutualistic strength, and δ corresponds to the mutualistic trade-off. The mutualistic strength is the per capita effect of a certain species on the per capita growth rate of their mutualistic partners. The mutualistic trade-off modulates the extent to which a species that interacts with few other species does it strongly, whereas a species that interacts with many partners does it weakly. This trade-off has been justified on empirical grounds (40, 41). The degree to which interspecific interactions y_{ij} are organized into a nested way can be quantified by the value of nestedness N introduced in (42).

We are interested in quantifying the extent to which network architecture (the combination of mutualistic strength, mutualistic trade-off, and nestedness) modulates the set of conditions compatible with the stable coexistence of all species—the structural stability. In the next sections, we explain how this problem can be split into two parts. First, we explain how the stability conditions can be disentangled from the feasibility conditions as it has already been shown for the two-species competition system. Specifically, we show that below a critical level of mutualistic strength ($\gamma_0 < \gamma_0^*$), any feasible equilibrium point is granted to be globally stable. Second, we explain how network architecture modulates the domain in the parameter space of intrinsic growth rates, leading to a feasible equilibrium under the constraints of being globally stable (given by the level of mutualistic strength).

Stability condition

We investigated the conditions in our dynamical system that any feasible equilibrium point needs to satisfy to be globally stable. To derive these conditions, we started by studying the linear Lotka-Volterra approximation ($h = 0$) of the dynamical model (Eq. 2). In this linear approximation, the model reads

$$\begin{bmatrix} \frac{dP}{dt} \\ \frac{dA}{dt} \end{bmatrix} = \text{diag}\left(\begin{bmatrix} P \\ A \end{bmatrix}\right) \times \left(\begin{bmatrix} \alpha^{(P)} \\ \alpha^{(A)} \end{bmatrix} - \underbrace{\begin{bmatrix} \beta^{(P)} & -\gamma^{(P)} \\ -\gamma^{(A)} & \beta^{(A)} \end{bmatrix}}_{:=B} \begin{bmatrix} P \\ A \end{bmatrix} \right) \quad (3)$$

where the matrix B is a two-by-two block matrix embedding all the interaction strengths.

Conveniently, the global stability of a feasible equilibrium point in this linear Lotka-Volterra model has already been studied (14–17, 43). Particularly relevant to this work is that an interaction matrix that is Lyapunov-diagonally stable grants the global stability of any potential feasible equilibrium (14–18).

Although it is mathematically difficult to verify the condition for Lyapunov diagonal stability, it is known that for some classes of matrices, Lyapunov stability and Lyapunov diagonal stability are equivalent conditions (44). Symmetric matrices and Z-matrices (matrices whose off-diagonal elements are nonpositive) belong to those classes of equivalent matrices. Our interaction strength matrix B is either symmetric when the mutualistic trade-off is zero ($\delta = 0$) or is a Z-matrix when the interspecific competition is zero ($\rho = 0$). This means that as long as the real parts of all eigenvalues of B are positive (18), any feasible equilibrium point is globally stable. For instance, in the case of $\rho < 1$ and $\gamma_0 =$

0, the interaction matrix B is symmetric and Lyapunov-diagonally stable because its eigenvalues are $1 - \rho$, $(S_A - 1)\rho + 1$, and $(S_P - 1)\rho + 1$.

For $\rho > 0$ and $\delta > 0$, there are no analytical results yet demonstrating that Lyapunov diagonal stability is equivalent to Lyapunov stability. However, after intensive numerical simulations we conjecture that the two main consequences of Lyapunov diagonal stability hold (45). Specifically, we state the following conjectures: Conjecture 1: If B is Lyapunov-stable, then B is D-stable. Conjecture 2: If B is Lyapunov-stable, then any feasible equilibrium is globally stable.

We found that for any given mutualistic trade-off and interspecific competition, the higher the level of mutualistic strength, the smaller the maximum real part of the eigenvalues of B (45). This means that there is a critical value of mutualistic strength (γ_0^*) so that above this level, the matrix B is not any more Lyapunov-stable. To compute γ_0^* , we need only to find the critical value of γ_0 at which the real part of one of the eigenvalues of the interaction-strength matrix reaches zero (45). This implies that at least below this critical value

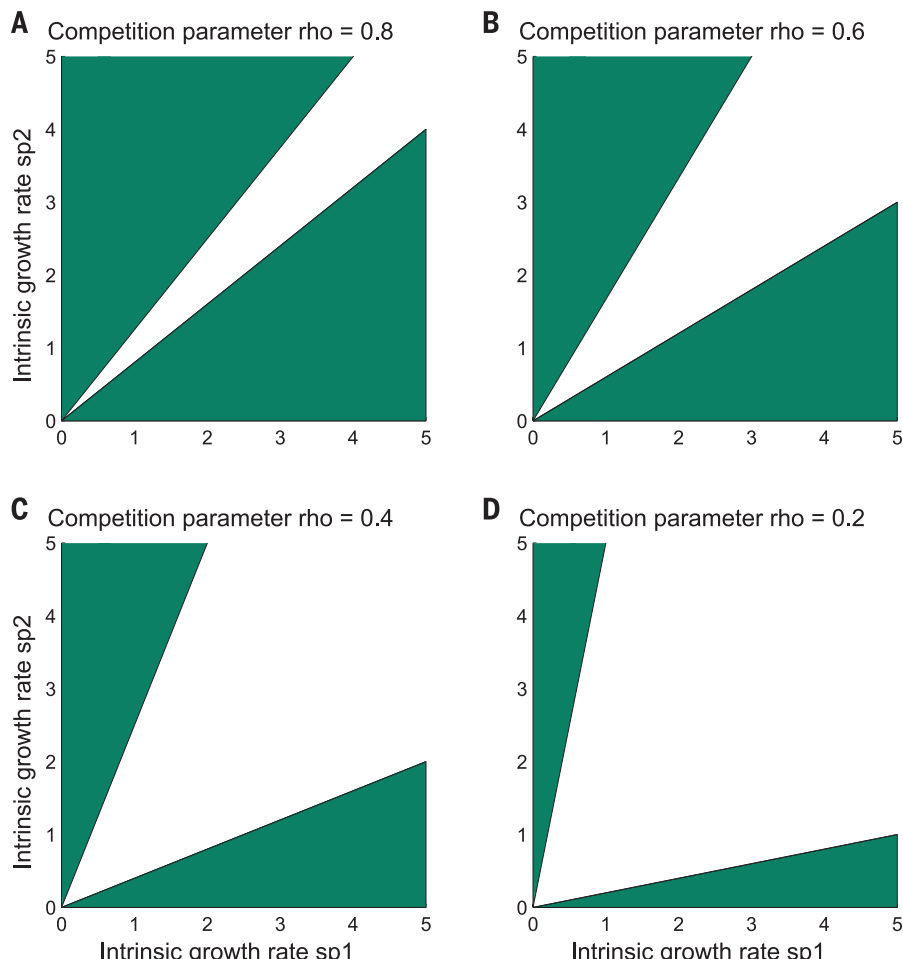


Fig. 3. Structural stability in a two-species competition system. The figure shows how the range of intrinsic growth rates leading to the stable coexistence of the two species (white region) changes as a function of the competition strength. (A to D) Decreasing interspecific competition increases the area of feasibility, and in turn, the structural stability of the system. Here, $\beta_{11} = \beta_{22} = 1$, and $\beta_{12} = \beta_{21} = \rho$. Our goal is extending this analysis to realistic networks of species interactions.

$\gamma_0 < \gamma_0^*$, any feasible equilibrium is granted to be locally and globally stable according to conjectures 1 and 2, respectively. We can also grant the global stability of matrix B by the condition of being positive definite, which is even stronger than Lyapunov diagonal stability (44). However, this condition imposes stronger constraints on the critical value of mutualistic strength than does Lyapunov stability (39).

Last, we studied the stability conditions for the nonlinear Lotka-Volterra system (Eq. 2). Although the theory has been developed for the linear Lotka-Volterra system, it can be extended to the nonlinear dynamical system. To grant the stability of any feasible equilibrium ($P_i > 0$ and $A_i > 0$ for all i) in the nonlinear system, we need to show that the above stability conditions hold on the following two-by-two block matrix (44, 43)

$$B_{nl} := \begin{bmatrix} \beta_{ij}^{(P)} & -\frac{\gamma_{ij}^{(P)}}{1 + h \sum_k \gamma_{ik}^{(P)} A_k} \\ -\frac{\gamma_{ij}^{(A)}}{1 + h \sum_k \gamma_{ik}^{(A)} P_k} & \beta_{ij}^{(A)} \end{bmatrix} \quad (4)$$

B_{nl} differs from B only in the off-diagonal block with a decreased mutualistic strength. This implies that the critical value of mutualistic strength for the nonlinear Lotka-Volterra system is larger than or equal to the critical value for the linear system (45). Therefore, the critical value γ_0^* derived from the linear Lotka-Volterra system (from the matrix B) is already a sufficient condition to grant the global stability of any feasible equilibrium in the nonlinear case. However, this does not imply that above this critical value of mutualistic

strength, a feasible equilibrium is unstable. In fact, when the mutualistic-interaction terms are saturated ($h > 0$), it is possible to have feasible and locally stable equilibria for any level of mutualistic strength (39, 45).

Feasibility condition

We highlight that for any interaction strength matrix B , whether it is stable or not, it is always possible to find a set of intrinsic growth rates so that the system is feasible (Fig. 2). To find this set of values, we need only to choose a feasible equilibrium point so that the abundance of all species is greater than zero ($A_i^* > 0$ and $P_j^* > 0$) and find the vector of intrinsic growth rates so that the right side of Eq. 2 is

$$\text{equal to zero: } \alpha_i^{(P)} = \sum_j \beta_{ij}^{(P)} P_j^* - \frac{\sum_j \gamma_{ij}^{(P)} A_j^*}{1 + h \sum_k \gamma_{ik}^{(P)} A_k} \text{ and } \alpha_i^{(A)} = \sum_j \beta_{ij}^{(A)} A_j^* - \frac{\sum_j \gamma_{ij}^{(A)} P_j^*}{1 + h \sum_k \gamma_{ik}^{(A)} P_k^*}. \text{ This recom-}$$

firms that the stability and feasibility conditions are different and that they need to be rigorously verified when studying the stable coexistence of species (3, 16, 17, 19). This also highlights that the relevant question is not whether we can find a feasible equilibrium point, but how large is the domain of intrinsic growth rates leading to a feasible and stable equilibrium point. We call this domain the feasibility domain.

Because the parameter space of intrinsic growth rates is substantially large (\mathbb{R}^S , where S is the total number of species), an exhaustive numerical search of the feasibility domain is impossible. However, we can analytically estimate the center of this domain with what we call the structural vector of intrinsic growth rates. For example, in

the two-species competition system of Fig. 4A the structural vector is the vector (in red), which is in the center of the domain leading to feasibility of the equilibrium point (white region). Any vector of intrinsic growth rates collinear to the structural vector guarantees the feasibility of the equilibrium point—that is, guarantees species coexistence. Because the structural vector is the center of the feasibility domain, then it is also the vector that can tolerate the strongest deviation before leaving the feasibility domain—that is, before having at least one species going extinct.

In mutualistic systems, we need to find one structural vector for animals and another for plants. These structural vectors are the set of intrinsic growth rates that allow the strongest perturbations before leaving the feasibility domain. To find these structural vectors, we had to transform the interaction-strength matrix B to an effective competition framework (45). This results in an effective competition matrix for plants and a different one for animals (6), in which these matrices represent respectively the apparent competition among plants and among animals, once taking into account the indirect effect via their mutualistic partners. With a nonzero mutualistic trade-off ($\delta > 0$), the effective competition matrices are nonsymmetric, and in order to find the structural vectors, we have to use the singular decomposition approach—a generalization of the eigenvalue decomposition. This results in a left and a right structural vector for plants and for animals in the effective competition framework. Last, we need to move back from the effective competition framework in order to obtain a left and right vector for plants ($\alpha_L^{(P)}$ and $\alpha_R^{(P)}$) and animals ($\alpha_L^{(A)}$ and $\alpha_R^{(A)}$) in the observed mutualistic framework. The full derivation is provided in the supplementary materials (45).

Once we locate the center of the feasibility domain with the structural vectors, we can approximate the boundaries of this domain by quantifying the amount of variation from the structural vectors allowed by the system before having any of the species going extinct—that is, before losing the feasibility of the system. To quantify this amount, we introduced proportional random perturbations to the structural vectors, numerically generated the new equilibrium points (21), and measured the angle or the deviation between the structural vectors and the perturbed vectors (a graphical example is provided in Fig. 4A). The deviation from the structural vectors is quantified, for the plants, by $\eta_P(\alpha^{(P)}) = [1 - \cos(\theta_L^{(P)}) \cos(\theta_R^{(P)})]/[\cos(\theta_L^{(P)}) \cos(\theta_R^{(P)})]$, where $\theta_L^{(P)}$ and $\theta_R^{(P)}$ are, respectively, the angles between $\alpha^{(P)}$ and $\alpha_L^{(P)}$ and between $\alpha^{(P)}$ and $\alpha_R^{(P)}$. The parameter $\alpha^{(P)}$ is any perturbed vector of intrinsic growth rates of plants. The deviation from the structural vector of animals is computed similarly.

As shown in Fig. 4B, the greater the deviation of the perturbed intrinsic growth rates from the structural vectors, the lower is the fraction of surviving species. This confirms that there is a restricted domain of intrinsic growth rates centered on the structural vectors compatible with the stable coexistence of species. The greater the tolerated deviation from the structural vectors

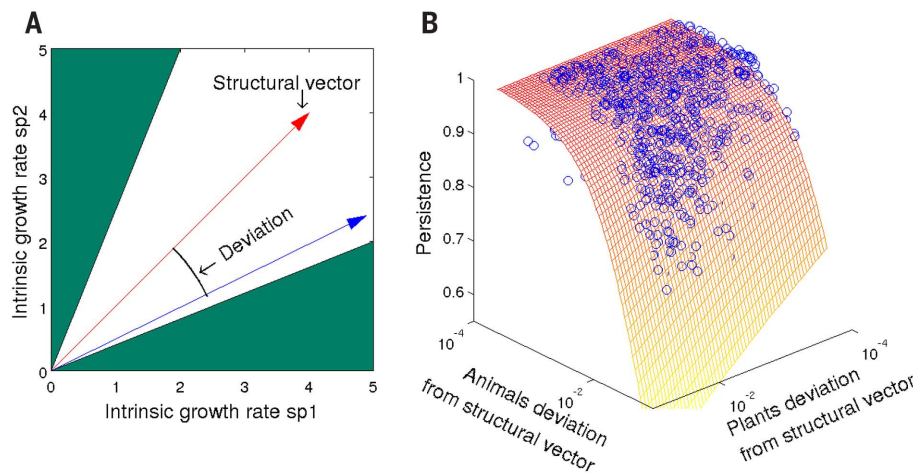


Fig. 4. Deviation from the structural vector and community persistence. (A) The structural vector of intrinsic growth rates (in red) for the two-species competition system of Fig. 1. The structural vector is the vector in the center of the domain leading to the feasibility of the equilibrium point (white region) and thus can tolerate the largest deviation before any of the species go extinct. The deviation between the structural vector and any other vector (blue) is quantified by the angle between them. (B) The effect of the deviation from the structural vector on intrinsic growth rates on community persistence defined as the fraction of model-generated surviving species. The example corresponds to an observed network located in North Carolina, USA (table S1), with a mutualistic trade-off $\delta = 0.5$ and a maximum level of mutualistic strength $\gamma_0 = 0.2402$. Blue symbols represent the community persistence, and the surface represents a fit of a logistic regression ($R^2 = 0.88$).

within which all species coexist, the greater the feasibility domain, and in turn the greater the structural stability of the system.

Network architecture and structural stability

To investigate the extent to which network architecture modulates the structural stability of mutualistic systems, we explored the combination of alternative network architectures (combinations of nestedness, mutualistic strength, and mutualistic trade-off) and their corresponding feasibility domains.

To explore these combinations, for each observed mutualistic network (table S1) we obtained 250 different model-generated nested architectures by using an exhaustive resampling model (46) that preserves the number of species and the expected number of interactions (45). Theoretically, nestedness ranges from 0 to 1 (42). However, if one imposes architectural constraints such as preserving the number of species and interactions, the effective range of nestedness that the network can exhibit may be smaller (45). Additionally, each individual model-generated nested architecture is combined with different levels of mutualistic trade-off δ and mutualistic strength γ_0 . For the mutualistic trade-off, we explored values $\delta \in [0, \dots, 1.5]$ with steps of 0.05 that allow us to explore sublinear, linear, and superlinear trade-offs. The case $\delta = 0$ is equivalent to the soft mean field approximation studied in (6). For each combination of network of interactions and mutualistic trade-off, there is a specific critical value γ_0^* in the level of mutualism strength γ_0 up to which any feasible equilibrium is globally stable. This critical value γ_0^* is dependent on the mutualistic trade-off and nestedness. However, the mean mutualistic strength $\bar{\gamma} = \langle \gamma_{ij} \rangle$ shows no pattern as a function of mutualistic trade-off and nestedness (45). Therefore, we explored values of $\gamma_0 \in [0, \dots, \gamma_0^*]$ with steps of 0.05 and calculated the new generated mean mutualistic strengths. This produced a total of 250×589 different network architectures (nestedness, mutualistic trade-off, and mean mutualistic strength) for each observed mutualistic network.

We quantified how the structural stability (feasibility domain) is modulated by these alternative network architectures in the following way. First, we computed the structural vectors of intrinsic growth rates that grant the existence of a feasible equilibrium of each alternative network architecture. Second, we introduced proportional random perturbations to the structural vectors of intrinsic growth rates and measured the angle or deviation ($\eta_{(A)}, \eta_{(P)}$) between the structural vectors and the perturbed vectors. Third, we simulated species abundance using the mutualistic model of (6) and the perturbed growth rates as intrinsic growth rate parameter values (21). These deviations lead to parameter domains from all to a few species surviving (Fig. 4).

Last, we quantified the extent to which network architecture modulates structural stability by looking at the association of community persistence with network architecture parameters,

once taking into account the effect of intrinsic growth rates. Specifically, we studied this association using the partial fitted values from a binomial regression (47) of the fraction of surviving species on nestedness (N), mean mutualistic strength ($\bar{\gamma}$), and mutualistic trade-off (δ), while controlling for the deviations from the structural vectors of intrinsic growth rates ($\eta_{(A)}, \eta_{(P)}$). The full description of this binomial regression and the calculation of partial fitted values are provided in (48). These partial fitted values are the contribution of network architecture to the logit of the probability of species persistence, and in turn, these values are positively proportional to the size of the feasibility domain.

Results

We analyzed each observed mutualistic network independently because network architecture is

constrained to the properties of each mutualistic system (11). For a given pollination system located in the KwaZulu-Natal region of South Africa, the extent to which its network architecture modulates structural stability is shown in Fig. 5. Specifically, the partial fitted values are plotted as a function of network architecture. As shown in Fig. 5A, not all architectural combinations have the same structural stability. In particular, the architectures that maximize structural stability (reddish/darker regions) correspond to the following properties: (i) a maximal level of nestedness, (ii) a small (sublinear) mutualistic trade-off, and (iii) a high level of mutualistic strength within the constraint of any feasible solution being globally stable (49).

A similar pattern is present in all 23 observed mutualistic networks (45). For instance, using three different levels of interspecific competition

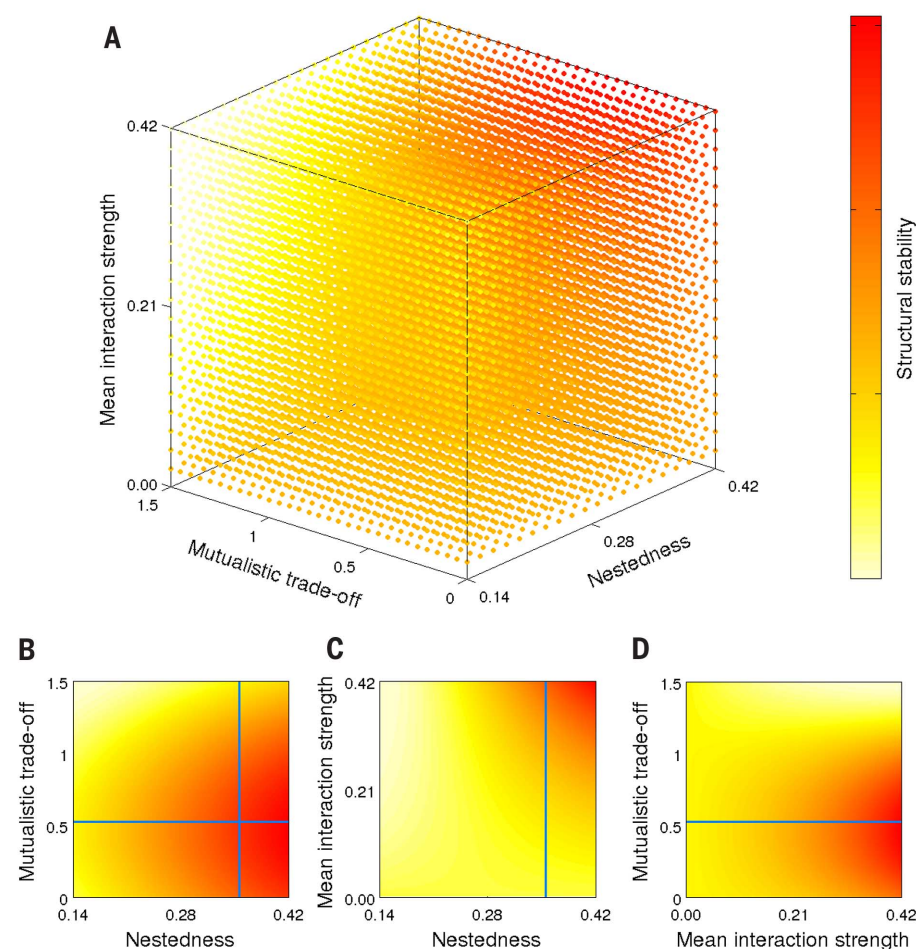


Fig. 5. Structural stability in complex mutualistic systems. For an observed mutualistic system with 9 plants, 56 animals, and 103 mutualistic interactions located in the grassland asclepiads in South Africa (table S1) (58), (A) corresponds to the effect—colored by partial fitted residuals—of the combination of different architectural values (nestedness, mean mutualistic strength, and mutualistic trade-off) on the domain of structural stability. The reddish/darker the color, the larger the parameter space that is compatible with the stable coexistence of all species, and in turn the larger the domain of structural stability. (B), (C), and (D) correspond to different slices of (A). Slice (B) corresponds to a mean mutualistic strength of 0.21, slice (C) corresponds to the observed mutualistic trade-off, and slice (D) corresponds to the observed nestedness. Solid lines correspond to the observed values of nestedness and mutualistic trade-offs.

($\rho = 0.2, 0.4, 0.6$), we always find that structural stability is positively associated with nestedness and mutualistic strength (45). Similarly, structural stability is always associated with the mutualistic trade-off by a quadratic function, leading quite often to an optimal value for maximizing structural stability (45). These findings reveal that under the given characterization of interspecific competition, there is a general pattern of

network architecture that increases the structural stability of mutualistic systems.

Yet, one question remains to be answered: Is the network architecture that we observe in nature close to the maximum feasibility domain of parameter space under which species coexist? To answer this question, we compared the observed network architecture with theoretical predictions. To extract the observed network

architecture, we computed the observed nestedness from the observed binary interaction matrices (table S1) following (42). The observed mutualistic trade-off δ is estimated from the observed number of visits of pollinators or fruits consumed by seed-dispersers to flowering plants (41, 50, 51). The full details on how to compute the observed trade-off is provided in (52). Because there is no empirical data on the relationship between competition and mutualistic strength that could allow us to extract the observed mutualistic strength γ_0 , our results on nestedness and mutualistic trade-off are calculated across different levels of mean mutualistic strength.

As shown in Fig. 5, B to D, the observed network (blue solid lines) of the mutualistic system located in the grassland asclepiads of South Africa actually appears to have an architecture close to the one that maximizes the feasibility domain under which species coexist (reddish/darker region). To formally quantify the degree to which each observed network architecture is maximizing the set of conditions under which species coexist, we compared the net effect of the observed network architecture on structural stability against the maximum possible net effect. The maximum net effect is calculated in three steps.

First, as outlined in the previous section, we computed the partial fitted values of the effect of alternative network architectures on species persistence (48). Second, we extracted the range of nestedness allowed by the network given the number of species and interactions in the system (45). Third, we computed the maximum net effect of network architecture on structural stability by finding the difference between the maximum and minimum partial fitted values within the allowed range of nestedness and mutualistic trade-off between $\delta \in [0, \dots, 1.5]$. All the observed mutualistic trade-offs have values between $\delta \in [0, \dots, 1.5]$. Last, the net effect of the observed network architecture on structural stability corresponds to the difference between the partial fitted values for the observed architecture and the minimum partial fitted values extracted in the third step described above.

Looking across different levels of mean mutualistic strength, in the majority of cases (18 out of 23, $P = 0.004$, binomial test) the observed network architectures induce more than half the value of the maximum net effect on structural stability (Fig. 6, red solid line). These findings reveal that observed network architectures tend to maximize the range of parameter space—structural stability—for species coexistence.

Structural stability of systems with other interaction types

In this section, we explain how our structural stability framework can be applied to other types of interspecific interactions in complex ecological systems. We first explain how structural stability can be applied to competitive interactions. We proceed by discussing how this competitive approach can be used to study trophic interactions in food webs.

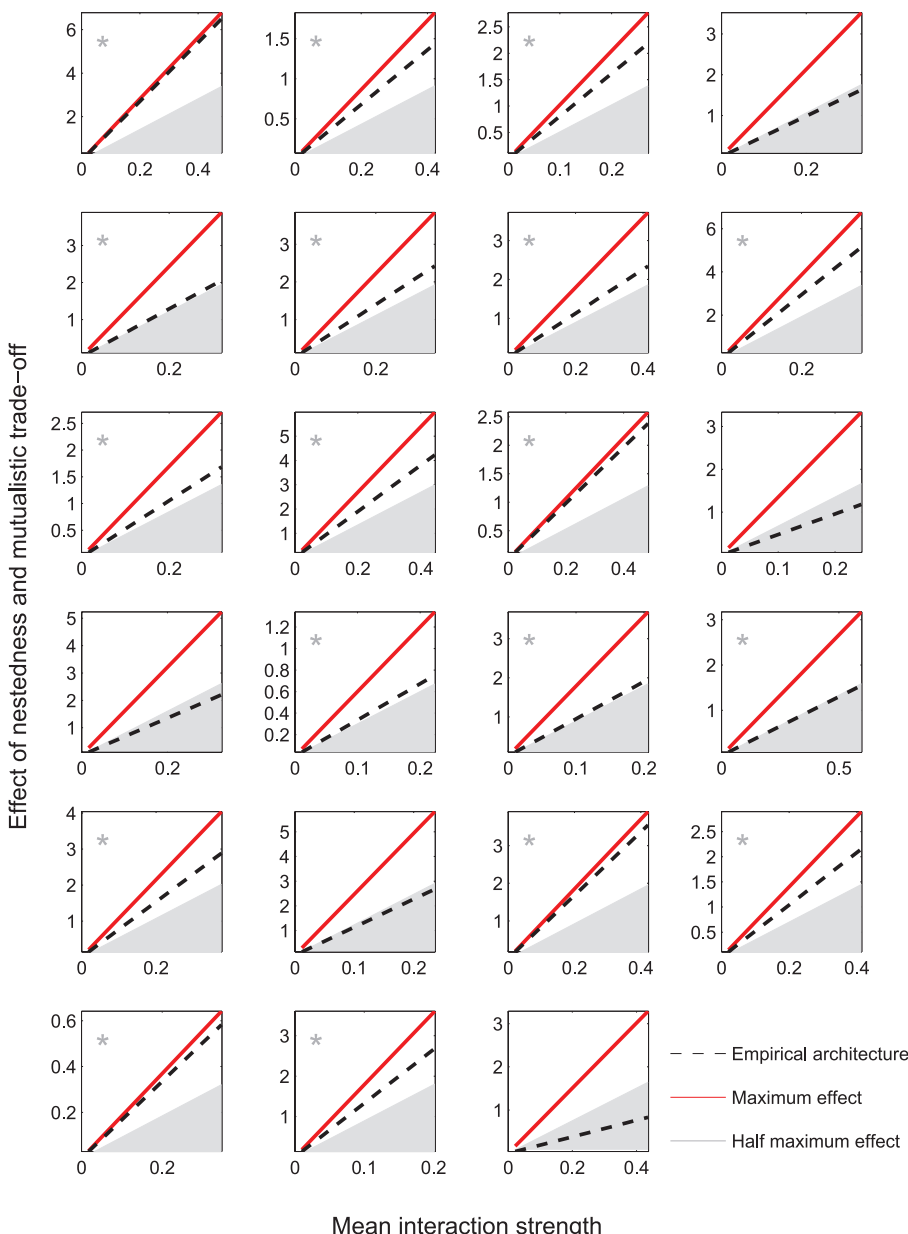


Fig. 6. Net effect of network architecture on structural stability. For each of the 23 observed networks (table S1), we show how close the observed feasibility domain (partial fitted residuals) is as a function of the network architecture to the theoretical maximal feasibility domain. The network architecture is given by the combination of nestedness and mutualistic trade-off (x axis) across different values of mean mutualistic strength (y axis). The solid red and dashed black lines correspond to the maximum net effect and observed net effect, respectively. In 18 out of 23 networks (indicated by asterisks), the observed architecture exhibits more than half the value of the maximum net effect (gray regions). The net effect of each network architecture is system-dependent and cannot be used to compare across networks.

For a competition system with an arbitrary number of species, we can assume a standard set of dynamical equations given by $\frac{dN_i}{dt} = N_i(\alpha_i - \sum_j \beta_{ij}N_j)$, where $\alpha_i > 0$ is the intrinsic growth rate, $\beta_{ij} > 0$ is the competition interaction strength, and N_i is the abundance of species i . Recall that the Lyapunov diagonal stability of the interaction matrix β would imply the global stability of any feasible equilibrium point. However, in nonsymmetric competition matrices Lyapunov stability does not always imply Lyapunov diagonal stability (53). This establishes that we should work with a restricted class of competition matrices such as the ones derived from the niche space of (54). Indeed, it has been demonstrated that this class of competition matrices are Lyapunov diagonally stable and that this stability is independent of the number of species (55). For a competition system with a symmetric interaction-strength matrix, the structural vector is equal to its leading eigenvector. For other appropriate classes of matrices, we can compute the structural vectors in the same way as we did with the effective competition matrices of our mutualistic model and numerically simulate the feasibility domain of the competition system. In general, following this approach we can verify that the lower the average interspecific competition, the higher is the feasibility domain, and in turn, the higher is the structural stability of the competition system.

In the case of predator-prey interactions in food webs, so far there is no analytical work demonstrating the conditions for a Lyapunov-diagonally stable system and how this is linked to its Lyapunov stability. Moreover, the computation of the structural vector of an antagonistic system is not a straightforward task. However, we may have a first insight about how the network architecture of antagonistic systems modulates their structural stability by transforming a two-trophic-level food web into a competition system among predators. Using this transformation, we are able to verify that the higher the compartmentalization of a food web, then the higher is its structural stability. There is no universal rule to study the structural stability of complex ecological systems. Each type of interaction poses their own challenges as a function of their specific population dynamics.

Discussion

We have investigated the extent to which different network architectures of mutualistic systems can provide a wider range of conditions under which species coexist. This research question is completely different from the question of which network architectures are aligned to a fixed set of conditions. Previous numerical analyses based on arbitrary parameterizations were indirectly asking the latter, and previous studies based on local stability were not rigorously verifying the actual coexistence of species. Of course, if there is a good empirical or scientific reason to use a specific parameterization, then we should take advantage of this. However, because the set of conditions present in a community can be constant-

ly changing because of stochasticity, adaptive mechanisms, or global environmental change, we believe that understanding which network architectures can increase the structural stability of a community becomes a relevant question. Indeed, this is a question much more aligned with the challenge of assessing the consequences of global environmental change—by definition, directional and large—than with the alternative framework of linear stability, which focuses on the responses of a steady state to infinitesimally small perturbations.

We advocate structural stability as an integrative approach to provide a general assessment of the implications of network architecture across ecological systems. Our findings show that many of the observed mutualistic network architectures tend to maximize the domain of parameter space under which species coexist. This means that in mutualistic systems, having both a nested network architecture and a small mutualistic trade-off is one of the most favorable structures for community persistence. Our predictions could be tested experimentally by exploring whether communities with an observed network architecture that maximizes structural stability stand higher values of perturbation. Similarly, our results open up new questions, such as what the reported associations between network architecture and structural stability tell us about the evolutionary processes and pressures occurring in ecological systems.

Although the framework of structural stability has not been as dominant in theoretical ecology as has the concept of local stability, it has a long tradition in other fields of research (29). For example, structural stability has been key in evolutionary developmental biology to articulate the view of evolution as the modification of a conserved developmental program (27, 28). Thus, some morphological structures are much more common than others because they are compatible with a wider range of developmental conditions. This provided a more mechanistic understanding of the generation of form and shape through evolution (56) than that provided by a historical, functionalist view. We believe ecology can also benefit from this structuralist view. The analogous question here would assess whether the invariance of network architecture across diverse environmental and biotic conditions is due to the fact that such a network structure is the one increasing the likelihood of species coexistence in an ever-changing world.

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- Any matrix B is called Lyapunov-stable if the real parts of all its eigenvalues are positive, meaning that a feasible equilibrium at which all species have the same abundance is at least locally stable. A matrix B is called D -stable if DB is a Lyapunov-stable matrix for any strictly positive diagonal matrix D . D -stability is a stronger condition than Lyapunov stability in the sense that it grants the local stability of any feasible equilibrium (15). In addition, a matrix B is called Lyapunov diagonally stable if there exists a strictly positive diagonal matrix D so that $DB + B^tD$ is a Lyapunov-stable matrix. This notion of stability is even stronger than D -stability in the sense that it grants not only the local stability of any feasible equilibrium point, but also its global stability (14). A Lyapunov-diagonally stable matrix has all of its principal minors positive. Also, its stable equilibrium (which may be only partially feasible; some species may have an abundance of zero) is specific (57). This means that there is no alternative stable state for a given parameterization of intrinsic growth rates (55). We have chosen the convention of having a minus sign in front of the interaction-strength matrix B . This implies that B has to be positive Lyapunov-stable (the real parts of all eigenvalues have to be strictly positive) so that the equilibrium point of species abundance is locally stable. This same convention applies for D -stability and Lyapunov-diagonal stability.
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48. For a given network, we look at the association of the fraction of surviving species with the deviation from the structural vector of plants $\eta_{(P)}$ and animals $\eta_{(A)}$, nestedness N , mutualistic trade-off δ , and mean level of mutualistic strength $\bar{\gamma} = \langle \gamma_{ij} \rangle$ using the following binomial generalized linear model (47):

$$\text{logit}(\text{probability of surviving}) \sim \log(\eta_{(A)}) + \log(\eta_{(P)}) + \bar{\gamma} + \bar{\gamma}^2 + \bar{\gamma} \cdot N + \bar{\gamma} \cdot N^2 + \bar{\gamma} \cdot \delta + \bar{\gamma} \cdot \delta^2.$$
 Obviously, at a level of mutualism of zero ($\gamma = 0$), nestedness and mutualistic trade-off cannot influence the probability of a species to survive. We have to include an interaction between the mean level of mutualism and the nestedness and mutualistic trade-off. We have also included a quadratic term in order to take into account potential nonlinear effects of nestedness, mutualistic trade-off, and mean level of mutualistic strength. The effect of network architecture is confirmed by the significant likelihood ratio between the full model and a null model without such a network effect ($P < 0.001$) for all the observed empirical networks. The effect of network architecture on structural stability can be quantified by the partial fitted values defined as follows: partial fitted values = $\hat{\beta}_1\bar{\gamma} + \hat{\beta}_2\bar{\gamma}^2 + \hat{\beta}_3\bar{\gamma} \cdot N + \hat{\beta}_4\bar{\gamma} \cdot N^2 + \hat{\beta}_5\bar{\gamma} \cdot \delta + \hat{\beta}_6\bar{\gamma} \cdot \delta^2$, where $\hat{\beta}_1 \dots \hat{\beta}_6$ are the fitted parameters corresponding to the terms $\bar{\gamma}$ to $\bar{\gamma} \cdot \delta^2$, respectively.
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52. To estimate the mutualistic trade-off, δ , from an empirical point of view, we proceed as follows. First, there is available data on the frequency of interactions. Thus, q_{ij} is the observed number of visits of animal species j on plant species i . This quantity has been proven to be the best surrogate of per capita effects of one species on another (γ_{ij}^P and γ_{ij}^A) (41). Second, the networks provide information on the number of species one species interacts with (its degree or generalization level; k_i^A and k_i^P). Following (41, 50, 51), the generalization level of a species has been found to be proportional to its abundance at equilibrium. Thus, the division of the total number of visits by the product of the degree of plants and animals can be assumed to be proportional to the interaction strengths. Mathematically, we obtain two equations, one for the effect of the animals on the plants and vice versa: $(q_{ij})/(k_i^P k_i^A) \propto \gamma_{ij}^P$ and $(q_{ij})/(k_i^A k_i^P) \propto \gamma_{ij}^A$. Introducing the explicit dependence between interaction strength and trade-off—for example, $\gamma_{ij}^P = \gamma_0/(k_i^P)^{\delta}$ and $(q_{ij})/(k_i^P k_i^A) \propto (\gamma_0)/[(k_i^P)^{\delta}]$ and $(q_{ij})/(k_i^A k_i^P) \propto (\gamma_0)/[(k_i^A)^{\delta}]$. In order to estimate the value of δ , we can just take the logarithm on both sides of the previous equations for the data excluding the zeroes. Then, δ is simply given by the slope of the following linear regressions:
- $$\log\left(\frac{q_{ij}}{k_i^P k_i^A}\right) = a^P - \delta \log\left(k_i^P\right) \quad \text{and} \quad \log\left(\frac{q_{ij}}{k_i^A k_i^P}\right) = a^A - \delta \log\left(k_i^A\right),$$
- where a^P and a^A are the intercepts for plants and animals, respectively. These two regressions are performed simultaneously by lumping together the data set. The intercept for the effect of animals on plants (a^P) may not be the same as the intercept for the effect of the plants on the animals (a^A).
53. The competition matrix given by $\beta = \begin{bmatrix} 3 & 3.1 & 1 \\ 5 & 5 & 6 \\ 4 & 1 & 3 \end{bmatrix}$ is Lyapunov-stable but not D-stable. Because Lyapunov diagonal stability implies D-stability, this matrix is also not Lyapunov-diagonally stable.
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SUPPLEMENTARY MATERIALS

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Figs. S1 to S14
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Supplementary Material for

On the structural stability of mutualistic systems

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S1 Local and global stability of a feasible equilibrium point

Ref. 14 introduced a Lyapunov function to grant the global stability of any feasible point under some constraints on interaction strengths. In the linear Lotka-Volterra version ($h = 0$), this theory works as follows. Let B be the interaction-strength matrix, i.e.,

$$B = \begin{bmatrix} \beta^{(P)} & -\gamma^{(P)} \\ -\gamma^{(A)} & \beta^{(A)} \end{bmatrix}.$$

Assuming that we can find a strictly positive diagonal matrix D , such that $DB + B^T D \succ 0$ (59), a Lyapunov function can be constructed to prove the global stability of any feasible equilibrium. In this case, the matrix B is called Lyapunov-diagonally stable. Practically, it is a complex task to find a suitable matrix D . A very strict condition would be to impose that the matrix B itself has to be positive definite.

On the other extreme, we can simply look at the local stability of a feasible equilibrium by looking at the Jacobian matrix. Assuming that \tilde{P} and \tilde{A} is a feasible equilibrium, the Jacobian is then given by

$$J = - \begin{bmatrix} \tilde{P}_1 & 0 & \cdots & \cdots & \cdots & 0 \\ 0 & \ddots & \ddots & & & \vdots \\ \vdots & \ddots & \tilde{P}_{S(P)} & \ddots & & \vdots \\ \vdots & & \ddots & \tilde{A}_1 & \ddots & \vdots \\ \vdots & & & \ddots & \ddots & 0 \\ 0 & \cdots & \cdots & \cdots & 0 & \tilde{A}_{S(A)} \end{bmatrix} \begin{bmatrix} \beta^{(P)} & -\gamma^{(P)} \\ -\gamma^{(A)} & \beta^{(A)} \end{bmatrix}.$$

Now, if the real parts of all eigenvalues are strictly negative, the feasible equilibrium is locally stable. The problem with this approach is that we have to evaluate the Jacobian and its eigenvalues for every possible feasible point. Fortunately, the concept of D -stability solves this issue (15,60,61). A matrix M is called D -stable if for every strictly positive diagonal matrix D , the real part of the eigenvalues of DM are strictly positive. Then, if the interaction-strength matrix B were to be D -stable, any feasible equilibrium is granted

to be at least locally stable. Note that Lyapunov-diagonal stability implies D -stability.

Importantly, all these different conditions on stability are imposing constraints on the interaction strengths. Assuming the strength of the competition matrices $\beta^{(P)}$ and $\beta^{(A)}$ are set such that without mutualism any feasible equilibrium is stable, then these different notions of stability constrain the level of mutualistic strength that the system can handle before losing the stability of a feasible equilibrium. Intuitively, in the linear Lotka-Volterra case, the mutualistic interactions have to be “weaker” than competitive interactions to have the stability of a feasible equilibrium.

S2 Numerical test of the two conjectures for the global stability of feasible equilibrium

We provide details on the numerical simulations supporting our two conjectures related to the stability of a feasible equilibrium. Note that for a level of mutualistic strength of zero, i.e., $\gamma_0 = 0$, any feasible equilibrium point is automatically globally stable. Indeed, the competition matrices are chosen positive definite, thus according to Ref. 14, any feasible equilibrium point is globally stable.

To grant the global stability of any feasible equilibrium when $\gamma_0 > 0$, one can use the Lyapunov function developed in Ref. 14. However, in practice, it is a very difficult task to find the positive diagonal matrix D such that $DB + B^t D \succ 0$, and even more difficult to prove it for a full family of matrices. D -stability is a more relaxed condition (15,60), but as before, it is very difficult to prove that a given matrix is D -stable. However, we find that a very general result can be postulated. First, we need to formulate two conjectures:

Conjecture 1: if the interaction strength matrix M is Lyapunov stable, then M is D -stable.

Conjecture 2: if the interaction strength matrix M is Lyapunov stable, then any feasible equilibrium is globally stable.

Recall that for a given network, mutualistic trade-off, and interspecific competition strength, a matrix B is Lyapunov stable when the level of mutualistic strength is below a given critical value $\gamma_0 < \gamma_0^r$. However, there exists a stronger constraint on B within which the two conjectures can be proven. If we impose B to be positive definite, i.e., $B+B^t$ to be Lyapunov stable, then it has already been proven that B is D -stable (15,60). This proves our Conjecture 1. Moreover, following Ref. 14, there exists a Lyapunov function for any feasible equilibrium point. This proves our Conjecture 2. As for Lyapunov stability, the matrix B is positive definite if the level of mutualistic strength is below a critical value $\gamma_0 < \gamma_0^s$, which is stronger than the the critical value for Lyapunov stability ($\gamma^s < \gamma_0^r$). Thus, our two conjectures are partially proven and need to be numerically check only over

a restricted range of mutualistic strength (from γ_0^s to γ_0^r).

For each of the observed networks, we explore the mutualistic trade-off of $\delta = 0, 0.1, 0.2, \dots, 1.5$, interspecific competition of $\rho = 0.2$, and 10 equidistant steps in the level of mutualistic strength between the two critical values γ_0^s and γ_0^r with the following numerical simulations. For conjecture 1, at each step we generated 1000 samples of strictly positive diagonal matrices D and tested whether DB is still Lyapunov stable. During this test, we find no counter-example, thus our conjecture 1 seems to hold.

For conjecture 2, in each step from above, we chose 1000 different feasible equilibria and 1000 different initial points for the numerical integrator, and tested whether the numerical simulations converge to the chosen feasible equilibria. We find the numerical simulations always converged to the same chosen feasible equilibria, indicating that our conjecture 2 also holds.

If it would be analytically proven that the interaction-strength matrix B is Lyapunov-diagonally stable when it is only Lyapunov stable, our two conjectures would become theorems. Indeed, in this case we would have equivalence among Lyapunov stability, D -stability, and Lyapunov-diagonal stability. Additionally, Lyapunov-diagonal stability implies global stability for any feasible equilibrium point. Mathematically, our interaction-strength matrix B can be seen as a two-by-two block Z -matrix, where the diagonal block contains the positive definite mean-field competition matrices and the off-diagonal blocks are non-positive matrices. The equivalence of the three types of stability is known for Z -matrices (40,60,61), but remains an open question for block Z -matrices.

Finally, we run numerical simulations to show that the stability conditions derived within the linear Lotka-Volterra model imply also stability for our nonlinear dynamical model. What we have to show is that if B is Lyapunov stable, then B_{nl} is also Lyapunov stable for any positive plant and animal abundance. As for the two conjectures above, we explored the mutualistic trade-off of $\delta = 0, 0.1, 0.2, \dots, 1.5$, interspecific competition of $\rho = 0.2$, and 10 equidistant steps in the level of mutualistic strength between zero and γ_0^r .

For each step we generate twice 1000 diagonal matrices where the elements are sampled from a uniform distribution between 0 and 1. Then, we multiply the two off-diagonal blocks of the matrix B by two of these random matrices. This procedure mimics the fact that in the nonlinear model the mutualistic interactions have a dominator greater than one. We find that all the generated matrices are Lyapunov stable. This indicates that the stability conditions derived in the linear version hold for the nonlinear version too.

S3 Critical value of mutualistic strength

The main consequence of our two conjectures for stability is that to determine the stability of a feasible equilibrium, it is enough to look at the eigenvalues of the interaction-strength matrix B . Thus, for computing the critical value in the level of mutualistic strength γ_0^r , we just need to find the critical value of γ_0 at which the real part of one of the eigenvalues reaches zero. Importantly, as shown in Fig. S1, we find that for any given mutualistic trade-off and interspecific competition, the higher the level of mutualistic strength, the smaller the real part of the eigenvalues of B .

Additionally, we find no particular pattern in how the critical level of mutualistic strength varies with the specific model parameterization. In Figures S2 and S3, we show how mutualistic trade-off δ and nestedness influence the critical value of mutualistic strength γ_0^r . For this, we randomize the observed networks and compute the slope of the linear regression between this level of mutualistic strength and nestedness. While in the majority of cases the slopes increase along with the mutualistic trade-off, we find that these slopes can be positive as well as negative, meaning that there is no general pattern.

S4 Stability and feasibility for strong mutualism

Our study of mutualistic networks is still limited to mutualistic interactions that are weaker than our critical value of mutualistic strength ($\gamma_0 < \gamma_0^r$). Recall that below this critical value, any feasible point is granted to be globally stable. Mathematically, we can disentangle the feasibility condition from the stability one, like in the two-species competition system. Here we explain how our dynamical system reacts to a mutualism that is stronger than our critical value of mutualistic strength—that corresponds to the strong mutualism regime (6,39). In the case where the handling time is zero ($h = 0$), the system usually blows up depending on the sign of the intrinsic growth rates (43). In the case where $h > 0$, it is always possible for any level of mutualistic strength to have both feasibility and local stability (39). Recall that any set of positive abundances can be turned into a feasible equilibrium point just by adjusting the intrinsic growth rates, such that the right side of Equation 2 vanishes.

Now, if we linearize the dynamics around this equilibrium point (A_i^*, P_i^*) , we get the following linear Lotka-Volterra system:

$$\begin{cases} \frac{dP_i}{dt} = P_i \left(\tilde{\alpha}_i^{(P)} - \sum_j \beta_{ij}^{(P)} P_j + \sum_j \tilde{\gamma}_{ij}^{(P)} A_j \right) \\ \frac{dA_i}{dt} = A_i \left(\tilde{\alpha}_i^{(A)} - \sum_j \beta_{ij}^{(A)} A_j + \sum_j \tilde{\gamma}_{ij}^{(A)} P_j \right), \end{cases}$$

where $\tilde{\alpha}_i^{(A)} = \sum_j \beta_{ij}^{(A)} \tilde{A}_i - \sum_j \gamma_{ij}^{(A)} P_i^*/(1 + h \sum_j \gamma_{ij}^{(A)} P_i^*)^2$ and the linearized interaction strength $\tilde{\gamma}_{ij}^{(A)} = \gamma_{ij}^{(A)}/(1 + h \sum_j \gamma_{ij}^{(A)} A_i^*)^2$ (similar expressions hold for plants). In this linearized version, the mutualistic interaction strengths are now dependent on the abundances at the equilibrium point. Moreover, the linearized interaction strengths are inversely proportional to the square of these equilibrium abundances. Then, for any level of mutualistic strength, if the abundances at the feasible equilibrium are large enough such that all eigenvalues of

$$\tilde{B} = \begin{bmatrix} \beta^{(P)} & -\tilde{\gamma}^{(P)} \\ -\tilde{\gamma}^{(A)} & \beta^{(A)} \end{bmatrix}$$

have a strictly positive real parts, this feasible equilibrium is locally stable. This simple

mathematical fact implies that now we cannot completely disentangle the feasibility conditions from the stability ones. There exists a proper subspace of the positive quadrant ($S \subset R_{\geq 0}^S$) such that when the feasible equilibrium is inside ($A_i^*, P_i^* \in S$), it is globally stable only inside the subspace S (14,43). In turn, this implies that if the intrinsic growth rates are too low, a feasible equilibrium may be unstable, and thus the domain of feasibility is also bounded by below. In general, we still have that for any level of mutualistic strength, the system can be at a feasible and locally stable equilibrium. However, it is more difficult to determine the complete set of conditions for feasibility and stability.

S5 Derivation of the structural vector

Here we provide the full analytic derivation of the structural vector. While the derivation is made with a linear version ($h = 0$) of the dynamical system (Equation 2), numerically we find that this is already a good approximation to the general solution (Fig. S4). Since we are interested in the existence of a feasible equilibrium point, i.e., $P_i^* > 0$ and $A_i^* > 0$ such that evaluated at that point the right side of Equation 2 vanishes, we have to solve the following linear system of equations:

$$\begin{bmatrix} \alpha^{(P)} \\ \alpha^{(A)} \end{bmatrix} = \left(\underbrace{\begin{bmatrix} \beta^{(P)} & 0 \\ 0 & \beta^{(A)} \end{bmatrix}}_{=C} - \underbrace{\begin{bmatrix} 0 & \gamma^{(P)} \\ \gamma^{(A)} & 0 \end{bmatrix}}_{=\Gamma} \right) \begin{bmatrix} P \\ A \end{bmatrix}. \quad (\text{S1})$$

We do this by diagonalizing the matrix $C - \Gamma$ per block, and obtaining two independent systems of equations, one for the plants and one for the animals. This is achieved by multiplying by the matrix $T = 1 + \Gamma C^{-1}$ both left sides of the equation. This is the same technique as used in Ref. 6. The new equivalent system is given by:

$$\underbrace{\begin{bmatrix} \alpha^{(P)} + \gamma^{(P)}(\beta^{(A)})^{-1}\alpha^{(A)} \\ \alpha^{(A)} + \gamma^{(A)}(\beta^{(P)})^{-1}\alpha^{(P)} \end{bmatrix}}_{= \begin{bmatrix} \alpha_{eff}^{(P)} \\ \alpha_{eff}^{(A)} \end{bmatrix}} = \underbrace{\begin{bmatrix} \beta^{(P)} - \gamma^{(P)}(\beta^{(A)})^{-1}\gamma^{(A)} & 0 \\ 0 & \beta^{(A)} - \gamma^{(A)}(\beta^{(P)})^{-1}\gamma^{(P)} \end{bmatrix}}_{= \begin{bmatrix} \beta_{eff}^{(P)} & 0 \\ 0 & \beta_{eff}^{(A)} \end{bmatrix}} \begin{bmatrix} P \\ A \end{bmatrix} \quad (\text{S2})$$

Using this mathematical transformation, we have moved from an observable parameterization space to an effective competition framework. Here, $\alpha_{eff}^{(P)}$ and $\alpha_{eff}^{(A)}$ are called effective intrinsic growth rates, and $\beta_{eff}^{(P)}$ and $\beta_{eff}^{(A)}$ are called effective competition rates. This is called so because the new linear system of equations, for the fixed point of plants and animals, is equivalent to the system of a purely competitive system within plants and within animals. Since the form of the equation is the same for the plants than for the animals, we will present the solution without the superscript (A) or (P) for the sake of clarity.

The system that we have to solve has the form $\alpha_{eff} = \beta_{eff}N$. However, here we have to keep in mind that we are interested in how the existence of feasible solutions, i.e., solutions with strictly positive species abundances ($N_i^* > 0$), are a function of α_{eff} . Since the matrix β_{eff} may not be symmetric, we solve the system using its singular value decomposition β_{eff} (59):

$$\beta_{eff} = \underbrace{\begin{bmatrix} u_{eff}^1 & u_{eff}^2 & \cdots & u_{eff}^S \end{bmatrix}}_{=U_{eff}} \underbrace{\begin{bmatrix} \lambda_1 & & & \\ & \lambda_2 & & \\ & & \ddots & \\ & & & \delta_S \end{bmatrix}}_{=D} \underbrace{\begin{bmatrix} - & v_{eff}^1 & - \\ - & v_{eff}^2 & - \\ \vdots & & \\ - & v_{eff}^S & - \end{bmatrix}}_{=V_{eff}}. \quad (\text{S3})$$

The columns of matrix U_{eff} are composed of orthonormal vectors called the left singular vectors, the rows of matrix V_{eff} are composed of orthonormal vectors called the right singular vectors. The elements of the diagonal matrix D , called the singular values, are positive and can be ordered without loss of generality, such that $\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_S \geq 0$. Assuming that all singular values are strictly positive, i.e., the matrix β_{eff} is nonsingular, the solution for the equilibrium point N^* is unique and given by

$$N^* = \beta_{eff}^{-1} \alpha_{eff} = V_{eff}^{-1} D^{-1} U_{eff}^{-1} \alpha_{eff} = \sum_{k=1}^S v_{eff}^k \frac{1}{\lambda_k} \langle u_{eff}^k | \alpha_{eff} \rangle, \quad (\text{S4})$$

where $\langle u^i | \alpha_{eff} \rangle$ denotes the scalar product between the vectors u^i and α_{eff} .

Now, we will explain how effective growth rates can be chosen such that all species have strictly positive abundances at the equilibrium point. This needs two extra assumptions on the matrix β_{eff} : all elements of matrices $\beta_{eff}^t \beta_{eff}$ and $\beta_{eff} \beta_{eff}^t$ should be positive. Then, using the Perron-Frobenius Theorem (theorem 8.4.4 in Ref. 60, this implies that all the elements of the singular vectors u^1 and v^1 are strictly positive. If we choose $\alpha_{eff} = u_{eff}^1$, then the equilibrium point, which is given by $N^* = \frac{1}{\lambda_1} v_{eff}^1$, is a feasible solution (i.e., $N_i^* > 0$). Note that any vector collinear to u_{eff}^1 leads to a feasible solution, i.e., u_{eff}^1 generates a vector of effective growth rate resulting in feasible solutions. Using effective growth rates on that specific vector, one can make any mutualistic network perfectly

persistent (see Figure ??). By imposing $N_i^* > 0$ to the solution in Equation S4, we derive the following inequality:

$$\frac{\sum_{k=1}^S \langle u_{eff}^k | \alpha_{eff} \rangle \langle v_{eff}^k | \alpha_{eff} \rangle - \langle u_{eff}^1 | \alpha_{eff} \rangle \langle v_{eff}^1 | \alpha_{eff} \rangle}{\langle u_{eff}^1 | \alpha_{eff} \rangle \langle v_{eff}^1 | \alpha_{eff} \rangle} \leq \frac{\lambda_2}{\lambda_1}. \quad (\text{S5})$$

This inequality is the generalization of Equation 20 in Ref. 6 to the case where the competition matrix β_{eff} is not necessary symmetric. This inequality holds for plants and animals and tell us that the higher the collinearity of the effective intrinsic growth rates α_{eff} with the leading singular left u_{eff}^1 and right v_{eff}^1 vectors, the higher is the chance of obtaining a solution where all species are persistent. The left side of this inequality shown in Equation S4 is called deviation from the structural vector and the sentence above can be rephrased as: the lower the deviation, the higher the chance that all species are persistent.

Effective intrinsic growth rates ($\alpha_{eff}^{(P)}$ and $\alpha_{eff}^{(A)}$) are not the observable parameters of intrinsic growth rates ($\alpha^{(P)}$ and $\alpha^{(A)}$) of plants and animals. Thus, one has to move back from the effective competition framework to the observable parameter space. This can be achieved by using the inverse of the matrix $T = 1 + \Gamma C^{-1}$ on the left and right leading singular vectors of $\beta_{eff}^{(P)}$ and $\beta_{eff}^{(A)}$:

$$\begin{bmatrix} \alpha_L^{(P)} \\ \alpha_L^{(A)} \end{bmatrix} = T^{-1} \begin{bmatrix} (u_{eff}^{(P)})^1 \\ (u_{eff}^{(A)})^1 \end{bmatrix} \quad \text{and} \quad \begin{bmatrix} \alpha_R^{(P)} \\ \alpha_R^{(A)} \end{bmatrix} = T^{-1} \begin{bmatrix} (v_{eff}^{(P)})^1 \\ (v_{eff}^{(A)})^1 \end{bmatrix} \quad (\text{S6})$$

This transformation generates results in a left and right vector of intrinsic growth rates for both the plants and the animals—what we call the structural vector. As in the effective framework, the higher the collinearity of the intrinsic growth rates with the vectors generated by these left and right structural vectors, the higher the number of persistent species.

S6 Resampling model

To study the effect of nestedness on the structural stability of a given network, we generate alternative network architectures using a resampling model. Traditionally, the Erdős-Rényi model, the swap (or fixed) model, and the probabilistic model have been used. However, as shown in Figure S8, these models generate networks with a very restricted range of nestedness.

Here, we use a resampling procedure that is able to generate a large range of nestedness. This is important in order to generate significantly different nested architectures, and in order to obtain better statistical estimates of the effect of nestedness on structural stability. This procedure is based on the matching-centrality statistical model (46). Moreover, these restrictive generative rules are biologically justified by the constraints imposed by the phylogeny (34).

The matching-centrality model aims to infer the probability of a link between two species (p_{ij} , i is an animal, and j is a plant) by assuming that the species are characterized by a latent trait of centrality (v_i^* and f_j^*) and a latent trait a matching (v_i and f_j). The model is given by

$$\text{logit}(p_{ij}) = -\kappa(v_i - f_j)^2 + \phi_1 v_i^* + \phi_2 f_j^* + m. \quad (\text{S7})$$

In this way, centrality traits quantify variability in degree, while matching traits quantify the assortative mating structure. The parameters κ , ϕ_1 , and ϕ_2 are positive scaling parameters that give the importance of the contributions of the terms. Although these latent traits are a priori unknown, they can be estimated from the network itself. Then, based on their estimation, the probability of a link between all pairs of plants and animals is estimated. Thus, a new network can simply be generated by drawing randomly the links based on those estimated interaction probabilities. The expected number of links, as well as the expected degree of the sampled network are equal to the ones of the

observed networks. However, this model, which can be viewed as a probabilistic version of the swap model, also generates a restricted range of nestedness. To relax this constraint, we generalize the model by introducing a control parameter, that we call temperature T , that modulates the level of stochasticity in the model:

$$\text{logit}(p(T)_{ij}) = \frac{1}{T} (-\kappa(v_i - f_j)^2 + \phi_1 v_i^* + \phi_2 f_j^*) + m(T). \quad (\text{S8})$$

Here, the intercept $m(T)$ is adjusted for each temperature value such that the expected number of links is equal to the observed one. When the temperature goes to infinite, our model converges to the Erdős-Rényi model, when the temperature goes to zero, the system freezes in the most probable configuration predicted by our model.

As it can be observed in Figure S4, when varying the temperature, our model can reproduce a larger range of nestedness compared to the ones generated by alternative sampling models. Importantly, our model predicts also a maximum and a minimum level of nestedness that the system can reach, when generative rules (represented by the matching and centrality traits) are introduced.

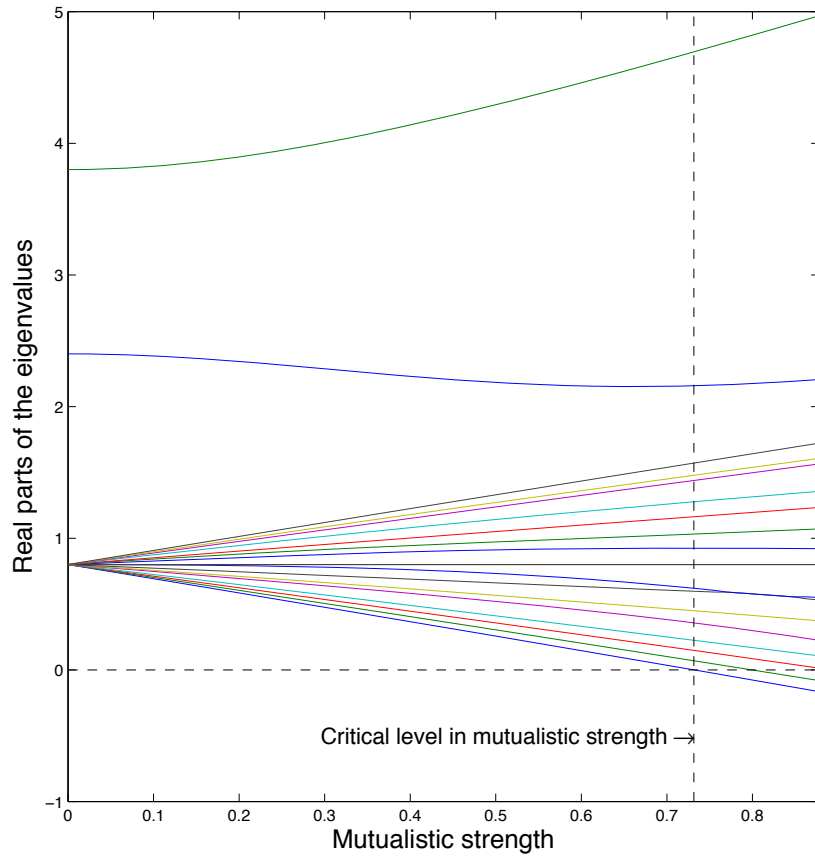


Fig. S1: Real part of the eigenvalues of matrix B as a function of the mutualistic strength (γ_0). Here we use the empirical network located in Yakushima Island, Japan (see Table S1), a mutualistic trade-off $\delta = 0.5$, and an interspecific competition $\rho = 0$. The vertical dashed line represents the critical value of mutualistic strength γ_0^r above which at least one of the eigenvalues has a negative real part.

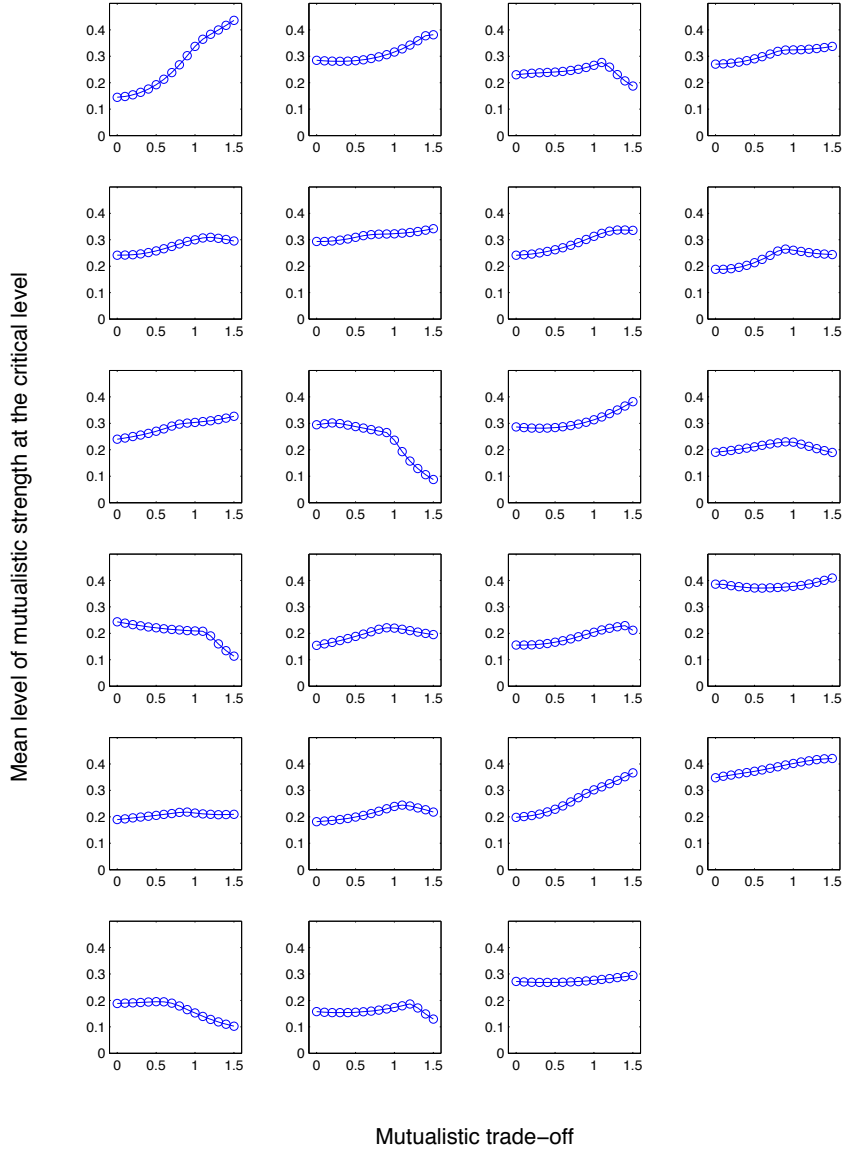


Fig. S2: Mutualistic trade-off and critical value of mutualistic strength. For each of 23 observed mutualistic networks (see Table S1), this figure illustrates how the mutualistic trade-off influences the maximum level of mean mutualistic strength that the system can handle to guarantee the stability of any feasible equilibrium. One can see that, in the majority of cases, the mutualistic trade-off enhances the level of mutualistic strength.

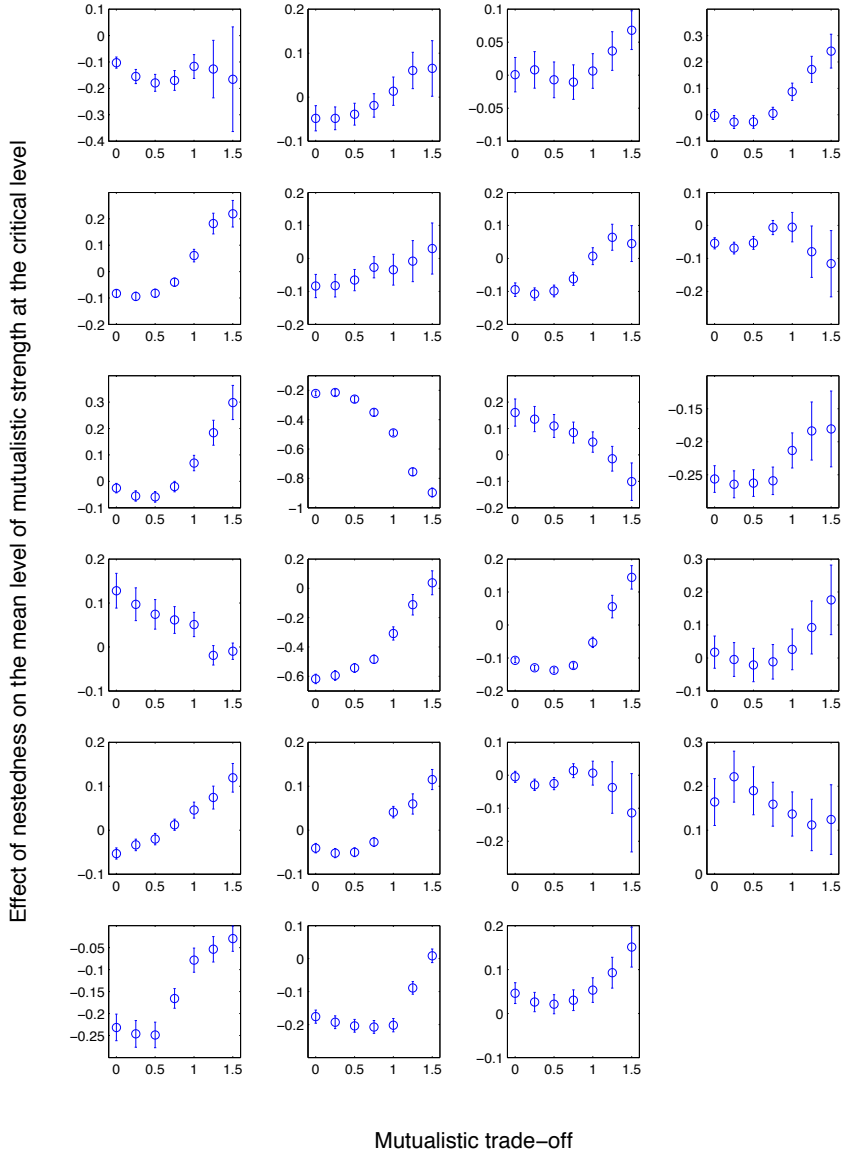


Fig. S3: Nestedness and critical value of mutualistic strength. For each of 23 observed mutualistic networks (Table S1), this figure illustrates, as function of the mutualistic trade off, the linear effect of nestedness on the maximum level of mean mutualistic strength that the system can handle to guarantee the stability of any feasible equilibrium. One can see that in the majority of cases, the slopes of this effect increase along with the mutualistic trade-off.

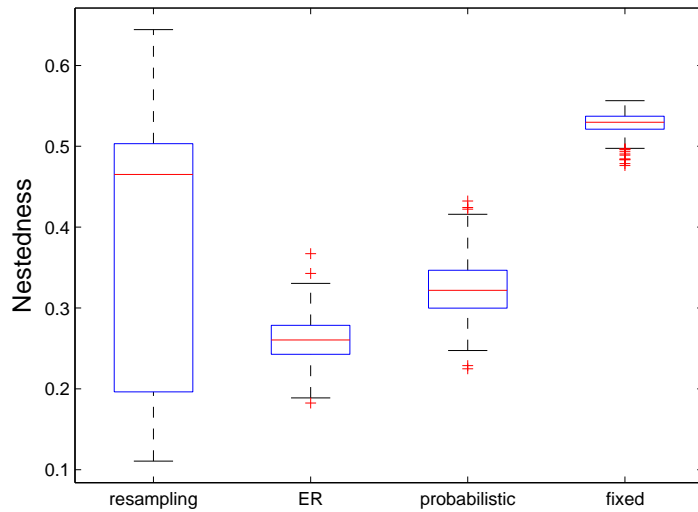


Fig. S4: Model-generated nested architectures. For the observed mutualistic network in the Snowy Mountains, Australia (see Table S1), the figure shows the different distributions of nestedness values generated by different models: our resampling model, the Erdős-Rényi model, the probabilistic model, and the fixed or swap model.

S7 Structural stability across observed mutualistic networks

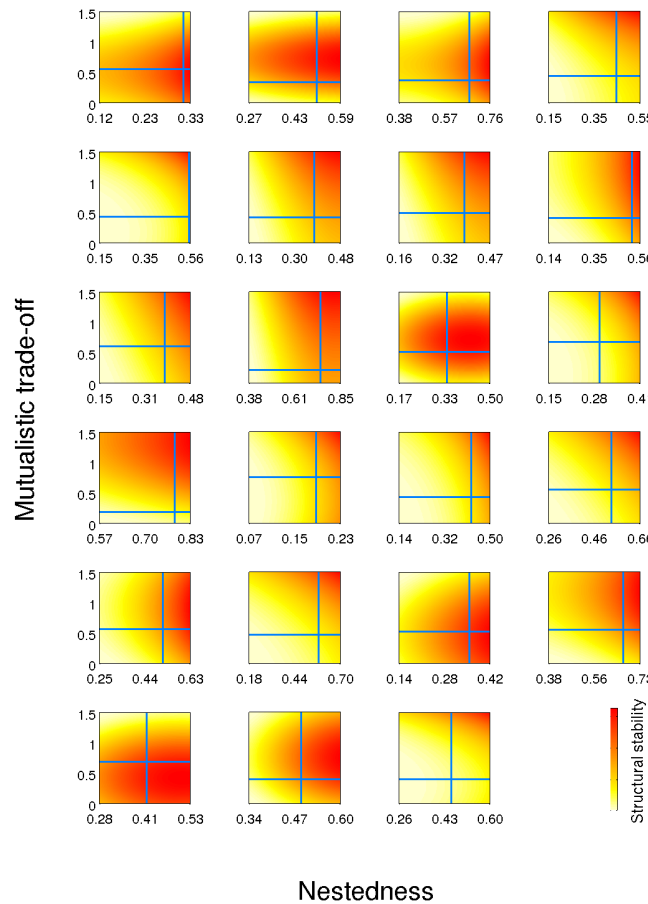
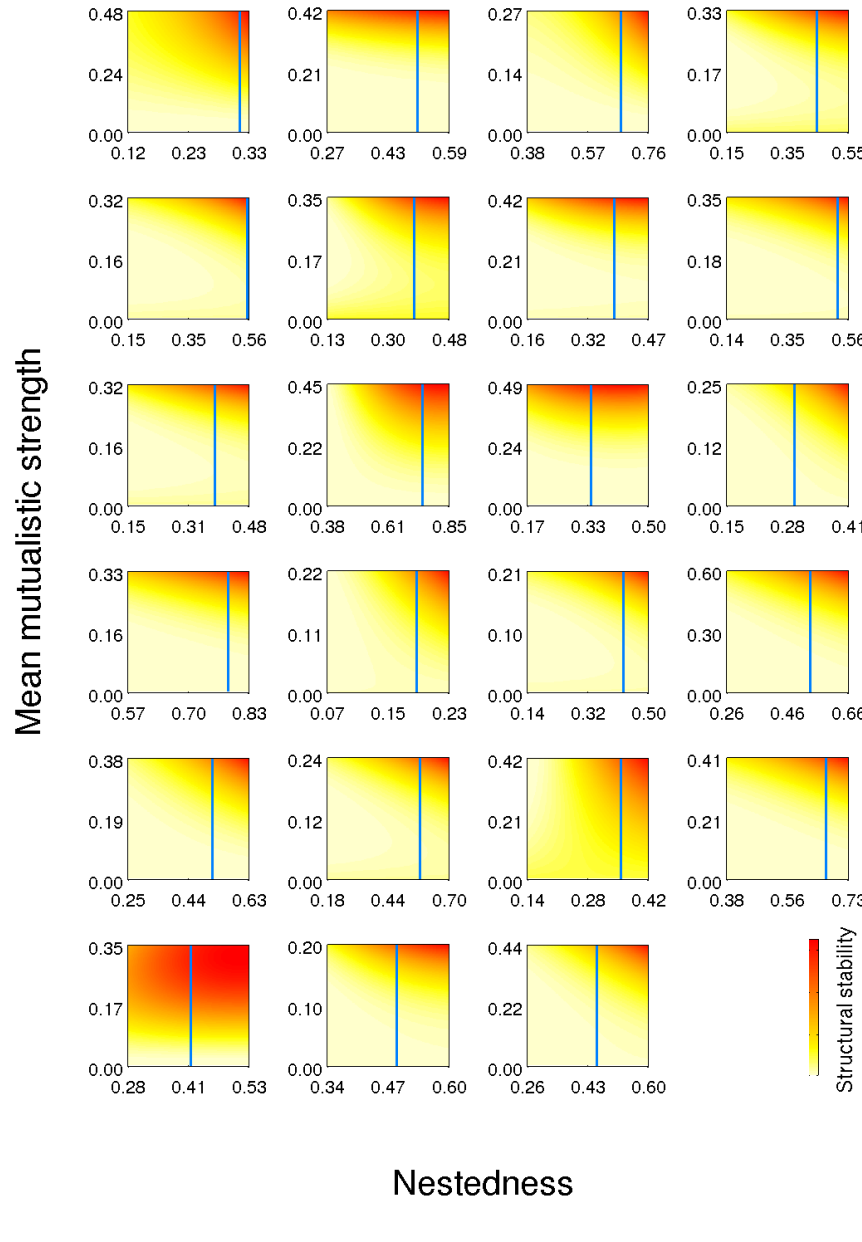


Fig. S5: Structural stability in all the observed networks. The darker the color, the larger the positive effect on the area of structural stability. The lightest regions correspond to architectures with a negative effect on this area. Dashed lines correspond to different values of mean strength level. Solid lines correspond to the observed values of nestedness and mutualistic trade-offs.



Nestedness

Fig. S6: Structural stability and nestedness. Keeping the observed mutualistic trade-off fixed, the figure shows the extent to which the observed nestedness (solid line) can modulate the area of structural stability (partial fitted residuals) across different values of mean mutualistic strength.

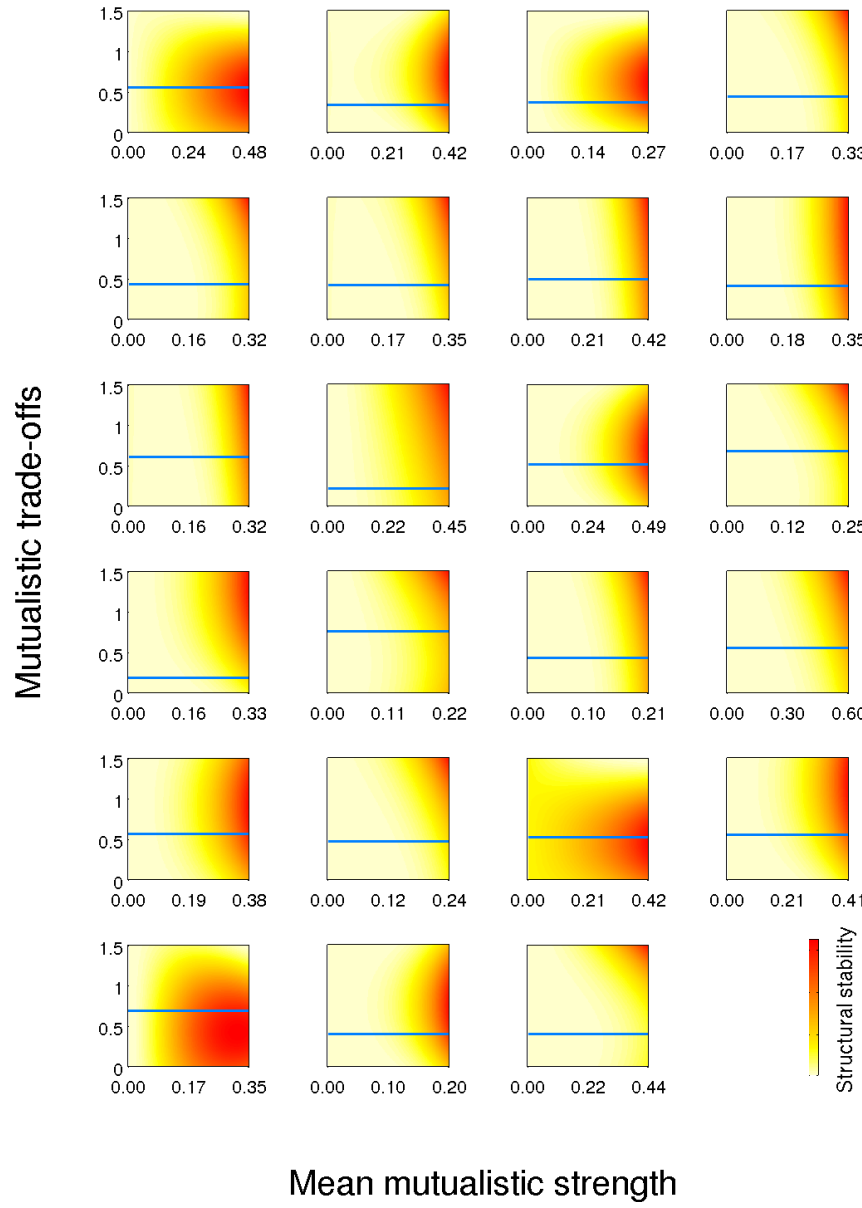


Fig. S7: Structural stability and mutualistic trade-offs. Keeping the observed nestedness fixed, the figure shows the extent to which the observed mutualistic trade-off (solid line) expands the area of structural stability (partial fitted residuals) across different values of mean mutualistic strength.

S8 Robustness of results to the choice of ρ and h

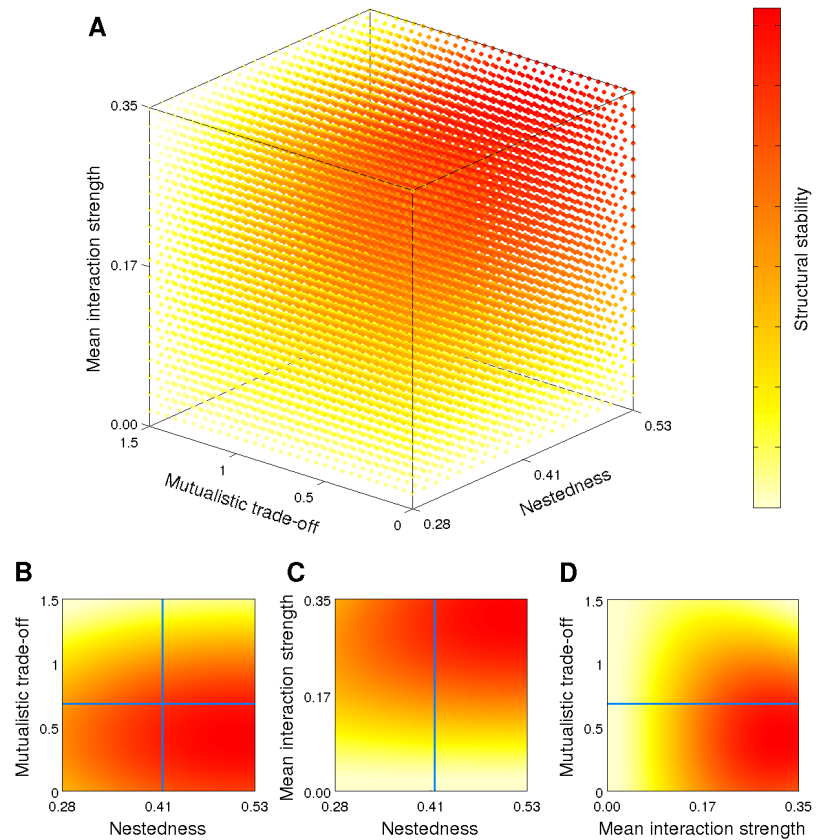


Fig. S8: Equivalent to Figure 5 in main text but where handling time is now sampled from a uniform distribution between 0.08 and 0.12 for each of the species.

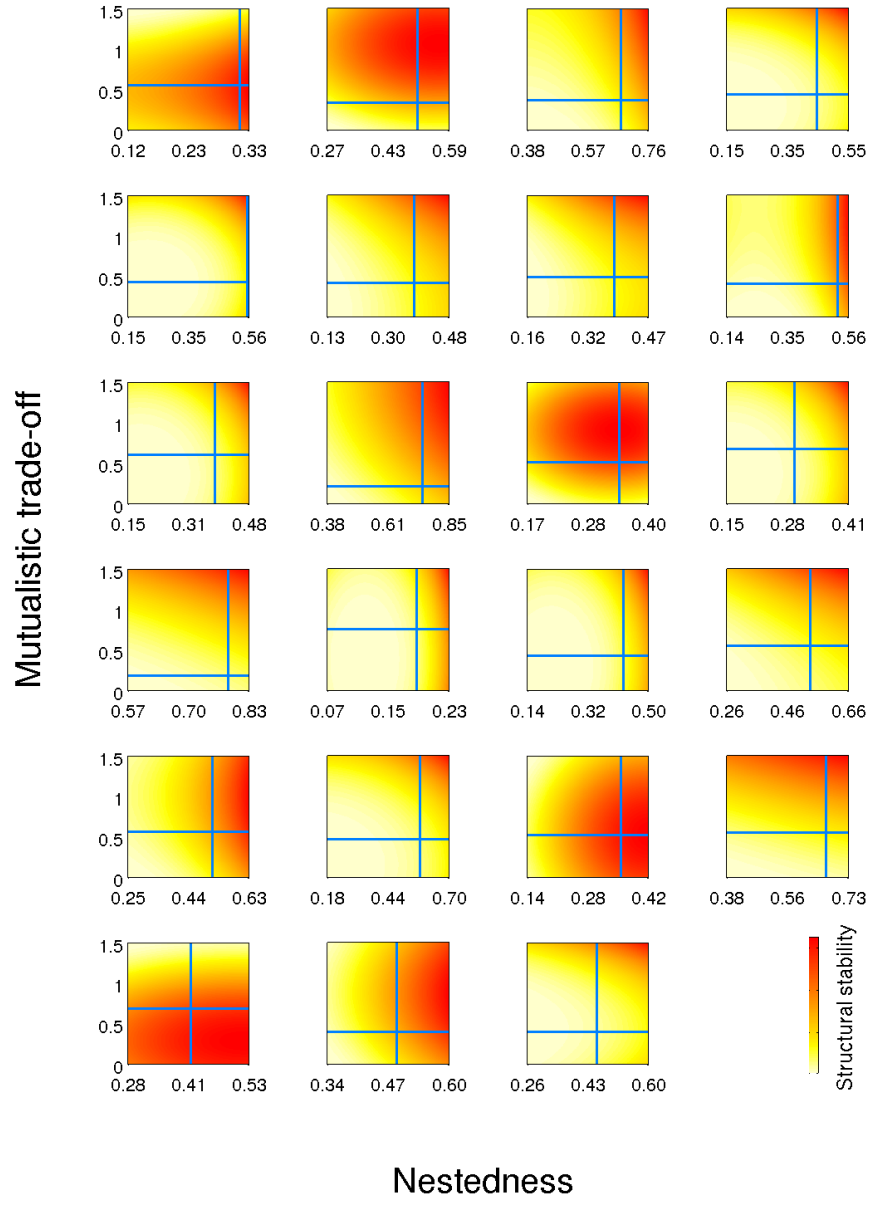


Fig. S9: Equivalent to Figure S5 in SM but with $\rho = 0.4$.

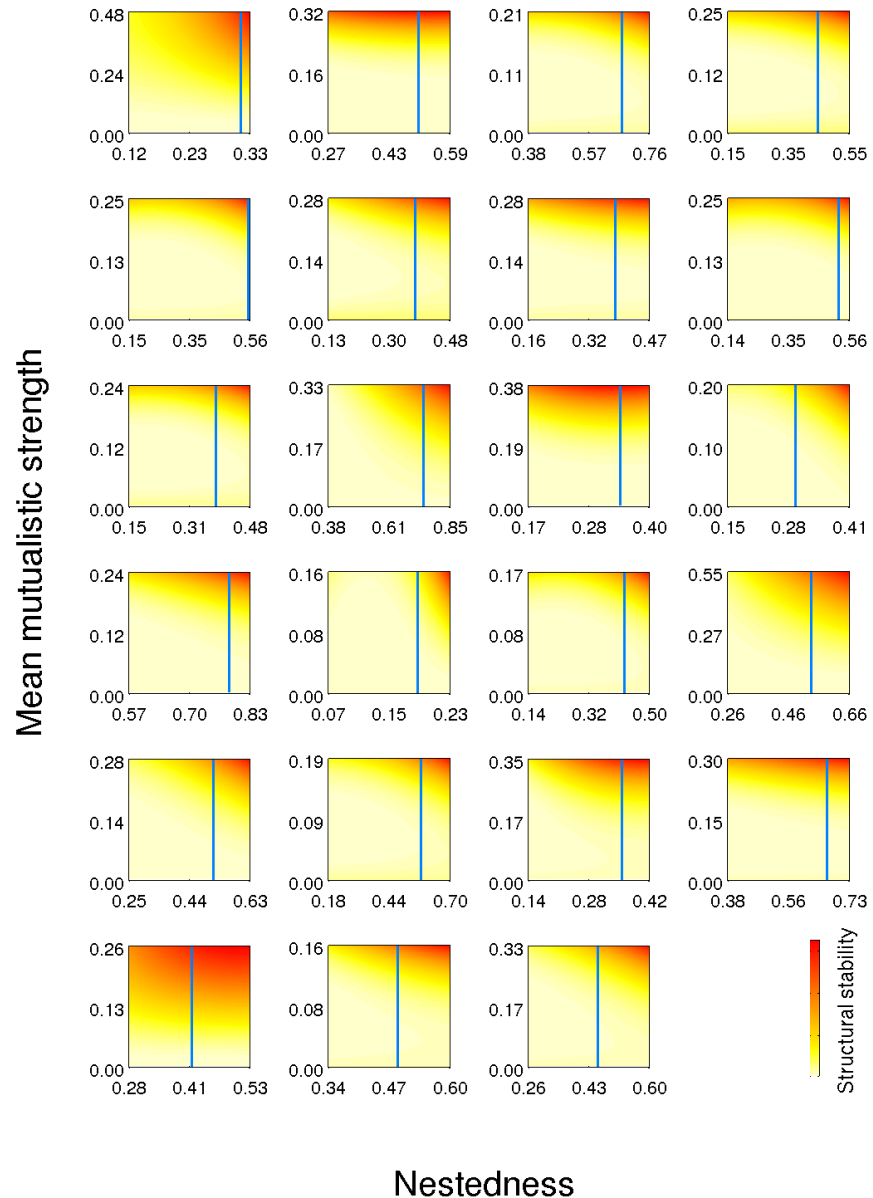


Fig. S10: Equivalent to Figure S6 in SM but with $\rho = 0.4$.

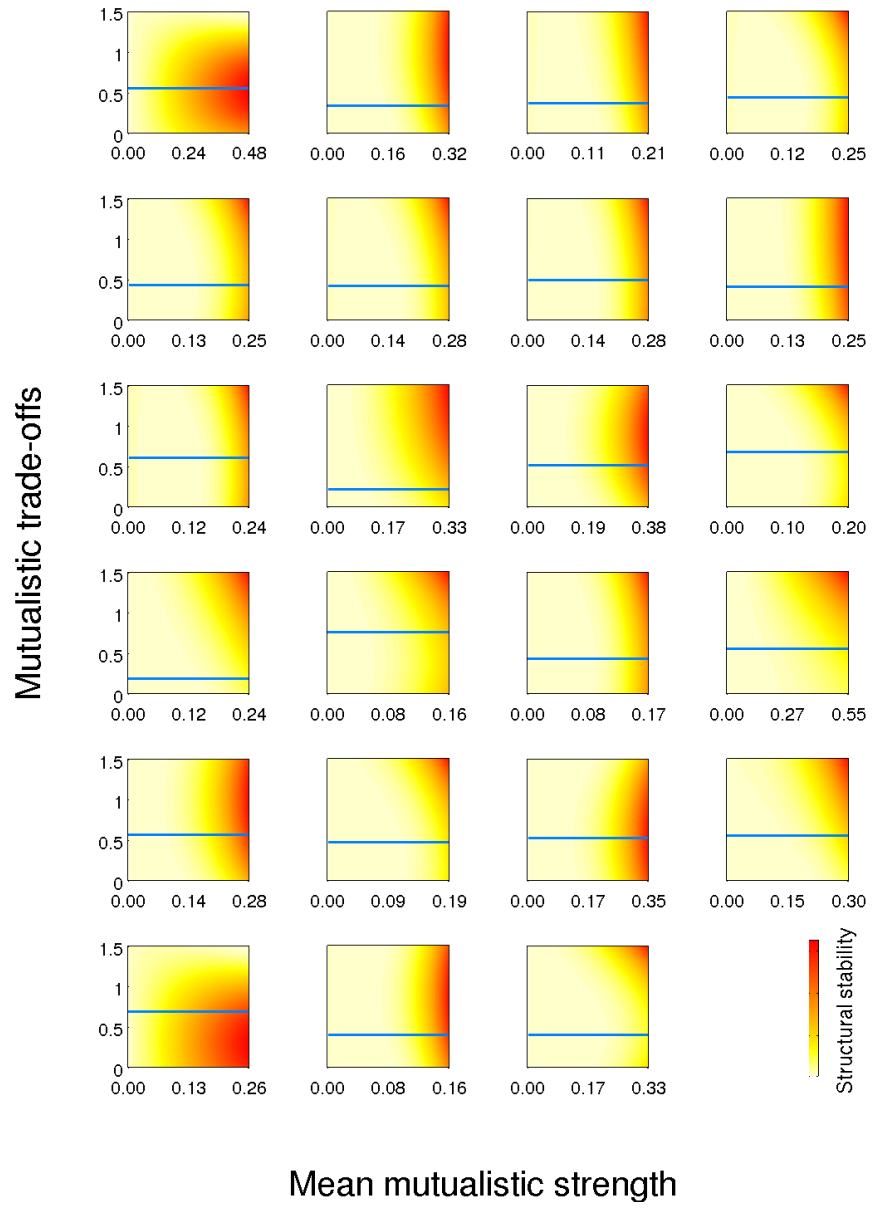


Fig. S11: Equivalent to Figure S7 in SM but with $\rho = 0.4$.

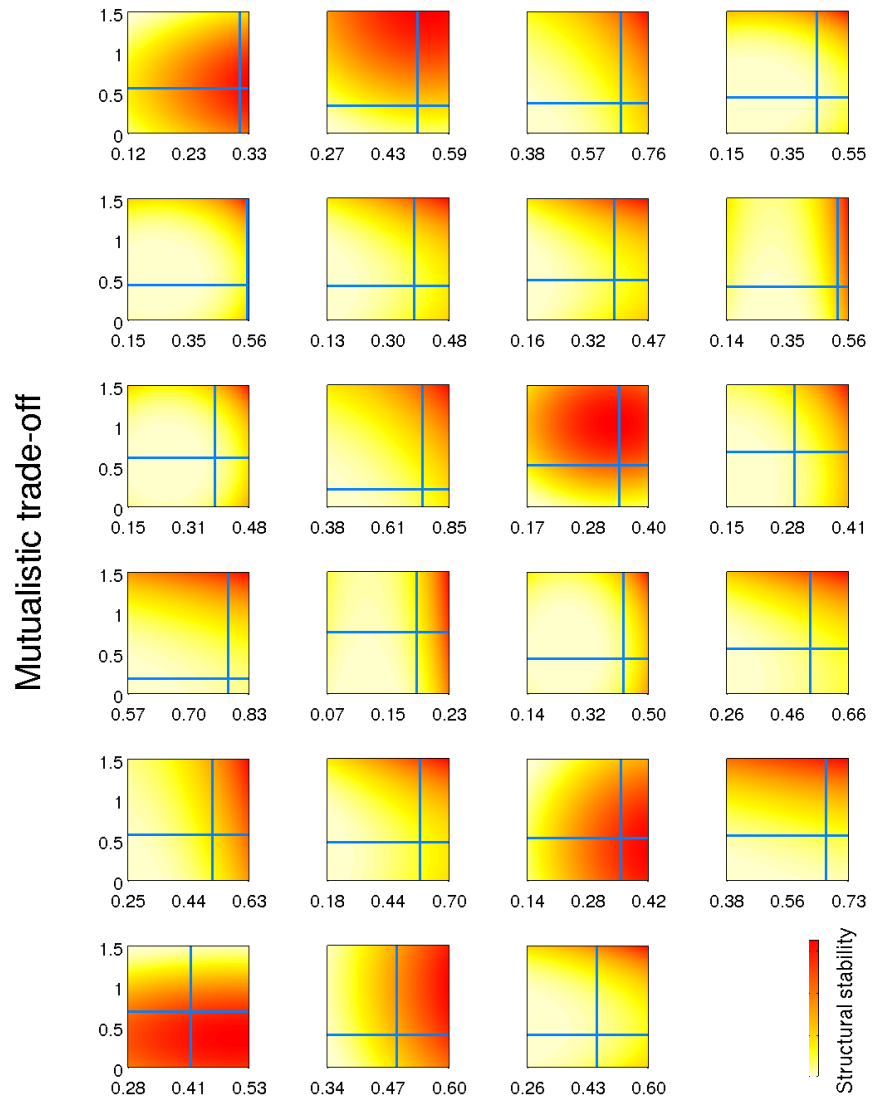


Fig. S12: Equivalent to Figure S5 in SM but with $\rho = 0.6$.

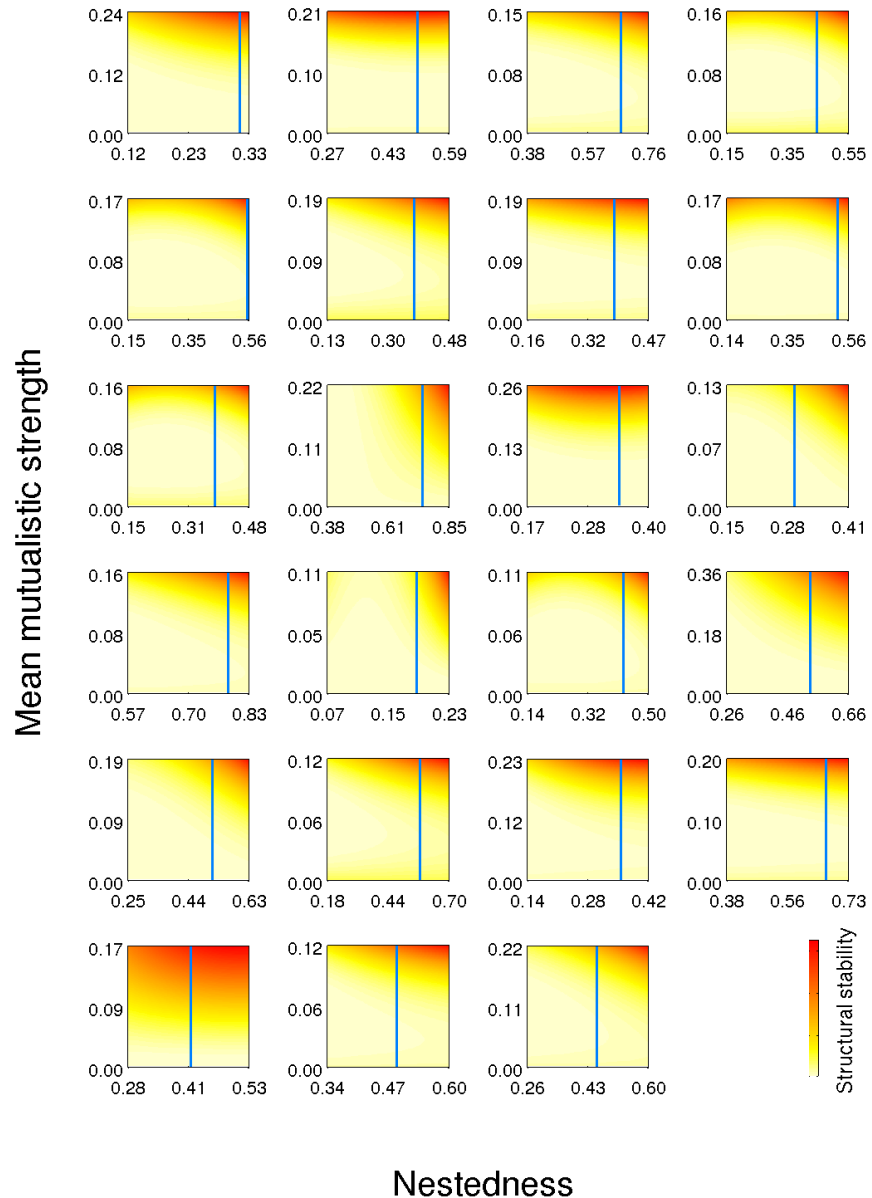


Fig. S13: Equivalent to Figure S6 in SM but with $\rho = 0.6$.

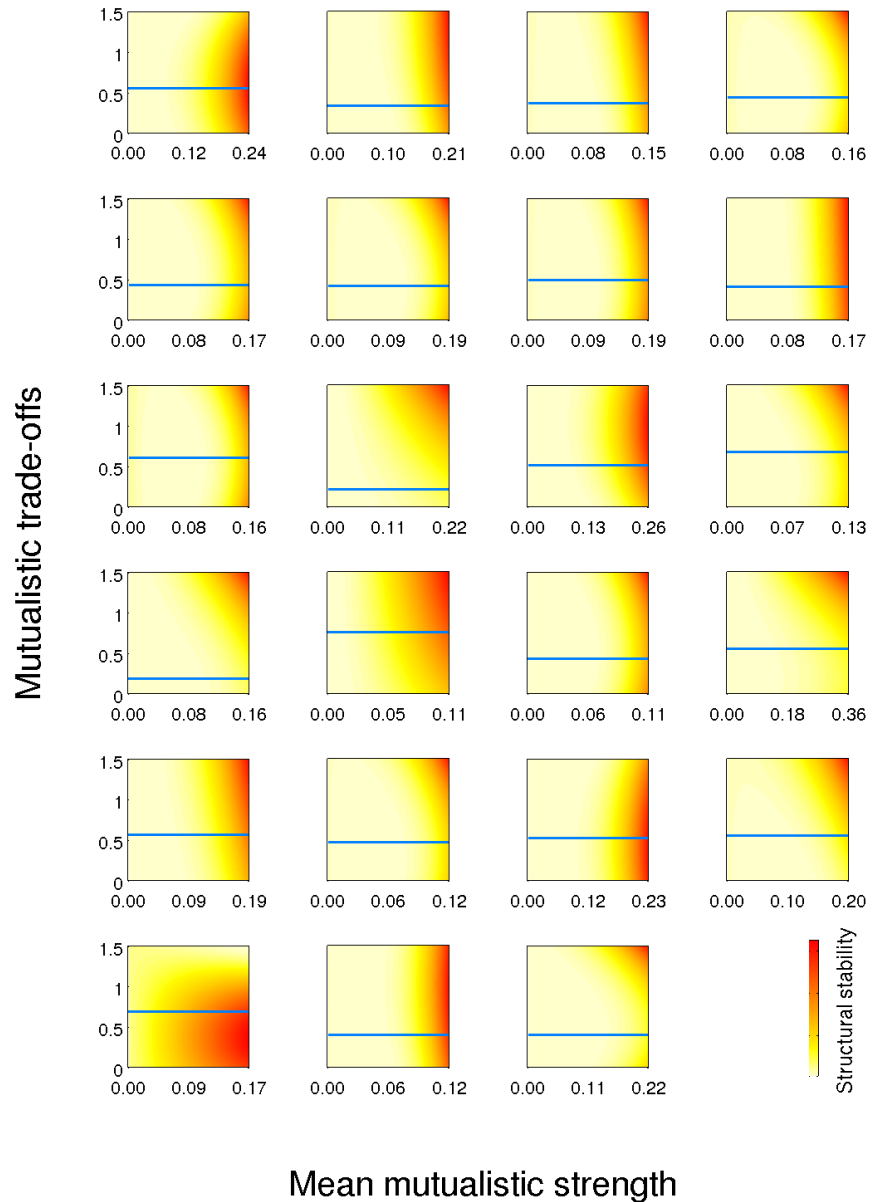


Fig. S14: Equivalent to Figure S7 in SM but with $\rho = 0.6$.

S9 Data

| Network | Animals | Plants | Interactions | Location |
|----------|---------|--------|--------------|---|
| M_PL_004 | 101 | 12 | 167 | Central New Brunswick, Canada |
| M_SD_001 | 21 | 7 | 50 | Princeton, Mercer, New Jersey, USA |
| M_SD_002 | 9 | 31 | 119 | Mont Missim, Morobe Prov., New Guinea |
| M_SD_003 | 16 | 25 | 68 | Caguana, Puerto Rico |
| M_SD_004 | 20 | 34 | 95 | Cialitos, Puerto Rico |
| M_SD_005 | 13 | 25 | 49 | Cordillera, Puerto Rico |
| M_SD_006 | 15 | 21 | 51 | Fronton, Puerto Rico |
| M_PL_006 | 61 | 17 | 146 | Hickling, Norfolk, UK |
| M_PL_007 | 36 | 16 | 85 | Shelfanger, Norfolk, UK |
| M_SD_008 | 10 | 16 | 110 | Mtunzini, South Africa |
| M_SD_009 | 18 | 7 | 38 | Santa Genebra Reserve T1 SE, Brazil |
| M_SD_010 | 14 | 50 | 234 | Tropical rainforest, Trinidad |
| M_SD_011 | 14 | 11 | 47 | Calton, UK |
| M_PL_013 | 56 | 9 | 103 | KwaZulu-Natal region, South Africa |
| M_SD_012 | 29 | 35 | 146 | Santa Genebra reserve T2. SE, Brazil |
| M_PL_017 | 79 | 25 | 299 | Bristol, UK |
| M_SD_014 | 17 | 16 | 121 | Hato Raton, Sevilla, Spain |
| M_PL_019 | 85 | 40 | 264 | Snowy Mountains, Australia |
| M_PL_024 | 18 | 11 | 38 | Melville, Island, Canada |
| M_PL_025 | 44 | 13 | 143 | North Carolina, USA |
| M_PL_033 | 34 | 13 | 141 | Ottawa, Canada |
| M_SD_020 | 33 | 25 | 150 | Nava Correhuelas, S. Cazorla, SE, Spain |
| M_SD_023 | 8 | 15 | 38 | Yakushima Island, Japan |

Table S1: Data set. We apply our study to the 23 quantitative mutualistic networks published at www.web-of-life.es at the time this analysis was conducted (see also Ref. 34). These networks represent diverse environmental and biotic conditions. Network identifiers consists on a combination of letters and a specific number: M_PL and M_SD stand for mutualistic networks of pollination and seed dispersal, respectively. The order of the networks matches the one used across the panels of Fig. 6.

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18. Any matrix B is called Lyapunov-stable if the real parts of all its eigenvalues are positive, meaning that a feasible equilibrium at which all species have the same abundance is at least locally stable. A matrix B is called D -stable if DB is a Lyapunov-stable matrix for any strictly positive diagonal matrix D . D -stability is a stronger condition than Lyapunov stability in the sense that it grants the local stability of any feasible equilibrium (15). In addition, a matrix B is called Lyapunov diagonally stable if there exists a strictly positive diagonal matrix D so that $DB + B^tD$ is a Lyapunov-stable matrix. This notion of stability is even stronger than D -stability in the sense that it grants not only the local stability of any feasible equilibrium point, but also its global stability (14). A Lyapunov-diagonally stable matrix has all of its principal minors positive. Also, its stable equilibrium (which may be only partially feasible; some species may have an abundance of zero) is specific (57). This means that there is no alternative stable state for a given parameterization of intrinsic growth rates (55). We have chosen the convention of having a minus sign in front of the interaction-strength matrix B . This implies that B has to be positive Lyapunov-stable (the real parts of all eigenvalues have to be strictly positive) so that the equilibrium point of species abundance is locally stable. This same convention applies for D -stability and Lyapunov-diagonal stability.
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21. All simulations were performed by integrating the system of ordinary differential equations by using the function `ode45` of Matlab. Species are considered to have gone extinct when their abundance is lower than 100 times the machine precision. Although we present the results of our simulations with a fixed value of $\rho = 0.2$ and $h = 0.1$, our results are robust to this choice as long as $\rho < 1$ (45). In our dynamical system, the units of the parameters are abundance for N , 1/time for α and h , and 1/(time · biomass) for β and γ . Abundance and time can be any unit (such as biomass and years), as long as they are constant for all species.
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48. For a given network, we look at the association of the fraction of surviving species with the deviation from the structural vector of plants $\eta_{(P)}$ and animals $\eta_{(A)}$, nestedness N , mutualistic trade-off δ , and mean level of mutualistic strength $\bar{\gamma} = \langle \gamma_{ij} \rangle$ using the following binomial generalized linear model (47):

$$\text{log it}(\text{probabilityofsurviving}) \sim \log(\eta_A) + \log(\eta_P) + \bar{\gamma} + \bar{\gamma}^2 + \bar{\gamma} \cdot N + \bar{\gamma} \cdot N^2 + \bar{\gamma} \cdot \delta + \bar{\gamma} \delta^2.$$

Obviously, at a level of mutualism of zero ($\gamma = 0$), nestedness and mutualistic trade-off cannot influence the probability of a species to survive. We have to include an interaction between the mean level of mutualism and the nestedness and mutualistic trade-off. We have also included a quadratic term in order to take into account potential nonlinear effects of nestedness, mutualistic trade-off, and mean level of mutualistic strength. The effect of network architecture is confirmed by the significant likelihood ratio between the full model and a null model without such a network effect ($P < 0.001$) for all the observed empirical networks. The effect of network architecture on structural stability can be quantified by the partial fitted values defined as follows:

$$\text{partialfittedvalues} = \hat{\beta}_1 \bar{\gamma} + \hat{\beta}_2 \bar{\gamma}^2 + \hat{\beta}_3 \bar{\gamma} \cdot N + \hat{\beta}_4 \bar{\gamma} \cdot N^2 + \hat{\beta}_5 \bar{\gamma} \cdot \delta + \hat{\beta}_6 \bar{\gamma} \cdot \delta^2,$$
where $\hat{\beta}_1 \dots \hat{\beta}_6$ are the fitted parameters corresponding to the terms $\bar{\gamma}$ to $\bar{\gamma} \cdot \delta^2$, respectively.

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52. To estimate the mutualistic trade-off, δ , from an empirical point of view, we proceed as follows. First, there is available data on the frequency of interactions. Thus, q_{ij} is the observed number of visits of animal species j on plant species i . This quantity has been proven to be the best surrogate of per capita effects of one species on another (γ_{ij}^P and γ_{ij}^A) (41). Second, the networks provide information on the number of species one species interacts with (its degree or generalization level; k_i^A and k_i^P). Following (41, 50, 51), the generalization level of a species has been found to be proportional to its abundance at equilibrium. Thus, the division of the total number of visits by the product of the degree of plants and animals can be assumed to be proportional to the interaction strengths.

Mathematically, we obtain two equations, one for the effect of the animals on the plants and vice versa: $(q_{ij})/(k_i^P k_j^A) \propto \gamma_{ij}^P$ and $(q_{ij})/(k_i^A k_j^P) \propto \gamma_{ij}^A$. Introducing the explicit dependence between interaction strength and trade-off—for example, $\gamma_{ij}^P = \gamma_0 / (k_i^P)^\delta$ and $\gamma_{ij}^A = \gamma_0 / (k_i^A)^\delta$ —we obtain $(q_{ij})/(k_i^P k_j^A) \propto (\gamma_0) / [(k_i^P)^\delta]$ and $(q_{ij})/(k_i^A k_j^P) \propto (\gamma_0) / [(k_i^A)^\delta]$. In order to estimate the value of δ , we can just take the logarithm on both sides of the previous equations for the data excluding the zeroes. Then, δ is simply given by the slope of the following linear regressions:

$$\log\left(\frac{q_{ij}}{k_i^P k_j^A}\right) = a^P - \delta \log(k_i^P) \text{ and } \log\left(\frac{q_{ij}}{k_i^A k_j^P}\right) = a^A - \delta \log(k_i^A), \text{ where } a^P \text{ and } a^A \text{ are the intercepts for plants and animals, respectively.}$$

These two regressions are performed simultaneously by lumping together the data set. The intercept for the effect of animals on plants (a^P) may not be the same as the intercept for the effect of the plants on the animals (a^A).

53. The competition matrix given by $\beta = \begin{bmatrix} 3 & 3.1 & 1 \\ 5 & 5 & 6 \\ 4 & 1 & 3 \end{bmatrix}$ is Lyapunov-stable but not D -stable.

Because Lyapunov diagonal stability implies D -stability, this matrix is also not Lyapunov-diagonally stable.

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