

Niche Theory for Mutualism: A Graphical Approach to Plant-Pollinator Network Dynamics

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ABSTRACT: Contemporary niche theory is a useful framework for understanding how organisms interact with each other and with their shared environment. Its graphical representation, popularized by Tilman's resource ratio hypothesis, facilitates analysis of the equilibrium structure of complex dynamical models, including species coexistence. This theory has been applied primarily to resource competition since its early beginnings. Here, we integrate mutualism into niche theory by expanding Tilman's graphical representation to the analysis of consumer-resource dynamics of plant-pollinator networks. We graphically explain the qualitative phenomena previously found by numerical simulations, including the effects on community dynamics of nestedness, adaptive foraging, and pollinator invasions. Our graphical approach promotes the unification of niche and network theories and deepens the synthesis of different types of interactions within a consumer-resource framework.

Keywords: contemporary niche theory, ecological networks, mutualistic interactions, pollination ecology, competition for pollination, competition for floral rewards.

Introduction

Mutualistic interactions pervade every type of ecosystem and level of organization on Earth (Boucher et al. 1982; Bronstein 2015). Mutualisms such as pollination (Ollerton et al. 2011), seed dispersal (Wang and Smith 2002), coral symbioses (Rowan 2004), and nitrogen-fixing associations between plants and legumes, bacteria, or fungi (Horton and Bruns 2001; van der Heijden et al. 2008) sustain the pro-

ductivity and biodiversity of most ecosystems on the planet and human food security (Potts et al. 2016; Ollerton 2017). However, ecological theory on mutualisms has been scarce and less integrated than for predation and competition, which hinders our ability to protect, manage, and restore mutualistic systems (Vandermeer and Boucher 1978; Bascompte and Jordano 2014; Bronstein 2015). This scarce theoretical development is of particular concern because several mutualisms (such as coral-algae and plant-pollinator mutualisms) that play a critical role in the functioning of ecosystems are currently under threat (Brown 1997; Rowan 2004; Goulson et al. 2015; Ollerton 2017). In particular, niche theory (MacArthur 1969, 1970; Tilman 1982; Leibold 1995; Chase and Leibold 2003) for mutualisms has only recently started to be developed (Peay 2016; Johnson and Bronstein 2019). Chase and Leibold (2003) suggest that contemporary niche theory can be expanded to mutualism, but such a suggestion has yet to be explored. Here, we expand niche theory to mutualistic networks of plant-pollinator interactions by further developing the graphical approach popularized by Tilman (1982) to analyze a consumer-resource dynamic model of plant-pollinator networks developed, analyzed, and tested by Valdovinos et al. (2013, 2016, 2018).

For about 70 years, theoretical research analyzing the population dynamics of mutualisms roughly only assumed Lotka-Volterra-type models (sensu Valdovinos 2019) to conduct their studies (e.g., Kostitzin 1934; Gause and Witt 1935; Vandermeer and Boucher 1978; Wolin and Lawlor 1984; Bascompte et al. 2006; Okuyama and Holland 2008; Bastolla et al. 2009). Those models represent mutualistic relationships as direct positive effects between species using a (linear or saturating) positive term in the growth equation of each mutualist that depends on the population size of its mutualistic partner. While this research increased our understanding of the effects of facultative, obligate, linear, and saturating mutualisms on the long-term stability of mutualistic systems, a more sophisticated understanding of their dynamics (e.g., transients) and of phenomena

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beyond the simplistic assumptions of the Lotka-Volterra-type models was extremely scarce. A more mechanistic consumer-resource approach to mutualisms was recently proposed by Holland and colleagues (Holland et al. 2005; Holland and DeAngelis 2010) and further developed by Valdovinos et al. (2013, 2016, 2018). This approach decomposes the net effects assumed to be always positive by Lotka-Volterra-type models into the biological mechanisms producing those effects, including the gathering of resources and exchange of services.

The key advance of the consumer-resource model developed by Valdovinos et al. (2013) is separating the dynamics of the plants' vegetative biomass from the dynamics of the plants' floral rewards. This separation allows (i) tracking the depletion of floral rewards by pollinator consumption, (ii) evaluating exploitative competition among pollinator species consuming the floral rewards provided by the same plant species, and (iii) incorporating the capability of pollinators (adaptive foraging) to behaviorally increase their foraging effort on the plant species in their diet with more floral rewards available. Another advance of this model is incorporating the dilution of conspecific pollen carried by pollinators, which allows tracking competition among plant species for the quality of pollinator visits (see the next section).

This contribution analyzes the dynamics of plant-pollinator networks when all of the above-mentioned biological mechanisms are considered. Specifically, we provide analytical understanding for the results found with extensive numerical simulations (Valdovinos et al. 2013, 2016, 2018; hereafter, "previous simulations") and generalize some of them beyond the original simulation conditions. By "analytical understanding" we refer to finding those results using a graphical approach whose geometry rigorously reflects mathematical analysis (Tilman 1982; Koffel et al. 2016; also provided in our appendices). Our methods describe Valdovinos et al.'s model and our graphical approach, including the conditions for coexistence among adaptive pollinators sharing floral rewards and how we use projections to analyze high-dimensional systems. Our results first demonstrate the effects of nestedness on species coexistence in networks without adaptive foraging found by previous simulations (Valdovinos et al. 2016). Nestedness is the tendency of generalists (species with many interactions) to interact with both generalists and specialists (species with one or a few interactions) and of specialists to interact with only generalists. Second, we demonstrate the effects of adaptive foraging on species coexistence in nested networks found by the same simulation study. Third, we demonstrate the impacts of pollinator invasions on native pollinators in nested networks with adaptive foraging found numerically by Valdovinos et al. (2018). Finally, we discuss how our approach helps to integrate niche and network theories and deep-

ens the synthesis of different types of interactions within a consumer-resource framework.

Methods

Dynamical Model of Plant-Pollinator Interactions

Valdovinos et al. (2013) model the population dynamics of each plant and pollinator species of the network as well as the dynamics of floral rewards and pollinators' foraging preferences (see table 1 for definitions of variables and parameters). Four functions define these dynamics. The function $V_{ij}(p_i, a_j) = \alpha_{ij}\tau_{ij}a_jp_i$ represents the visitation rate of animal species j to plant species i and connects the dynamics of plants, animals, rewards, and foraging preferences. An increase in visits increases plant growth rate via pollination and animal growth rate via rewards consumption but decreases rewards availability. The function

$$\sigma_{ij}(p_k) = \frac{\alpha_{ij}\tau_{ij}p_i}{\sum_{k \in P_i} \alpha_{kj}\tau_{kj}p_k}$$

represents the fraction of j 's visits that successfully pollinate plant i and accounts for the dilution of plant i 's pollen when j visits other plant species. Pollinators visiting many different plant species carry more diluted pollen (low-quality visits) than the pollen carried by pollinators visiting only one plant species (high-quality visits). The function $\gamma_i(p_k) = g_i(1 - \sum_{l \neq i \in P} u_l p_l - w_i p_i)$ represents the germination rate of the seeds produced by the successful pollination events, where g_i is the maximum fraction of i

Table 1: Model variables and parameters

Symbol	Meaning
Variables:	
p_i	Plant abundance per unit area ([ind.]/m ²)
a_j	Animal (pollinator) abundance per unit area ([ind.]/m ²)
R_i	Reward abundance per unit area (g/m ²)
α_{ij}	Foraging preference (dimensionless)
Parameters:	
g_i	Maximum germination rate ([ind.]/[seeds])
u_i	Plant interspecific competition (m ² /[ind.])
w_i	Plant intraspecific competition (m ² /[ind.])
e_{ij}	Expected number of seeds per pollination event ([seeds]/[visits])
τ_{ij}	Visitation efficiency ([visits]m ² /[ind.] ² year)
$\mu_i^{P/A}$	Mortality rates (1/year)
c_{ij}	Conversion efficiency of rewards into animal abundance ([ind.]/g)
b_{ij}	Per-visit rewards extraction (1/[visits])
β_i	Per-plant reward production (g/[ind.]/year)
ϕ_i	Self-limitation of reward production (1/year)
G_j	Adaptation rate (dimensionless)

recruits subjected to both interspecific (u_i) and intraspecific (w_i) competition. Finally, the function

$$f_{ij}(R_i/p_i) = b_{ij} \frac{R_i}{p_i}$$

represents the rewards consumption by animal j in each of its visits to plant i . These functions capturing the above mentioned biological processes lead to the following equations:

$$\frac{dp_i}{dt} = \overbrace{\gamma_i(p_k)}^{\text{germination rate}} \sum_{j \in A_i} \overbrace{e_{ij} \sigma_{ij}(p_k) V_{ij}(p_i, a_j)}^{\text{seed production}} - \overbrace{\mu_i^p p_i}^{\text{mortality}}, \quad (1)$$

$$\frac{da_j}{dt} = \sum_{i \in P_j} \overbrace{c_{ij} V_{ij}(p_i, a_j) f_{ij}(R_i/p_i)}^{\text{rewards consumption}} - \overbrace{\mu_j^A a_j}^{\text{mortality}}, \quad (2)$$

$$\frac{dR_i}{dt} = p_i \left[\overbrace{\beta_i - \phi_i \frac{R_i}{p_i}}^{\text{per-plant rewards production}} \right] - \sum_{j \in A_i} \overbrace{V_{ij}(p_i, a_j) f_{ij}(R_i/p_i)}^{\text{rewards consumption}}, \quad (3)$$

$$\frac{d\alpha_{ij}}{dt} = \frac{G_j \alpha_{ij}}{a_j} \left(\overbrace{c_{ij} V_{ij}^s(p_i, a_j) f_{ij}(R_i/p_i)}^{\text{rewards consumption as specialist}} - \sum_{k \in P_j} \overbrace{c_{kj} V_{kj}(p_k, a_j) f_{kj}(R_k/p_k)}^{\text{actual rewards consumption}} \right), \quad (4)$$

where $V_{ij}^s = \tau_{ij} a_j p_i$ is the visitation rate of animal species j to plant species i under a pure specialist strategy $\alpha_{ij} = 1$. That is, the preference of animal j for plant i increases when the rewards that could be extracted from plant species i by application of full foraging effort to that plant ($\alpha_{ij} = 1$) exceed the rewards currently obtained from all plants in j 's diet. The preference decreases in the opposite case, where the rewards obtainable by exclusive foraging on plant i are lower than the current rewards uptake level. Note that the terms in equation (4) have been rearranged from previous publications of this model to emphasize the coupling of the four equations through the visitation rates V_{ij} . We use parentheses that include the variables determining each of the functions in the equations to distinguish functions from parameters, but in the text those parentheses are excluded for improved readability. The visitation rate V_{ij} and the rewards extracted per visit f_{ij} can also be modeled by a saturating function following Holling's type II functional response (Holling 1959), as discussed in appendix D (apps. A–D are available online).

The sums in equations (1)–(4) are taken over the sets of A_i and P_j of pollinator species that are capable of vis-

iting plant i and plant species that can be visited by pollinator j , respectively. Those sets are defined by the network structure taken as model input. Finally, the dynamic preferences of equation (4) model adaptive foraging. These preferences are restricted by $\sum_{k \in P_j} \alpha_{kj} = 1$. When adaptive foraging is not considered, foraging preferences are fixed to

$$\alpha_{ij} = \frac{1}{P_j}, \quad (5)$$

where P_j here represents the number of plant species visited by pollinator species j .

Niche Theory for Plant-Pollinator Dynamics

Niche is a central concept in ecology, significantly clarified and refined over the past 50 years (MacArthur 1969, 1970; Tilman 1982; Leibold 1995; Chase and Leibold 2003). We analyze the niche of plant and pollinator species within their mutualistic interactions, assuming all of their other niche variables (e.g., soil nutrients, water, temperature, nesting sites) constant and sufficient for supporting their populations. There are two reasonable choices for the definition of environment space in plant-pollinator systems. First, on short timescales (i.e., within a flowering season; fig. 1a, "Rewards Space"), the plant populations can be regarded as constant and the relevant environmental factors are the floral rewards. Second, on longer timescales (i.e., across several flowering seasons; fig. 1b, "Plant Space"), plant populations represent the axes for the environment space, letting the reward levels implicitly determine the value of each plant population as a food source. Table 2 summarizes both representations in terms of the model parameters. This section explains both representations to provide a broader picture of niche theory applied to plant-pollinator systems, but we obtain our results in rewards space.

The requirement niche of each pollinator species j ($j = 1, 2, \dots, A$) in either rewards or plant space can be encoded by a zero net growth isocline (ZNGI; Tilman 1982; Leibold 1995). The ZNGI is a hypersurface that separates the environmental states where the growth rate is positive from the states where it is negative. Environmental states along the ZNGI support animal reproduction rates that exactly balance mortality rates, leading to constant population sizes. Adaptive foraging allows the ZNGIs in rewards space to dynamically rotate in the direction of the most abundant rewards. The ZNGIs are dynamic in plant space (even in the absence of adaptive foraging) because the contribution each plant makes to the animal growth rate depends on the current reward level.

The impact niche of each pollinator species is represented by an impact vector, which specifies the magnitude

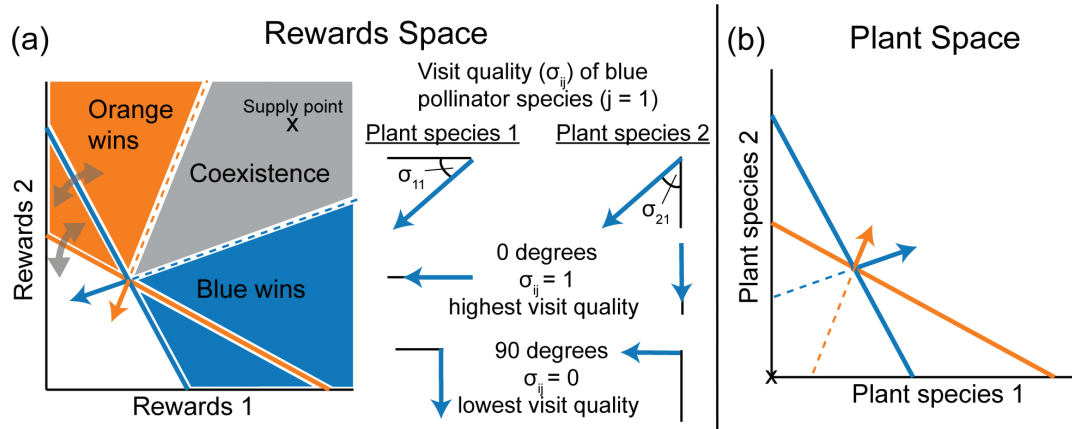


Figure 1: Niche theory for mutualism. *a*, Representation of plant-pollinator system as a standard consumer-resource type model for timescales on which plant populations are approximately constant. Impact vectors and zero net growth isoclines (ZNGIs) are shown for two pollinator species (blue and orange) competing for the rewards of two plant species. Adaptive foraging causes the ZNGIs and impact vectors to rotate in the direction of the most abundant resource, as discussed in detail in appendix A. The angle between the impact vector and a given rewards axis affects the pollinator’s visit quality for the corresponding plant, with zero degrees corresponding to $\sigma_{ij} = 1$ (highest visit quality) and 90 degrees corresponding to $\sigma_{ij} = 0$ (lowest visit quality). See figure D2 for a detailed discussion of the angle-quality relationship. *b*, Representation in terms of plant populations for analysis of longer timescales, where the mutualism becomes visible. The supply point is now located at the origin, and the pollinator impacts are necessary to sustain nonzero plant abundance. The location of the ZNGIs depends on the current nutritional value of each plant species, which is lower for species whose floral rewards are more depleted. The impact vectors (see table 2) depend on both the visit quality and the per capita visit frequency of each pollinator species (σ_{ij} and V_{ij}/a_j of eq. [1], respectively) and encode each pollinator’s contribution to the total number of seedlings in the next generation.

and direction of the environmental change induced by an average individual of the species (Tilman 1982; Leibold 1995). In rewards space, the impact of a pollinator species is the rate at which it depletes the floral rewards, just as in traditional models of resource competition, but its angle takes on a new importance in connection with the visit quality σ_{ij} . A nearly perpendicular impact vector to a given rewards axis means that only a small fraction of the pollinator’s visits are allocated to the corresponding plant, and most of the pollen carried by this pollinator belongs to other plant species. A plant species will eventually be-

come extinct if all of its visits have such low quality (see below). Note that the exact mapping from the angle to the visit quality depends on the foraging strategy, the number of plant species, and plant abundances (see fig. D2; figs. D1–D4 are available online). In plant space, the positive effects of plant-pollinator mutualisms are directly visible in the impact vectors pointing to larger plant population sizes (as opposed to pointing to smaller population sizes in the traditional models of resource competition) and represent the number of successful pollination events caused by each pollinator species.

Table 2: Mapping elements of the model to niche theory concepts

Niche concept	Description	Mathematical expression
Rewards space:		
ZNGI	Reproduction/mortality balance	$\sum_{i \in P_j} c_{ij}(V_{ij}/a_i)f_{ij} = \mu_j^A$
Impact vector	Per capita rewards consumption	$-(V_{ij}/a_i)f_{ij}$
Supply point	Rewards equilibrium without animals	$\beta_i p_i / \phi_i$
Plant space:		
ZNGI	Reproduction/mortality balance	$\sum_{i \in P_j} c_{ij}(V_{ij}/a_i)f_{ij} = \mu_j^A$
Impact vector	Plant production	$\gamma_i e_{ij} \sigma_{ij}(V_{ij}/a_j)$
Supply point	Plant equilibrium without animals	0

Note: ZNGI = zero net growth isocline.

The environment also has its intrinsic dynamics, represented by a supply vector (Tilman 1982; Chase and Leibold 2003). In rewards space, the supply vector points toward the supply point, where the rewards reach equilibrium in the absence of pollinators (as in traditional models of resource competition). However, the supply point itself is determined by the plant populations, which depend on pollination activity for their long-term survival. Extinction of a plant species (e.g., because of low visit quality) causes the supply point to drop to zero along the corresponding rewards axis, leading to a cascade of ecological reorganization and a new equilibrium (see below). In plant space, the equilibrium point in the absence of pollinators is always at the origin, since all plants require pollination services to avoid extinction.

These three quantities (ZNGIs, impact vectors, and supply point) define the conditions for stable coexistence. Pollinator populations reach equilibrium when all of the corresponding ZNGIs pass through the current environmental state. In addition, the combined impact of all pollinator species must exactly cancel the supply for the environment to remain in this state. This total impact is found by multiplying each impact vector by the corresponding population density and then summing the results. Whenever the supply point lies within the cone formed by extending all of the impact vectors backward (fig. 1), a set of population densities can be found with a total impact equal and opposite to the supply. Each potentially stable set of coexisting species is thus represented by an intersection of ZNGIs, and coexistence is achieved whenever the supply point falls within the corresponding coexistence cone.

Conditions for Adaptive Pollinator Coexistence on Shared Rewards

The full equilibrium of the model also requires that adaptive foraging dynamics have reached a steady state. This requirement is satisfied with additional restrictions on the parameter values, which we derive by setting the pollinator growth rate $da_j/dt = 0$ in equation (2) and substituting into the adaptive foraging equation (4). We find the following equilibrium condition:

$$0 = \frac{G_j \alpha_{ij}}{a_j} (c_{ij} V_{ij}^s f_{ij} - \mu_j^A). \quad (6)$$

The term in parentheses is what the growth rate da_j/dt for animal species j would be if it were a specialist on plant species i , with $V_{ij} = V_{ij}^s$ and $\alpha_{ij} = 1$. Equation (6) requires that this term vanish at equilibrium for all plant-animal pairs i, j , where $\alpha_{ij} \neq 0$. Substituting in the expres-

sions for V_{ij} and f_{ij} from the first section of “Methods,” we find the equilibrium rewards abundance R_i^* :

$$R_i^* = \frac{\mu_j^A}{c_{ij} \tau_{ij} b_{ij}}. \quad (7)$$

This result imposes a strict constraint on the animal mortality rates μ_j^A and the reward uptake efficiencies $c_{ij} \tau_{ij} b_{ij}$, requiring that both terms vary in the same way from species to species for all animals that share rewards from the same plant species i (i.e., for all animals with $\alpha_{ij} \neq 0$). Pacciani-Mori et al. (2020) suggests that this required correlation between mortality rates and ingestion rates is consistent with allometric scaling relationships (Yodzis and Innes 1992). However, it is still unknown whether this relationship holds at the species level. Hereafter, we assume that the pollinators’ ZNGIs intersect, acknowledging that the mechanism for coexistence is not present in our model.

Appendix A shows that R_i^* is the rotation center for the ZNGIs, and therefore the shared R_i^* remains the point of intersection for all of the ZNGIs over the entire course of adaptive foraging dynamics.

Using Projections to Analyze High-Dimensional Ecosystems

The graphical analysis described above is easily visualized for environmental spaces with two dimensions. Plant-pollinator networks, however, contain tens to hundreds of plant species. In this full space, the ZNGIs are no longer lines but hypersurfaces of dimension $P - 1$ (fig. 2b, where P is the number of plant species in the network). The intersections among these hypersurfaces determine the points of potential coexistence. We extend our graphical approach to many dimensions and analyze the conditions for coexistence among the species whose ZNGI hypersurfaces intersect by using projections of the coexistence cone onto two-dimensional slices through the full environmental space.

We consider the two-dimensional slice where two of the rewards (or plant) abundances are allowed to vary (gray plane in fig. 2b), while all other abundances are held fixed at the values where the intersection occurs. We then create a diagram like those of figure 1 by drawing the lines where the ZNGIs intersect this slice and projecting the impact vectors and supply point onto this slice (i.e., taking the component parallel to the slice’s surface). The species do not coexist if the projection of the supply point lies outside the projection of the coexistence cone (e.g., fig. 2a–2c) because this can happen only when the supply point lies outside the full coexistence cone. But the supply point may still lie outside the cone (along one of the directions

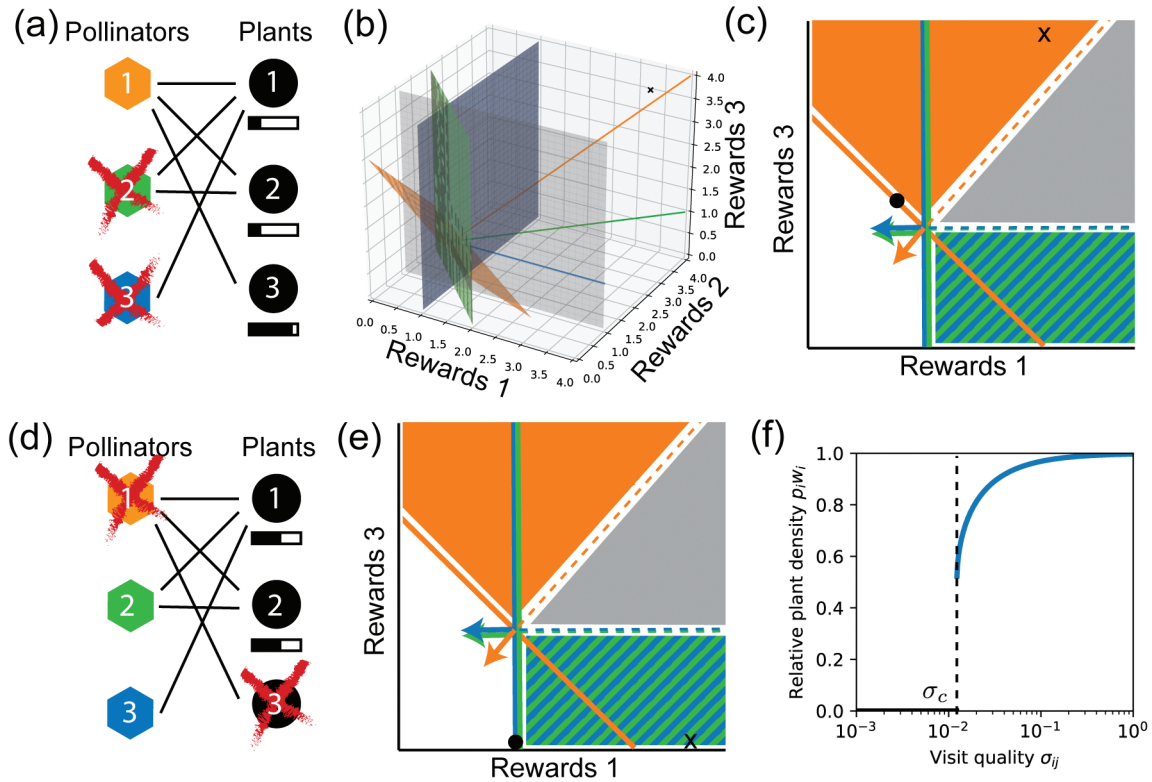


Figure 2: Effects of nestedness without adaptive foraging. *a*, Nested network with three pollinator (polygons) and three plant (circles) species. Shaded bars indicate rewards abundance at the equilibrium point in *c* and *e*, with differences among species exaggerated for clarity. The red X indicates extinction at equilibrium. *b*, Three-dimensional zero net growth isoclines (ZNGIs), impact vectors, and supply point of this network. *c*, ZNGIs and impact vectors projected onto the rewards 1/rewards 3 plane (gray transparent plane in *b*). Pollinator species 2 and 3 have the same projections onto this plane because both visit plant species 1 and none visit plant species 3 (see other projections in fig. D3). The black circle indicates rewards at equilibrium. Specialist pollinators 2 and 3 become extinct because the supply point (black X) falls in the orange zone. *d*, Specialist plant species 3 becomes extinct when the quality of visits it receives is lower than the threshold σ_c of *f*. *e*, Supply point drops to zero along the rewards 3 axis when plant species 3 becomes extinct, which results in the extinction of the generalist pollinator species 1. *f*, Dependence of specialist plant abundance p_i on visit quality σ_{ij} , using equation (C5). The minimal visit quality σ_c required for plant persistence is indicated by the dashed line. Parameters values are taken from Valdovinos et al. (2013), with $\tau_{ij} = 1$, $e_{ij} = 0.8$, $\mu_i^p = 0.008$, $c_{ij} = 0.2$, $\mu_j^A = 0.004$, $b_{ij} = 0.4$, $g_i = 0.4$, $w_i = 1.2$, $\beta_i = 0.2$, and $\phi_{ij} = 0.04$. Plant abundance is measured in units of the plant's carrying capacity $1/w_i$, so that the maximum possible value equals 1.

that has been projected out) even if the projected supply point lies inside the projected coexistence cone. To guarantee coexistence, one must examine all possible two-dimensional projections and ensure that the supply point is inside the cone in every projection (fig. D3).

Results

Effects of Nestedness on Network Dynamics without Adaptive Foraging

Most plant-pollinator networks exhibit a nested structure (definition and citations are provided in the introduction). The implications of nestedness for the stability

of these networks have been a topic of study for over a decade (Bastolla et al. 2009; Allesina and Tang 2012; reviewed in Valdovinos 2019). Valdovinos et al. (2016) provide a more mechanistic framework to evaluate the effects of nestedness on the dynamics of plant-pollinator networks. This section analytically confirms their numerical results when pollinators are fixed foragers (eq. [5]) and provides criteria for plant survival not found by previous work (see the next section for adaptive foragers).

We perform our graphical analysis using two-dimensional slices through the full rewards space of a nested three-plant/three-pollinator-species network (fig. 2*a*), which has sufficient complexity to illustrate all of the relevant projections for arbitrarily large networks. Figure 2*b* shows the

three-dimensional rewards space, with the three colored planes being the ZNGIs of the three pollinator species (derived from table 2). The coexistence cone is the three-sided solid bounded by planes connecting the backward extensions of the impact vectors (colored lines). We project this cone onto the gray transparent plane composed by rewards 1 and 3. This projection is depicted in figure 2c, which shows the asymmetric shape of the coexistence cone, bounded on one side by the impact vectors of the specialist pollinators (green and blue vectors parallel to rewards axis 1) and on the other by the impact vector of the generalist pollinator species (diagonal orange vector). This asymmetric shape is characteristic of nested networks since nestedness increases the diet overlap between specialist and generalist species. This is one of only three possible cone shapes in a two-dimensional projection (see fig. D4) regardless of the full environment dimension.

Valdovinos et al. (2016) show that increasing nestedness increases the extinction of specialist species in networks without adaptive foraging. Our graphical approach explains this result by demonstrating that the asymmetric coexistence cone found most frequently in nested networks favors the extinction of specialist pollinators. To show this, we note that obtaining a supply point in the orange region of figure 2c (where both specialist pollinator species become extinct) requires only that the supply level $\beta_3 p_3 / \phi_3$ of rewards 3 is greater than the supply of rewards 1. This happens half of the time when the plant parameters are randomly chosen (as they were in the previous simulations). But for the supply point to reach the blue and green region, where one or both of the specialist pollinator species persist, the supply of rewards 3 must drop below the ZNGI intersection. This is a much more stringent condition, and in practice it is satisfied only when the specialist plant (here, plant species 3) becomes extinct (fig. 2e).

To elucidate the conditions for plant extinction, we distinguish two drivers of species elimination: competitive exclusion by other plant species for resources other than pollination and failure to receive sufficient pollination. Plant competition is modeled with a Lotka-Volterra-type competition matrix, and standard techniques from coexistence theory can be employed to study this aspect (see app. B). We focus on the second driver by assuming that intraspecific competition is much stronger than interspecific competition, which effectively gives each plant species its own niche. This leaves pollination—particularly visit quality (σ_{ij} ; see “Methods”)—as the sole determinant of plant survival. Specialist plants receive the lowest quality of visits in nested networks because they are visited only by generalist pollinators that carry diluted pollen from many other species. We find the criteria for plant survival by calculating the plant population size p_i as a function of the visit quality σ_{ij} for a perfectly specialist plant (visited by only one

pollinator species). We obtained an exact analytic expression for this relationship (eq. [C5]), which is depicted in figure 2f. This relationship shows that each plant species remains near its maximum abundance ($1/w_i$) as long as the visit quality they receive is above a threshold σ_c , but it suddenly drops to zero when the visit quality drops below this threshold.

Effects of Adaptive Foraging

Adaptive foraging (eq. [4]) rotates the ZNGIs and impact vectors in the direction of the more plentiful floral rewards (see “Methods”). This section explains the consequences of this rotation for species coexistence and provides analytical understanding for the result found by previous simulations showing that adaptive foraging increases the species persistence of nested networks (Valdovinos et al. 2016).

Figure 3 shows how adaptive foraging changes the result illustrated in figure 2a–2c. The supply point lies just outside the coexistence cone, and the equilibrium state with fixed foraging preferences gives plant species 3 a higher equilibrium concentration of floral rewards. This means that the generalist pollinators will begin to focus their foraging efforts on plant species 3, resulting in a rotation of the ZNGI and impact vector to become more like those pollinators specialized on plant species 3 (i.e., a horizontal line and vertical arrow in this visualization). This rotation opens up the coexistence cone until it engulfs the supply point. The resource abundances then relax to the coexistence point (R_1^*, R_2^*, R_3^*), where all plants are equally good food sources, and adaptation stops. This process allows the coexistence of all pollinator species.

Adaptive foraging increases coexistence among plant species in nested networks by causing pollinator species to focus their foraging efforts on more specialist plant species (fig. 3a), increasing the visit quality they receive (see the angle of the orange impact vector becoming more parallel to the rewards 3 axis in the sequence of fig. 3b and 3c). This rotation in ZNGIs in turn decreases the visit quality that the generalist plants receive from the generalist pollinators (see the angle of the orange impact vector becoming more perpendicular to the rewards 1 axis in the sequence of fig. 3b and 3c). The generalist plant species will still persist despite this reduction in visit quality by generalist pollinators because they still receive perfect visit quality from specialist pollinators that visit only them (e.g., pollinator species 3 in fig. 3a) and that cannot shift their foraging effort to other plant species. Overly connected networks (i.e., with many more interactions than the ones found in empirical networks) lack these perfect specialists, and therefore the average quality of visits received by generalist plant

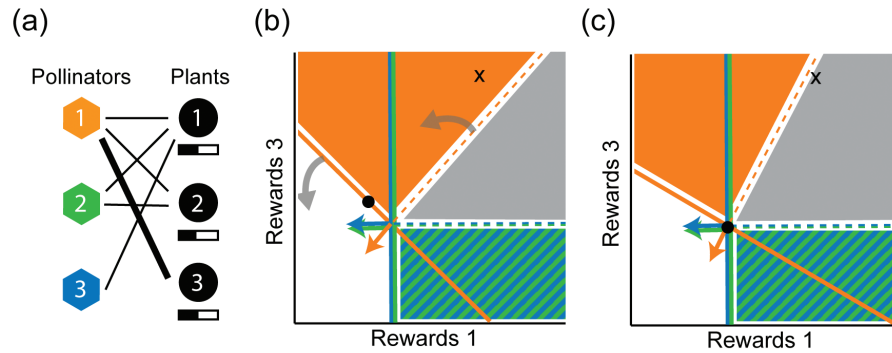


Figure 3: Effects of adaptive foraging. *a*, Adding adaptive foraging to the nested network allows the generalist pollinators to focus their foraging effort on the plant species with more abundant floral rewards (thick line connecting pollinator 1 to plant 3). *b*, Adaptive foraging causes the zero net growth isocline of the generalist pollinator species and its impact vector to rotate counterclockwise (toward the most plentiful rewards 3). The black circle represents the equilibrium state of figure 2c, with more available rewards in plant species 3 than in species 1. *c*, The rotation of the impact vector expands the coexistence cone, making it engulf the supply point, so that all three species coexist in the new equilibrium (black circle). This rotation also reduces the angle between the impact vector and the rewards 3 axis, increasing the quality of visits by the generalist pollinators to these plants while decreasing their quality of visits to the other plant species.

species drops below the threshold σ_c (fig. 2c) and they become extinct, as observed in previous simulations.

Impact of Pollinator Invasions on Native Species

This final section analyzes the consequences of pollinator invasions on species coexistence in networks with adaptive foraging and provides analytical understanding for the results found numerically by Valdovinos et al. (2018). We assume that exotic species come from a different regional pool, with consumption and mortality rates not following the strict relationship imposed on the native species by equation (7). This results in the exotic’s ZNGI not passing through the natives’ common ZNGI intersection (fig. 4b, 4d) but instead intersecting different native ZNGIs at different points. The resulting proliferation of possible coexistence points and cones impede the analysis of high-dimensional systems using the method of projections employed above. Therefore, we focus on a network similar to that in previous sections but with only two (instead of three) plant species.

Exotic pollinators will invade the network whenever the native coexistence point R_i^* falls on the positive growth rate side of the exotic’s ZNGI, regardless of the number of plant species the exotic visits. This corresponds to the case of efficient foragers reported in previous simulations (i.e., with higher foraging efficiency than natives), which were the only exotic pollinators invading the networks studied by Valdovinos et al. (2018). The impact of the invader on native species will depend on how the exotic’s ZNGI alters the coexistence points, which in turn depends on the network structure.

A network structure with native pollinator species visiting only plant species visited by the efficient invader (fig. 4a) has three possible outcomes depending on the position of the supply point: (i) native specialists become extinct when the supply point falls in the invader-generalist coexistence cone (cone 1 in fig. 4b), (ii) generalists become extinct when the supply point falls in the invader-specialist coexistence cone (cone 2 in fig. 4b), and (iii) all native pollinator species become extinct when the supply point falls in the gap between the two coexistence cones (dark region in fig. 4b). This third outcome (illustrated in fig. 4a) happens when all plant species have similar properties (as assumed in previous simulations), which results in a supply point near the diagonal of the rewards space.

A network structure where native pollinators visit plant species not visited by the invader results in a coexistence between the invader and the natives that have access to those alternative resources. For example, the pollinator species 1 coexists with the invader if the invader only interacts with plant species 1. This results in plant species 2 having higher rewards than species 1 at the new coexistence point, which makes pollinator species 1 shift its foraging effort to plant species 2 until it becomes a pure specialist (fig. 4c). Conversely, all three pollinator species coexist as specialists on plant species 1 if the invader interacts only with plant species 2.

This analysis suggests that native pollinators only visiting plants visited by the invader will typically be driven extinct in larger networks because the supply point will most likely fall in the gap between the high-dimensional coexistence cones. But if a pollinator species interacts with at least one plant species not visited by the invader, it will

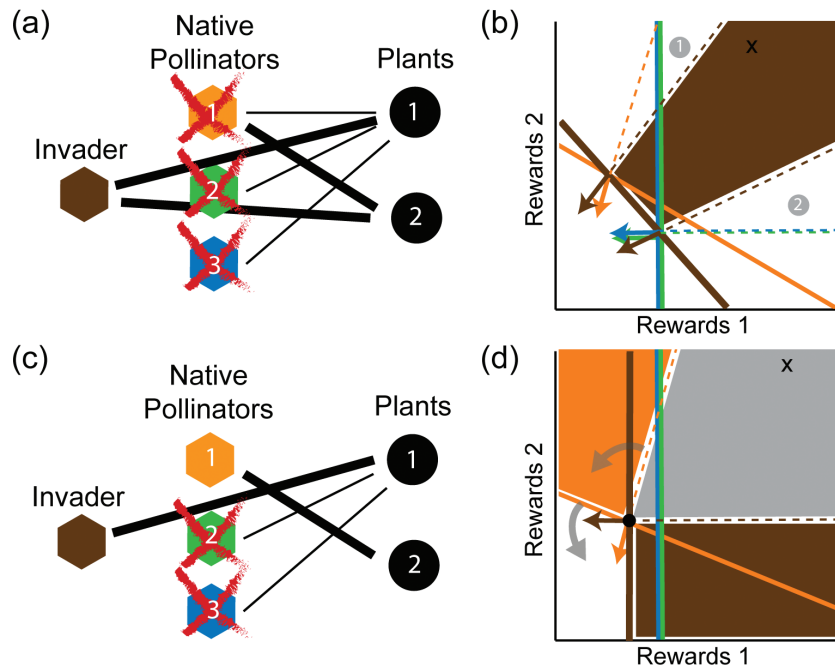


Figure 4: Pollinator invasions. *a*, The brown polygon represents an exotic pollinator species with higher visit efficiency than natives, visiting the two plant species. *b*, If plant species have similar abundances (as in previous simulations), the supply point falls in the gap between the two coexistence cones, and only the invader survives at equilibrium. *c*, The invader does not interact with plant species 2. *d*, The supply point now falls inside the coexistence cone 1 and the invader coexists with pollinator species 1. Adaptive foraging drives the native species to become a pure specialist on plant species 2 (which had more rewards). This results in plant species 2 receiving better visits and in pollinator species 1 reducing its population size. The relative abundances can be estimated from the position of the supply point within the cone. For example, only a small contribution will be required from a pollinator species to achieve perfect cancellation of the supply if one of the other impact vectors points almost directly away from the supply point. The invader's impact vector points in slightly different directions at the two coexistence points. This results from the factor of R_i contained in the f_{ij} term of the impact vector as given in table 2, which biases the vector in the direction of the more abundant reward. Plant extinctions do not occur under these conditions.

survive and transfer all of its foraging effort to these plants. This agrees with previous simulations.

Discussion

Previous studies of species coexistence in plant-pollinator systems mainly consisted of work developing conceptual (e.g., Palmer et al. 2003; Mitchell et al. 2009) and mathematical (e.g., Levin and Anderson 1970; De Mazancourt and Schwartz 2010; Johnson and Bronstein 2019) frameworks for analyzing conditions at which species can coexist as well as reviews of empirical cases showing competition among plant species for pollination services (e.g., Mitchell et al. 2009; Morales and Traveset 2009) and among pollinator species for floral rewards (e.g., Palmer et al. 2003). Contemporary niche theory allows a synthesis of all of this information in one framework and makes quantitative predictions about community dynamics, including species coexistence. We expand this theory by incorporating plant-pollinator systems. Our contributions consist of con-

sidering short- and long-term dynamics of plant-pollinator interactions, depicting the requirement and impact niches of pollinators, and demonstrating the effect of adaptive foraging and network structure on those niches. We applied these advances to the understanding of pollinator invasions. We next explain each of these contributions and contextualize them with previous literature.

Explicit Consideration of Two Timescales: Rewards and Plant Spaces

Explicit consideration of timescales has been recently highlighted as paramount for analyzing ecological systems, especially when evaluating management strategies (Callicott 2002; Hastings 2016) where the time frame of action determines the ecological outcome. This is particularly the case in plant-pollinator systems, whose dynamics can be distinctively divided into at least two timescales: the short-term dynamics occurring within a flowering season and the long-term dynamics occurring across flowering seasons. We

developed our graphical approach for these short- and long-term dynamics by representing the pollinators' niches in rewards and plant spaces, respectively. Rewards space assumes approximately constant plant populations, analyzing the dynamics occurring during a flowering season where plants do not reproduce but produce floral rewards that are depleted by pollinators in a matter of hours or days. Plant space represents the longer timescale at which the quality and quantity of pollinator visits impact plant populations represented on the axes.

The other work we know of that expands contemporary niche theory to mutualisms uses a more classic consumer-resource space (Peay 2016), where niche axes represent resources in the soil used by plant species indistinctly of the timescale. That work shows how the plants' ZNGIs change when the mycorrhizal mutualism is added, but the axes are still resources in the soil, not mutualists. In our work, by contrast, the axes are the abundances of the mutualistic partners themselves (plant space) or the rewards produced by them (rewards space).

Depicting the Pollinators' Requirement and Impact Niches

Analysis of the requirement niches of species sharing resources has long been used to study species coexistence (MacArthur 1970; Tilman 1982; Leibold 1995; Chase and Leibold 2003). Only recently has such analysis been applied to mutualistic systems. Johnson and Bronstein (2019) applied Tilman's resource ratio theory to two pollinator species competing for the rewards provided by one plant species and when an abiotic resource is added. Our results expand this work by extending to networks with larger numbers of plant and pollinator species, where nestedness and adaptive foraging become relevant properties. However, we do not explicitly consider resources or abiotic limitations other than floral rewards that species might require to survive (e.g., nesting sites, water), which represents an important avenue for future work.

We study the pollinators' impact niche corresponding to the change induced on plant and reward abundances. In plant space, the mutualism is directly visible in the impacts, which represent the number of successful pollination events caused by each pollinator, and the impact vectors point in the direction of larger plant population sizes. This space shows a main difference between resource competition in classic consumer-resource and mutualistic systems. Consumers in classic consumer-resource systems can only affect each other negatively through depleting their shared resource, while consumers in mutualistic systems can also benefit each other through benefiting their shared mutualistic partner. In rewards space, the impact of a pollinator species is simply the rate at which it depletes the floral re-

wards, just as in a classic model of resource competition. An important difference, however, is the representation of the visit quality of a particular pollinator species to a particular plant species in terms of the angle between its impact vector and the rewards axis corresponding to the plant species. The analysis of this representation advances another subject that has captured the attention of ecologists for over a century—plant competition for pollination (reviewed in Mitchell et al. 2009). This large body of research has shown that plant species sharing the same pollinator species potentially compete not only for the pollinators' quantity of visits but also for their quality of visits. Our approach provides means for analyzing plant competition for quantity and quality of visits quantitatively and therefore complements previous empirical and conceptual approaches.

Finally, the strict constraint on pollinator parameter values given by equation (7) highlights the intrinsic incompleteness of any model (including ours) that focuses exclusively on plant-pollinator interactions, which are only a subset of the full ecosystem (Hale et al. 2020). Questions on how many pollinator species can coexist or how to prevent competitive exclusion (Gause and Witt 1935; Levin 1970; McGehee and Armstrong 1977) present interesting avenues for further study in models that consider the broader ecological and evolutionary context of plant-pollinator interactions.

Effects of Network Structure and Adaptive Foraging on Species Coexistence

The network structure of plant-pollinator systems influences community dynamics and species coexistence by determining who interacts with whom and which mutualistic partners are shared between any two given species. We analyzed the effects of nestedness on species persistence in these networks by depicting the dynamics occurring in systems where generalist and specialist pollinators share the floral rewards of generalist plants while specialist plants are visited only by generalist pollinators. We provided analytical understanding to results found by previous simulations by showing how nestedness with its increased niche overlap produces an asymmetric coexistence cone that causes the extinction of specialist species.

We demonstrated that adaptive foraging rotates the pollinators' ZNGIs and impact vectors toward the most abundant rewards, promoting pollinator coexistence in nested networks through niche partitioning and plant coexistence through the increased visit quality to specialist plants. We anticipate that our graphical representation of adaptive foraging can be applied to other types of consumer-resource systems, such as food webs, where the effects of adaptive foraging have been extensively studied theoretically (reviewed in Valdovinos et al. 2010). For

example, Kondoh (2003) shows how adaptive foraging causes many species to coexist in complex food webs. Key to this result is the “fluctuating short-term selection on trophic links,” which effectively reduces the realized food web connectance. That is, adaptive foraging allows the rare prey to recover by making the consumers effectively specialize on the most abundant prey, which results in the rare prey becoming more abundant and the abundant prey becoming more rare, causing the adaptive consumers to switch their preferences again. This is similar to our result of generalist pollinators becoming effectively specialized on specialist plants with initially higher reward abundance, but it is also different because our plant-pollinator model does not exhibit fluctuations in foraging preferences. This difference is explained by the inherent timescales of rewards and prey dynamics, where the rewards are produced and consumed at the same short timescale while the production of new prey are lagged behind the consumption by predators. We anticipate that our graphical approach will deepen the conceptual unification of theory on mutualistic systems and theory on food webs by providing analytical understanding of species coexistence in consumer-resource systems and incorporating the effects of adaptive foraging and network structure, both critical for the dynamics of those two types of consumer-resource systems.

Conclusion

Our graphical approach promotes the unification of niche and network theories by incorporating network structure and adaptive foraging into the graphical representation of species’ niches. This approach also deepens the synthesis of mutualistic and exploitative interactions within a consumer-resource framework by including both in the graphical representation of pollinators’ niches. This research may promote further development of ecological theory on mutualisms, which is crucial for answering fundamental questions and informing conservation efforts.

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Statement of Authorship

F.S.V. and R.M. conceived and designed the study, conducted the analyses, and wrote the manuscript.

Data and Code Availability

Code producing the figures is available at github.com/robertsviii/niche-mutualism and on Zenodo (Marsland 2020).

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