Predicting invasion success in complex ecological networks

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A central and perhaps insurmountable challenge of invasion ecology is to predict which combinations of species and habitats most effectively promote and prevent biological invasions. Here, we integrate models of network structure and nonlinear population dynamics to search for potential generalities among trophic factors that may drive invasion success and failure. We simulate invasions where 100 different species attempt to invade 150 different food webs with 15–26 species and a wide range (0.06–0.32) of connectance. These simulations yield 11 438 invasion attempts by non-basal species, 47 per cent of which are successful. At the time of introduction, whether or not the invader is a generalist best predicts final invasion success; however, once the invader establishes itself, it is best distinguished from unsuccessful invaders by occupying a lower trophic position and being relatively invulnerable to predation. In general, variables that reflect the interaction between an invading species and its new community, such as generality and trophic position, best predict invasion success; however, for some trophic categories of invaders, fundamental species traits, such as having the centre of the feeding range low on the theoretical niche axis (for non-omnivorous and omnivorous herbivores), or the topology of the food web (for tertiary carnivores), best predict invasion success. Across all invasion scenarios, a discriminant analysis model predicted successful and failed invasions with 76.5 per cent accuracy for properties at the time of introduction or 100 per cent accuracy for properties at the time of establishment. More generally, our results suggest that tackling the challenge of predicting the properties of species and habitats that promote or inhibit invasions from food web perspective may aid ecologists in identifying rules that govern invasions in natural ecosystems.

Keywords: food web; invasibility; generality; nonlinear bioenergetic dynamic model; niche model; species invasions

1. INTRODUCTION

The Global Invasive Species Database (http://www.isssg.org) suggests that there are no natural rules that govern the processes of invasion, which have any real predictive value (Bright 1998) because any ‘generalizations about invaders over too wide a taxonomic range, such as all species, or all insects, or all angiosperms, invariably (lead to) too many exceptions to be useful’ (Williamson 1996). They list propagule pressure, suitability of the habitat and previous success (Williamson 1996) as the only predictors of invasion success, which are of general predictive value. Similarly, Jeschke & Strayer (2006) have suggested that the only consistent predictors of invasion success for mammals, birds and fishes are propagule pressure and human affiliation. Despite this well-founded pessimism, developing better models to predict when and where species will successfully invade is imperative (Enserink 1999).

Here, we describe a successful exploration for general rules that govern invasion success within models of ecological networks that integrate models of food web structure and nonlinear dynamics (Williams & Martinez 2004b; Brose et al. 2005, 2006). The biological foundation of these models is the fundamental requirement of metabolic energy for all life (Brown et al. 2004) and organisms’ different feeding roles in consuming and providing that energy (Pascual & Dunne 2006). Based on this foundation, we...
develop a food web theory of species’ invasions, which describes how each invader’s trophic function is mediated by other species’ trophic activities and the structural topology of the invaded food web. Our hope is that general rules governing invasion success in Nature will emerge from computational explorations of invasions in theory.

Our approach builds on an extensive literature on structural food web models (Cohen et al. 1990; Dunne 2006) from which we select the niche model (Williams & Martinez 2000), an empirically successful stochastic food web model that uses species richness (S) and directed connectance (C) as input parameters to construct food webs (Williams & Martinez 2000, 2008; Dunne et al. 2004; Stouffer et al. 2006). Our work also uses a nonlinear model (Brose et al. 2005; Martinez et al. 2006) that extends earlier work on bioenergetic dynamics (Yodzis & Innes 1992; McCann & Yodzis 1995; McCann & Hastings 1997; McCann et al. 1998) to n species and arbitrary functional responses (Williams & Martinez 2004b). The niche model successfully predicts the network structure of the largest and most complex food webs in the primary literature (Williams & Martinez 2000, 2008; Camacho et al. 2002; Dunne et al. 2004; Stouffer et al. 2007). The bioenergetic model of population dynamics component successfully simulates persistent and non-persistent stable, cyclic and chaotic dynamics (Williams & Martinez 2004b) that are often found in natural systems (Kendall et al. 1998) as well as in experimental microcosms (Fussmann et al. 2000; Shertzer et al. 2002). Invasions are simulated by integrating the structural niche model with the extended model of bioenergetic dynamics to create models of ecological networks with and without an invader. Prediction of invasion success within our simulations is evaluated using fundamental-niche properties of the invasive species that are independent of interactions with its biotic environment, realized-niche properties of the invader, which capture the trophic interactions between the invader and its biotic environment, and food web properties of the biotic environment independent of the invader. This approach allows us to explore many different structural and dynamic aspects of invasion success before and after invasion attempts.

2. MATERIAL AND METHODS

‘Fundamental-niche’ properties of invasive species, hereafter called ‘invaders’, are characterized by three stochastically generated parameters that dictate the location of an invader i on a theoretical niche axis (ni) as well as the centre (ci) and width (ri) of its feeding range (Williams & Martinez 2000; figure 1). The simulated native community is characterized by 17 ‘web’ properties including species richness (S), connectance (C=LS/S) where L is the number of feeding relationships among species, Martinez (1991)) and 15 other network properties that describe the topological and trophic structure of the webs. The relationship between the invader and its newly invaded community is characterized by a five ‘realized’-niche properties, including the generality (Inv-Gen) and vulnerability (Inv-Vul) of the invader (e.g. the invaders normalized prey (generality) and predator (vulnerability) counts), which are determined by the interaction between an invader’s ‘fundamental’-niche properties and the rest of the species’ niches in the food web.

![Figure 1. Diagram of the niche model and the invasion sequence. (a) S (trophic species richness) and C (connectance) are set at the desired values. Each of S species (here S=7, shown by inverted triangles) is assigned a ‘niche value’ (n) drawn uniformly from the interval (0, 1). Species i consumes all species falling in a range (r) that is placed by uniformly drawing the centre of the range (c) from the interval (r/2, n). Thus, in this diagram, species i consumes four species (grey and black triangles) including itself. The size of r is assigned by using a beta function to randomly draw values from the interval (0, 1) whose expected value is 2C and then multiplying that value by n to obtain a web with C that matches the desired C. These rules stochastically assign each invader three fundamental niche values (ni, ri, ci). These values determine the invader’s fundamental niche and, in concert with the fundamental niches of species in the invaded web, determine the realized niche of the invader. Thus, for example, an invader i with a specific ri and ci has higher generality in an invaded web when relatively many species’ ni fit within i’s feeding range than when invading a web with relatively few species’ ni fitting within i’s ri. (b) Example of attempted invasions by two different invaders into the same web. Invader 1 cannot invade because no species fall within its feeding range. Invader 2 can invade as it has prey (five grey triangles and itself) and therefore is allowed. Our invasion simulations were constructed in three steps. The first step specifies the structure of food webs and the fundamental niche of the invaders using the niche model (figure 1; Williams & Martinez 2000). The second step computes the dynamics of a network for 2000 time steps (t=2000) using the structure–dynamic integrated model (Yodzis & Innes 1992; Williams & Martinez 2004b; Brose et al. 2005, 2006) to generate a collection of dynamically persistent webs. The third step adds invaders to persistent webs at t=2000 (time of introduction) and computes the dynamics of both invaded and corresponding uninvaded (control) webs for another 2000 time steps until t=4000 (time of establishment).

(a) The niche model

The niche model (Williams & Martinez 2000, 2008; Dunne et al. 2004) uses species richness, S, and the connection probability of any pair of species known as directed connectance, C (C=LS/S, Martinez 1991) as input parameters to predict the structure of feeding relationships within a community (figure 1). Each i of S species is assigned a randomly drawn ‘niche value’ (ni) from the interval (1, 0). Each i species is then constrained to consume all prey species within a range of beta-distributed values (ri) whose mean is C and whose randomly chosen centre (ci) is less than the consumer’s niche value. The placement of the feeding range allows up to half a consumer’s range to include species

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with higher niche values than the consumer, thus allowing looping and cannibalism. The consumer is constrained to feed only on those species that fall within its feeding range. While several modifications of the niche model have been described (Cattin et al. 2004; Stouffer et al. 2005; Allesina et al. 2008), we continue to use the original niche model (Williams & Martinez 2000) because it appears to provide the most accurate overall fit to the empirical structure of complex food webs in terms of both the central tendencies and the variability of food web structure (Dunne et al. 2004; Fox 2006; Martinez & Cushing 2006; Stouffer et al. 2006, 2007; Williams & Martinez 2008). We generate invaders by parametrizing the niche model with $C=0.15$ and $S=30$ and then generating 100 species with this model to define each invader’s corresponding $n_i$, $r_i$, and $e_i$ without ever constructing the web that would result from these parameters.

(b) Bioenergetic model of nonlinear food web dynamics

The dynamic model closely follows previous work (Yodzis & Innes 1992; McCann & Yodzis 1995; McCann & Hastings 1997; McCann et al. 1998) but is generalized to $n$ species and arbitrary functional responses (Williams & Martinez 2004b; Brose et al. 2006; Martinez et al. 2006). Extending the earlier notation (Yodzis & Innes 1992) to $n$-species systems, the variation of $B_i$, the biomass of species $i$ over time $t$, is given by

$$\frac{dB_i(t)}{dt} = G_i(B) - x_i B_i(t) + \sum_j (x_{ij} y_{ij} F_{ij}(B) B_j(t)),$$

$$- x_{ij} y_{ij} F_{ij}(B) B_j(t)/(e_{ij}).$$

The first term on the right-hand side $G_i(B)=r_i B_i(t) (1-B_i/(K_i))$ is the net primary production rate of a basal species $i$. $r_i$ is the intrinsic growth rate that is non-zero only for basal species and $K_i$ is the carrying capacity; the second term is metabolic loss where $x_i$ is the mass-specific metabolic rate; the third and fourth terms are gains from resources and losses to consumers, respectively, where $y_{ij}$ is the maximum rate at which species $i$ assimilates species $j$ per unit metabolic rate of species $i$; $e_i (1-\beta_i$. The non-dimensional functional response that may depend on resource and consumer species’ biomasses, $F_{ij}(B)$, gives the realized fraction of the maximum ingestion rate of predator species $i$ consuming prey species $j$; $e_{ij}$ is the conversion efficiency with which the biomass of species $j$ lost due to consumption by species $i$ is converted into the biomass of species $i$. Dividing the last term by $e_{ij}$ converts the biomass assimilated by consumer $i$ into biomass lost by resource $j$. The parameters in these equations have been estimated from empirical measurements (Yodzis & Innes 1992; Brose et al. 2005, 2006). We simplified the dynamic model by assigning the following fixed parameter values: $K_i=1; r_i=1; x_i=0.5; y_{ij} = 6; e_{ij} = 1$; and $B_{0ij} = 0.5$. Previous analyses using this integrated structural-dynamic approach (Martinez et al. 2006) have shown that simulations that draw these parameters from normal distributions with specified means and standard deviations ($e_{ij}>1$ not allowed) gave similar results to fixed parameter simulations.

While a wide variety of functional responses have been proposed in the literature (Holling 1959a; Hassell & Varley 1969; Beddington 1975; DeAngelis et al. 1975; Skalski & Gilliam 2001), our model uses a modified “type II.2” functional response that is close to a type II response while being intermediate between type II and III functional responses (Holling 1959b) and providing much of the stability of type III response (Williams & Martinez 2004b). This response models consumption of resource $j$ by consumer $i$ as

$$F_{ij}(B) = \frac{B_i}{B_i^{1/3}} + \frac{B_j}{B_j^{1/3}}.$$

where $B_i$ is the half saturation density and $q=0.2$ (equation (2.2) is a type III response when $q=1$). A consumer consumes all available resources at a rate equal to its maximum consumption rate times the functional response. The amount that each resource loses to that consumer is equal to the resource’s density divided by the sum of all the densities of the consumer’s resources times the consumer’s rate of consumption. This allows a generalist to consume at its maximal rate even if only one of its prey species has high biomass (Williams 2008). Such biologically reasonable consumption rates are much higher than usual. Instead, most commonly used functional responses limit generalists to consuming any one resource species to no more than their total maximum consumption rate divided by the number of the consumer’s prey species no matter how abundant that one prey species is (e.g. Martinez et al. 2006; Rooney et al. 2006, see Williams 2008).

(c) Generating persistent webs

We constructed three sets of 250 niche model webs with $S=30$ and $C=0.05, 0.15$ or 0.30 (total 750 webs) as input parameters. Web structure is biologically implausible energy flow patterns, such as loops with no external energy source (e.g. cannibals with no other food source), were excluded from further consideration. This yielded 171, 162 and 147 webs, respectively, for each of the three categories of $C$ (0.05, 0.15 and 0.30). We then assigned each species a uniformly random biomass between 0.5 and 1 at $t=0$ and simulated the dynamics of these webs until $t=2000$. Species were eliminated from the web if their abundance decreased below our extinction threshold of $10^{-10}$. Mean species richness (s.d.) of the webs at $t=2000$ is 22 (3.66), 15 (4.44) and 10 (4.70) for each group of webs among $C=0.05, 0.15$ and 0.30 webs, respectively. Fifty webs with $S \geq 15$ were randomly selected and labelled ($v_j$ from 1 to 50 for each of the three $C$ categories (0.05, 0.15 and 0.30) resulting in a total of 150 persistent webs that were used for further simulations.

(d) Invasion simulations

Each invasion was simulated by adding one invader with a uniformly random initial biomass ($Inv-Bio$) between 0.5 and 1 to a persistent food web at $t=2000$. Invaded webs were discarded if the invader was trophically unconnected to the invaded web or if the invasion caused biologically implausible energy flows such as loops discussed above. Acceptable invaded webs and the same webs lacking invaders continued to be dynamically simulated until $t=4000$, employing the same extinction threshold applied during $t<2000$. The webs without invaders will be used in future work exploring the effects of invasions on webs and will not be considered further here. We repeated this procedure for each of 150 persistent webs with the 100 model invaders yielding a total of 15 000 invasion scenarios. This set of 15 000 invasion scenarios yielded 11 483 invasions after eliminating those with biologically implausible food web structures (196 cases) and those where the invader was a basal species (3231 cases). Invasions by basal species were eliminated because the constant growth term of basal species’ within our model makes it almost impossible for basal species to go extinct and usefuly inform our study of differential invasion success. All simulations are programmed in Java v. 1.4.1 and run using fifty 64 bit processors operating within a large cluster running...
(e) Food web and invader properties
We use 17 properties to describe food web structure. One property is simply the number of species within the food web (S). Two other properties are standard measures of food web trophic interaction richness (Martinez 1991, 1992): links per species (L/S) also referred to as link density; and directed connectance \( C = L/S^2 \) that equals the proportion of all possible trophic links that are actually realized. Five more properties indicate the fraction of the following types of species in a food web: top (%T; species that have resource species but lack any consumer species, Briand & Cohen (1984)); intermediate (%I, species that have both resource and consumer species, Briand & Cohen (1984)); basal species (%B, species that have consumer species but lack resources species, e.g. plants, Briand & Cohen 1984); cannibals (%C, species that eat themselves, Williams & Martinez (2000)); and omnivores (%Omn, species that eat species at different trophic levels (TLs), Williams & Martinez (2000)).

Four more properties indicate the fraction of links between the following types of species (Cohen & Briand 1984): top–basal (%T–B); top–intermediate (%T–I); intermediate–intermediate (%I–I); and intermediate–basal (%I–B). Mean TL was calculated in two ways (Williams & Martinez 2004a). One is the mean of the shortest chain length (SCL) between each species and basal species (MSCL). The other is the mean of each species’ prey-averaged TL (MTL) that is the mean of each species’ resource species plus 1. Two additional properties are the standard deviation of mean generality (GenSD) and vulnerability (VulSD) among species which quantify the variability of species’ normalized predator and prey counts, respectively (Schoener 1989; Williams & Martinez 2000).

Trophic similarity of a pair of species is the number of predators and prey shared in common divided by the pair’s total number of predators and prey (Martinez 1991, 1992), Solow & Beet 1998; Yodzis & Winemiller 1999). The average of all species’ similarity indices is the property called mean similarity (MSim). The final web property is a ‘small-world’ (Watts & Strogatz 1998) property called the clustering coefficient (CC), which is the mean fraction of species pairs connected to the same species and each other (Camacho et al. 2002; Dunne et al. 2002; Montoya & Solé 2002; Williams et al. 2002a,b).

Nine properties of the invader were evaluated. Three properties describe the fixed properties \((n_h, r, c_i; \text{figure 1})\) that characterize the invader’s fundamental niche and five properties describe the realized niche of the invader within the invaded food web. The fundamental and realized niches of the invader differ because the invader’s realized niche depends on the niches of other species in the invaded web whereas the fundamental niche does not. The invader’s realized-niche properties of generality \((Inv–Gen)\), vulnerability \((Inv–Vul)\), and omnivory \((Inv–Omn)\) are defined as, respectively, the invaders normalized prey (generality) and predator (vulnerability) counts (Williams & Martinez 2000), and the standard deviation of the prey-averaged TL of the invader’s prey (Williams & Martinez 2004a). Generality and vulnerability are normalized, respectively, by dividing the number of prey and predator species by the number of the species in the food web (S). Prey-averaged TL \((Inv–TL)\) equals 1 plus the mean TL of all of the invader’s resource species (Williams & Martinez 2004a). Short-chain TL \((Inv–SCL)\) equals 1 plus the SCL from the consumer taxon to a basal taxon (Williams & Martinez 2004a).

The final invader property was the initial biomass of the invader \((Inv–Bio)\), which was randomly assigned from a uniform distribution between 0.5 and 1. Initial invader biomass was, on average, 15 times higher than the equilibrium biomass of single species in the webs at \(t=2000\), but was equivalent to the average initial biomass of each species in the webs (0.75) at \(t=0\). Owing to the restricted range of values of invader biomass chosen, we expected minimal effects of invader biomass on invasion success.

Properties of the webs and invaders (when present) were calculated at \(t=0\) (initial conditions), \(t=2000\) (introduction) and \(t=4000\) (establishment; electronic supplementary material). All 3321 invasions by basal species \((SCL=0)\) were successful and, as mentioned above, removed from our results prior to the analyses described below. Unsuccessful invaders persisted an average of 199 time steps (s.d. = 195) ranging from \(t=2030\) to 3974.

(f) Statistical analysis
We used discriminant analysis (DA) to statistically model invasion success. DA models responses of a categorical dependent variable, in this case invasion success or failure, in terms of a series of other properties that might influence such success. We define success and failure as the persistence or not (failure) of the invader maintaining more than \(10^{-10}\) biomass for the 2000 time steps between \(t=2000\) and \(t=4000\). DA produces a set of coefficients that define the single linear combination of properties (the discriminant function), which best differentiates the successful from unsuccessful invasions. We used forward stepwise selection to choose the property that best discriminates between the successful and unsuccessful invasions at each step of the DA. This is akin to linearly regressing invasion success against each property and choosing the property whose regression explains the most variation in success. The residuals from that chosen regression are then evaluated in the next step to similarly choose the property that explains most of the variation of the residuals (i.e. yields the smallest Wilk’s lambda) or, in other words, explains most of the variation unexplained by the previously chosen property. The procedure is stopped when none of the properties explain 5 per cent or more of the remaining variation.

DA was used to assess both which variables best predict invasion success and how accurate that prediction is given 27 potential properties of both invader and web observed at two different times: \(t=2000\) (time of introduction) and \(t=4000\) (time of establishment). The time of establishment model was based on the web and invader properties at \(t=4000\) except for failed invaders in which case their properties are those at the time step that they dipped below the extinction threshold. Biomass of the invader was excluded as it would trivially define the categories. All 3321 invasions by basal species \((SCL=0)\) were successful and, as mentioned above, removed from our results prior to the analyses described below. Unsuccessful invaders persisted an average of 199 time steps (s.d. = 195) ranging from \(t=2030\) to 3974.
We tested the DA model's ability to predict invasion success by comparing discriminant functions from DAs of a randomly chosen 75 per cent of cases with discriminate functions from DAs of the remaining 25 per cent of cases. Discriminant functions estimated on one set of simulation results are expected to more poorly predict invasion success in a different set of results because the latter set plays no role in estimating the functions. We report results for DA models of the 75 per cent of cases unless specified otherwise as the cross-validation results that refer to the results from the 25 per cent of cases.

3. RESULTS

(a) Predicting invasion success from properties at time of introduction

Among all 11,483 analysed invasions, 47 per cent were successful (i.e. survive until \( t = 4000 \); figure 2a). DA revealed that the properties at the time of introduction, which predict invasion success most accurately, are high generality (\( \text{Inv-Gen} \)), having the centre of the feeding range lower on the niche axis (\( c_z \)), and a SCL leading to basal resources. Successful invaders also tended to invade webs with greater numbers of species (\( S \)) and webs with high variability of generality (\( \text{GenSD} \)) indicating that the webs are composed of both highly specialized and highly generalized species. A discriminant function using 15 out of the 27 properties retained in the stepwise DA discriminated between failed and successful invaders with 76.5 per cent accuracy (75% on cross-validation) and showed that invasion success was most accurately predicted by a relatively complex model that includes 8 out of the 18 web properties, 3 out of the 3 fundamental-niche properties of the invader and 4 out of the 5 realized-niche properties of the invader (table 1).

More insight into these general patterns comes from exploring in more detail how invasion success is affected by the highly variable properties of the invaded webs and the invader itself. Food web properties varied among the webs’ initial values \( (t = 0) \), their values at persistence \( (t = 2000 \); Martinez et al. 2006) when the webs were invaded and their values at \( t = 4000 \) when the invasion scenarios end (electronic supplementary material). Realized-niche properties of the invader also varied between their introduction at \( t = 2000 \) and persisting or not until \( t = 4000 \) (electronic supplementary material). We focus our results on the following questions: (i) which and how well properties of webs and invaders at the time of introduction (i.e. \( t = 2000 \)) predict invasion success, (ii) which are the differences between such predictions and those based on properties at the time of establishment \( (t = 4000) \), and (iii) do our results systematically vary according to trophic category?

(i) DA model for invasion success within \( C \) categories

Invasion success is over twice as high in webs with low \( C \) compared with those with high \( C \) decreasing from 70 per cent in the \( C = 0.05 \) webs to 42 per cent in the \( C = 0.30 \) webs (figure 2a). Invader generality (\( \text{Inv-Gen} \)) most accurately predicted invasion success in all three \( C \) categories. As \( C \) increased, DA retained web properties less often and the invaders’ fundamental- as opposed to realized-niche properties better predicted invasion success (table 1). The discriminant function using 14 out of the 27 properties retained for \( C = 0.05 \) webs discriminated between failed and successful invaders with 72.5 per cent accuracy (70% on cross-validation). As connectance increased, the DA retained fewer variables dropping sharply to eight properties predicting invasion success with 74 per cent accuracy (76.5% on cross-validation) in the \( C = 0.15 \) webs, to only six properties predicting success with 68 per cent accuracy (65.5% on cross-validation) in the \( C = 0.30 \) webs. This increase in model parsimony occurs concurrent with the increased importance of fundamental-niche properties (5, 7 and 29% of variability explained among \( C = 0.05 \), 0.15 and 0.30 webs, respectively) and decreased importance of web properties (18, 14 and 3% among \( C = 0.05 \), 0.15 and 0.30 webs, respectively). Most importantly, the fundamental-niche property, \( c_z \), becomes a more accurate predictor of invasion success than the realized-niche property, invader generality (\( \text{Inv-Gen} \)) in \( C = 0.30 \) webs. Despite these differences, properties relating to high invader generality remain the most accurate predictors of invasion success within all \( C \) categories.

Figure 2. Invasion success. (a) Fraction of the total number of successful invasions across all \( C \) categories (black bars, all \( C \)) and in \( C = 0.05 \) (white bars), 0.15 (light grey bars) and 0.30 (dark grey bars) webs. (b) Fraction of successful invasions according to trophic category across all \( C \) categories (black bars, all \( C \)) and in \( C = 0.05 \) (white bars), 0.15 (light grey bars) and 0.30 (dark grey bars) webs. Trophic categories are defined according to the SCL and TL of the invader: herbivore (SCL = 1, TL < 2), herbivorous omnivore (SCL = 1, TL < 2), secondary consumer (SCL = 2) and tertiary consumer (SCL = 3). Fractions of successful and failed invasions are shown for \( C = 0.05 \), 0.15 and 0.3.
Table 1. DA by $C$ at $t=2000$ among all simulations combined (all) as well as within each $C$ category separately for web properties and fundamental and realized-niche properties of the invader for predicting invasion success showing effect size ($F$), p-value ($p$) and the standardized function coefficients (coeff.). (Blank spaces indicate that the variable was not selected in forward stepwise DA. Positive coefficients are associated with traits that are associated with invasion failure while negative values are associated with invasion success. $F_t$ failed invasion; $S$, successful invasion.)

| web properties | all $C$ | $C=0.05$ | $C=0.15$ | $C=0.3$ |
|----------------|---------|----------|----------|---------|
| $S$            | 90.127  | $<0.001$ | $-0.023$ | 33.009  | $<0.001$ | $-0.018$ |
| $C$            | 5.936   | 0.015    | $-0.672$ |         |          |          |
| $L/S$          | 20.811  | $<0.001$ | 0.065    | 18.658  | $<0.001$ | $-0.119$ | 4.395   | 0.036   | $-0.018$ |
| $O$            | 29.727  | $<0.001$ | 0.282    | 33.583  | $<0.001$ | 0.531    |          |         |          |
| $I$            | 46.763  | $<0.001$ | 0.546    |          |          |          |          |         |          |
| $C$            | 25.633  | $<0.001$ | $-0.382$ |          |          |          |          |         |          |
| $T$            | 5.683   | 0.017    | 0.381    |          |          |          |          |         |          |
| $I-B$          | 44.377  | $<0.001$ | 0.327    | 10.173  | $<0.001$ | 0.208    |          |         |          |
| $I-I$          | 70.305  | $<0.001$ | $-0.432$ |          |          |          |          |         |          |
| $T-B$          | 88.399  | $<0.001$ | 0.281    |          |          |          |          |         |          |
| $O$            | 4.724   | 0.030    | 0.173    |          |          |          |          |         |          |
| $MSL$          | 88.909  | $<0.001$ | $-0.357$ |          |          |          |          |         |          |
| $Mim$          | 84.989  | $<0.001$ | 4.676    |          |          |          |          |         |          |
| $CC$           | 4.933   | 0.026    | $-0.163$ |          |          |          |          |         |          |

fundamental-niche properties

| $n_i$          | 7.738   | 0.005    | $-0.067$ | 11.323  | $<0.001$ | $-0.138$ |          |         |          |
| $r_i$          | 76.668  | $<0.001$ | $-0.449$ | 5.847   | 0.016    | $-0.237$ | 15.318  | $<0.001$ | $-0.363$ | 6.505   | 0.011   | $-0.227$ |
| $c_i$          | 180.891 | $<0.001$ | 0.392    | 32.621  | $<0.001$ | 0.279    | 116.256 | $<0.001$ | 0.336    | 294.019 | $<0.001$ | 0.569    |

realized-niche properties

| $Int-Gen$      | 557.810 | $<0.001$ | $-0.127$ | 188.430 | $<0.001$ | $-0.099$ | 279.224 | $<0.001$ | $-0.245$ | 146.874 | $<0.001$ | $-0.294$ |
| $Int-Vul$      | 5.945   | 0.015    | 0.028    |          |          |          |          |          |          |          |          |          |
| $Int-TL$       | 9.397   | 0.002    | $-0.032$ | 10.272  | 0.001    | $-0.078$ |          |          |          |          |          |          |
| $Int-SCL$      | 123.971 | 0.000    | 0.176    | 60.903  | $<0.001$ | 0.235    | 137.198 | $<0.001$ | 0.198    | 20.257  | $<0.001$ | 0.074    |
| $Int-OMN$      | 30.258  | $<0.001$ | $-0.128$ | 10.229  | $<0.001$ | $-0.135$ |          |          |          |          |          |          |

| eigenvalue     | 0.548   | 0.408    | 0.452    | 0.310   |          |          |          |          |          |          |          |          |
| canonical R    | 0.595   | 0.538    | 0.558    | 0.487   |          |          |          |          |          |          |          |          |
| Wilk’s lambda  | 0.646   | 0.710    | 0.689    | 0.763   |          |          |          |          |          |          |          |          |

$p$ values:

| $F$            | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ | $<0.001$ |

(iv) DA models for invaders within trophic-level categories
Omnivorous herbivores were the most successful trophic category of invader with invasion success occurring in 64 per cent of cases followed by herbivores (47%), secondary consumers (19%) and tertiary consumers (14%; figure 2b). Both successful non-omnivorous (herbivores that only eat basal species ($SCL=1, TL=2$)) and omnivorous herbivores ($SCL=1, TL>2$) tend to invade webs with more species ($S$), have a fundamental-niche constraining them to eat low on the niche dimension ($c_i$) and a realized-niche that includes more resource species (higher $Int-Gen$). Successful non-omnivorous herbivores differ slightly from omnivorous herbivores by tending to invade webs with fewer numbers of links per species ($L/S$) whereas successful omnivorous herbivores tend to have larger fundamental feeding ranges ($r_i$). The DA retained 15 out of the 27 properties and predicted invasion success by non-omnivorous herbivores with 72 per cent accuracy (73.5% cross-validation; table 2) and, for omnivorous herbivores, retained 14 properties that predicted success with 72.5 per cent accuracy (75% cross-validation).

Unlike herbivores, invasion success of primary carnivores ($SCL=2$) is most accurately predicted by a single realized-niche property: high invader generality ($Int-Gen$). However, the DA also retained 11 out of the 18 web properties and a fundamental-
Table 2. DA by SCL at t=2000 with web properties and fundamental and realized-niche properties of the invader for predicting invasion success showing effect size (F), p-value (p) and the standardized function coefficients (coefficient). (Blank spaces indicate that the variable was not selected in forward stepwise DA. Positive coefficients are associated with traits that are associated with invasion failure while negative values are associated with invasion success. F, failed invasion; S, successful invasion.)

|                  | SCL=1 | TL=2 |         | SCL=1 | TL<2 | SCL=2 | SCL=3 |
|------------------|-------|------|---------|-------|------|-------|-------|
|                  |       |      |         |       |      |       |       |
| web properties   |       |      |         |       |      |       |       |
| S                | 130.129 | <0.001 | -0.032 | 168.026 | <0.001 | -0.026 | 15.751 | <0.001 | -0.010 | 3.986 | 0.049 | 0.022 |
| L/S              | 51.618 | <0.001 | 0.116   | 166.535 | <0.001 | 0.085   | 86.498 | <0.001 | -1.200 |       |       |       |
| %B               | 29.308 | <0.001 | 0.387   | 6.941  | 0.008 | 0.208  | 11.662 | 0.001  | 0.263  |       |       |       |
| %O               | 16.231 | <0.001 | 0.320   | 6.305  | 0.012 | 0.236  | 24.581 | <0.001 | -1.598 |       |       |       |
| %I               | 12.345 | <0.001 | -0.528  | 21.876 | <0.001 | -0.852 | 38.087 | <0.001 | -5.140 |       |       |       |
| %T               | 7.048  | 0.008 | 0.203   | 7.404  | 0.007 | 0.405  | 68.060 | <0.001 | 7.545  |       |       |       |
| %I–T             | 50.203 | <0.001 | -0.564  | 56.458 | <0.001 | -0.699 | 8.518  | 0.004  | -0.379 |       |       |       |
| %T–I             | 14.546 | <0.001 | -0.287  | 17.137 | <0.001 | -0.644 | 19.660 | <0.001 | -1.560 |       |       |       |
| GenSD            | 50.742 | <0.001 | 0.358   | 50.551 | <0.001 | 0.338   | 20.548 | <0.001 | 0.230  | 13.994 | <0.001 | 1.793 |
| VulSD            | 6.265  | 0.012 | -0.120  | 6.644  | 0.011 | 0.376  | 91.117 | <0.001 | -0.344 |       |       |       |
| MSL              | 38.839 | <0.001 | 0.419   | 18.043 | <0.001 | 0.376   | 91.117 | <0.001 | -0.344 |       |       |       |
| MTI              | 19.211 | <0.001 | -0.162  | 42.904 | <0.001 | -0.263 | 91.117 | <0.001 | -0.344 |       |       |       |
| Mism             | 14.857 | <0.001 | -1.518  | 15.553 | <0.001 | -0.211 |       |       |       |       |       |       |
| fundamental-niche properties |      |       |         |       |      |       |       |
| n_i              | 8.208  | 0.004 | -0.078  | 6.878  | 0.009 | -0.080 |       |       |       |       |       |       |
| r_i              | 116.813 | <0.001 | -0.671  | 126.561 | <0.001 | -0.747 |       |       |       |       |       |       |
| c_i              | 374.799 | <0.001 | 0.599   | 266.935 | <0.001 | 0.555   | 50.356 | <0.001 | -0.211 |       |       |       |
| realized-niche properties |      |       |         |       |      |       |       |
| Inv–Gen          | 291.168 | <0.001 | -0.108  | 157.167 | <0.001 | -0.088 | 846.182 | <0.001 | -0.298 |       |       |       |
| Inv–Vul          | 4.631  | 0.031 | 0.029   |       |       |       |       |       |       |       |       |       |
| Inv–OMN          | 22.635 | <0.001 | -0.129  |       |       |       |       |       |       |       |       |       |
| eigenvalue       | 0.398  |       | 0.452   | 0.622  |       | 1.066  |       |       |       |       |       |       |
| canonical R      | 0.534  |       | 0.558   | 0.619  |       | 0.718  |       |       |       |       |       |       |
| Wilk’s lambda    | 0.715  |       | 0.689   | 0.617  |       | 0.484  |       |       |       |       |       |       |
| P                | <0.001 | <0.001 | <0.001  | <0.001 | <0.001 | <0.001 |       |       |       |       |       |       |

| F | S | F | S | F | S | F | S |
|---|---|---|---|---|---|---|---|
| 78 | 68 | 60 | 85 | 95 | 55 | 98 | 80 |
| 78 | 67 | 65 | 85 | 96 | 47 | 92 | 100 |

Niche property, c_i, as significant, albeit very minor, predictors of invasion success. The DA for primary carnivores discriminated between failed and successful invasions with 75 per cent accuracy (71.5 per cent cross-validation).

Unlike lower trophic categories for which success is most accurately predicted by realized and fundamental-niche properties, invasion success of secondary carnivores (SCL=3) is most accurately predicted by web properties. Specifically, successful secondary carnivores invaded webs with few links between other top species and intermediate species (%T–I), webs that had few top species (%T) and cannibals (%C), and webs that have species with similar numbers of consumers (low VulSD). The DA for secondary carnivores discriminated between failed and successful invasions with 89 per cent accuracy (95 per cent cross-validation).

(iii) Invader biomass

Inv–Bio at t=2000 was unrelated to invasion success. Among successful invasions, both mean biomass of the species in the web and final Inv–Bio was the highest in C=0.05. Inv–Bio of these successful invaders at t=4000 averaged 0.01 in the C=0.30 webs, 0.021 in the C=0.15 webs and 0.062 in the C=0.05 webs (electronic supplementary material).

Final mean invader biomass was lower than final mean biomass for each species in the web for all C categories.
Table 3. DA by $C$ at $t=4000$ across all simulations combined (all) as well as within each $C$ category separately for web properties and fundamental and realized-niche properties of the invader for predicting invasion success showing effect size ($F$), p-value ($p$) and the standardized function coefficients (coeff.). (Blank spaces indicate that the variable was not selected in forward stepwise DA. Positive coefficients are associated with traits that are associated with invasion failure while negative values are associated with invasion success. $F$, failed invasion; $S$, successful invasion.)

| Web properties | $F$ | $p$-value | coeff. | $F$ | $p$-value | coeff. | $F$ | $p$-value | coeff. | $F$ | $p$-value | coeff. |
|----------------|-----|----------|--------|-----|----------|--------|-----|----------|--------|-----|----------|--------|
| $S$            | 3.623 | 0.057 | -0.001 | 15.488 |<0.001 | 0.017 | 8.263 | 0.004 | 0.658 | 11.690 |<0.001 | 0.046 |
| $C$            | 81.368 |<0.001 | -0.433 |          |        |        |       |        |        |       |        |        |
| L/S            | 48.560 |<0.001 | -0.060 | 189.273 |<0.001 | 0.687 | 188.004 |<0.001 | 0.573 |          |        |        |
| $P$            | 446.791 | 0.001 | 0.573 | 25.128 | 0.001 | 0.194 | 41.896 |<0.001 | 0.399 |          |        |        |
| $Wilk$         | 29.164 | 0.001 | 0.082 | 1.102 | 0.292 | 0.130 | 6.899  | 0.009 | 0.033 |          |        |        |
| $Inv-OMN$      | 15.549 |<0.001 | -0.134 |          |        |        | 4.080 | 0.043 | 0.012 |          |        |        |
| $Inv-SCL$      | 7.542  | 0.006 | 0.032 |          |        |        | 7.230 | 0.007 | 0.039 |          |        |        |
| $Inv-TL$       | 474.830 |<0.001 | -0.150 | 162.422 |<0.001 | 0.430 | 11.581 | 0.001 | 0.073 | 174.095 |<0.001 | 0.134 |
| $MTL$          | 18.646 |<0.001 | 0.092 | 124.950 | 0.001 | 0.443 | 822.751 |<0.001 | 0.304 | 1004.292 |<0.001 | 0.271 |
| $Msim$         | 55.493 |<0.001 | 0.449 | 48.353 |<0.001 | 0.843 |          |        |        |          |        |        |
| $CC$           | 7.542  | 0.006 | 0.032 |          |        |        | 7.230 | 0.007 | 0.039 |          |        |        |

| Fundamental-niche properties | $n_i$ | 474.830 |<0.001 | -0.150 | 105.600 |<0.001 | -0.128 | 290.156 |<0.001 | -0.189 | 444.991 |<0.001 | -0.191 |
|                             | $r_i$ | 23.574 |<0.001 | 0.061 | 19.058 |<0.001 | 0.122 | 26.552 |<0.001 | 0.103 | 17.611 |<0.001 | 0.063 |
|                             | $c_i$ | 3069.823 |<0.001 | 0.370 | 1249.502 |<0.001 | 0.443 | 822.751 |<0.001 | 0.304 | 1004.292 |<0.001 | 0.271 |

| Realized-niche properties | $Inv-Gen$ | 1477.750 |<0.001 | -0.048 | 493.735 |<0.001 | -0.044 | 442.934 |<0.001 | -0.080 | 800.336 |<0.001 | -0.188 |
|                          | $Inv-Vid$ | 1740.905 |<0.001 | -0.119 | 138.880 |<0.001 | -0.053 | 1153.384 |<0.001 | -0.167 | 2583.355 |<0.001 | -0.254 |
|                          | $Inv-TL$ | 2687.111 |<0.001 | -0.183 | 14938.383 |<0.001 | -0.313 | 499.277 |<0.001 | -0.130 | 62.427 |<0.001 | -0.044 |
|                          | $Inv-SCL$ | 799.516 |<0.001 | -0.203 | 547.200 |<0.001 | -0.255 | 984.770 |<0.001 | -0.383 |          |        |        |
|                          | $Inv-OMN$ | 9.594  | 0.002 | -0.293 | 133.757 |<0.001 | 0.136 | 822.751 |<0.001 | 0.304 | 1004.292 |<0.001 | 0.271 |
| $F$                      | $S$     | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    |
| $F$                      | $S$     | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    | 100    |

(b) Predicting invasion success from properties
at establishment

When the establishment-web and establishment-niche properties of the invasion scenarios are used ($t=4000$) to model invasion success across all $C$ categories, the DA model discriminates between successful and failed invasions with 100 per cent accuracy (cross-validation 100%). Once established, accurate prediction of invasion success is dominated by fundamental and realized-niche properties of the invaders (table 3). Specifically, low $c_i$, $Inv-TL$ and $Inv-Vid$ are the properties that most accurately predict invasion success. However, the most accurate predictors of invasion success also differ within the $C$ categories. In $C=0.05$ webs, successful invaders are most accurately predicted by having a low $TL$ ($Inv-TL$). In $C=0.15$ and 0.30 webs, the most accurate predictor of invasion success is invulnerability to predation ($Inv-Vid$). While web properties remain accurate, albeit weak, predictors of invasion success at $t=4000$, niche properties of the invader accounted for the majority of the predictive accuracy of the models for both among all $C$ categories and within all $C$ categories.

4. DISCUSSION

As indicated by the high and repeatable explanatory abilities of the DA models, invasion success was accurately predicted among our simulations despite highly variable food web structure and invader identity.
This was especially true based on the properties at the end of the simulations ($t=4000$) compared with those based on the properties at the start of the invasions ($t=2000$). At $t=4000$, fundamental-niche properties of the invader, realized-niche properties of the invader and web properties discriminated between successful and unsuccessful invasions with 100 per cent accuracy. However, the corresponding $t=2000$ model was also highly accurate with 76.5 per cent of successful and failed invasions predicted accurately.

To successfully invade a habitat, a species must successfully complete three steps: introduction; establishment; and spread (Jeschke & Strayer 2005). Our network simulations focused on the first and second steps: properties at introduction and those at establishment. At $t=2000$, the fundamental-niche properties of the invader described by $n_z$, $r_z$ and $c_z$ interact with the niche properties of all other species in the web to create a set of realized-niche properties that describe how the invader initially interacts within the web. Final conditions at $t=4000$ represent the combined effects of the invader's fixed fundamental-niche properties, direct interactions with the invader's predators and prey, and indirect interactions with other species and the web as a whole.

The models for $t=2000$ and $4000$ capture complementary information about how invaders succeed in invading complex ecological networks. The most accurate predictors of invasion success at $t=2000$ are having high generality (Inv-Gen), having the centre of their feeding range ($c_z$) low on the niche axis, and being herbivorous ($SCL=1$, $TL\geq2$). However, while trophic category remains a highly accurate predictor of invasion success at $t=4000$, generality is no longer as accurate a predictor of invasion success as vulnerability to predation except in low connectance webs. These results suggest several potential rules about invasion success. Perhaps the most obvious example is that interactions between an invading species and its new community ultimately determine invasion success. More surprising is that, even at time of introduction, it may be possible to relatively accurately predict invasions success from a food web perspective.

Recent reviews of invasion ecology have suggested that models for species invasions must include traits of both the species and the environment, while recognizing that the interaction between an invasive species and other species in community is what ultimately determines invasion success (Lodge 1993; Shea & Chesson 2002; Ruesink 2005). However, there also appear to be fundamental properties associated with successful invaders that are not as dependent on the interaction of an invader with its new community. In our simulations, the most important fundamental property was the location of the centre of a species feeding range ($c_z$) on the theoretical niche axis, which was an accurate predictor of invasion success, particularly in webs with higher connectance. Specifically, at $t=2000$ in $C=0.30$ webs, $c_z$ contributes 59 per cent of the explained variance to the model and, at $t=4000$, $c_z$ contributes 27 per cent of the explained variance to the model across all $C$ categories. While $c_z$ tends more towards intermediate values in webs with higher $C$ in order to keep the larger $r_z$ from extending beyond the niche dimension, it is unclear why eating low on the niche axis is a more accurate predictor of invasion than the corresponding realized-niche properties such as SCL or Inv-TL. Realized-niche properties such as low TL (Inv-TL) were initially expected to account for practically all variability because fundamental-niche properties appear to have no other function with our models other than to define realized niches. Further exploration needs to determine the source of this curious result. Not all properties of species' niches were observed in our models. Unobserved properties including ratios and other functions of observed properties may be discovered the correlate strongly with $c_z$ and better elucidate $c_z$’s role in invasion success.

Invasion success decreases sharply as connectance increases from 70 per cent in the $C=0.05$ webs to 42 per cent in the $C=0.15$ webs to 27 per cent in the $C=0.30$ webs. This suggests that less complex food webs have less crowded niche space with less competition for food resources, which facilitates invasion success. The higher biomass of species in $C=0.05$ webs further supports this interpretation making it an important and intuitively appealing hypothesis to explore further. However, empirical variability in $C$ is typically confounded by differences in methodology among different investigators of different food webs (Martinez 1991, 1993b). Fortunately, consistent methodology (Havens 1992; Martinez 1993a; Srinivasan et al. 2007), use of trophic species (Martinez et al. 1999; Srinivasan et al. 2007) and moderate amounts of sampling effort (Martinez et al. 1999) may yield surprisingly robust measures of ecological complexity in terms of $C$. Applying such improved methodology to more and less invaded natural ecosystems could form an important test of whether ecological complexity in terms of trophic connectance strongly and consistently affects invasion success as suggested by our study.

Trophic position, and in particular the SCL of the invader, which tends to describe the dominant energy pathway of species (Williams & Martinez 2004a), accurately predicts invasion success in the introduction phase of our simulated invasions. However, as the invasion progresses, having a low SCL becomes less important than simply occupying a low trophic position. The distinction being that some species with high trophic positions may have a low SCL even though they feed mostly on high-trophic-position prey. The transition of importance from SCL to trophic position reflects the success of some of the primary carnivores that were successful in 19 per cent of invasion events and the secondary carnivores that were successful in 14 per cent of invasion events, which had high SCL but relatively low trophic positions.

These surprisingly intuitive and relatively simple results contradicted our expectation that invasion success and failure among our simulations would be as highly idiosyncratic similar to invasions in Nature (Williamson 1996). That is, each invasion outcome was thought to depend on fine nuances of each situation with invasion success based on the topology of the food web, the invaders fundamental and realized-niches, and fundamental and realized-niches of the other species in the web. Such idiosyncrasy could essentially preclude predictive simple regularities...
among invasions in our simulations and in Nature. However, instead of such idiosyncrasy, regularities identified by only a few predictive variables appeared among our simulations during both the introduction and the establishment stages of the invasions.

(a) Simulation robustness and further questions

The generality of our results are limited by the initial densities of invaders being randomly assigned from the same distribution as the initial densities of the species in the webs (e.g. between 0.5 and 1 for all species). This choice of such high initial invader biomass probably prevented us from detecting the effects of low invader biomass. Another explanation is that our observed insensitivity of invasion success to invader biomass mirrors the independence of invasion success to invader density found by recent meta-analyses that have found propagule pressure, but not invader density, to be one of the few general predictors of invasion success (Lockwood et al. 2005; Jeschke & Strayer 2006). Another possibility is that the relatively high initial invader biomass in our simulations insulates invaders from initial vulnerability to predators and weakens the role of invader vulnerability among properties at \( t=2000 \) compared with its role at \( t=4000 \). Large invader biomass increases the chance that invaders overcome a well-known strong resistance to invasions of single species at low densities (Post & Pimm 1983; Drake 1990; Hewitt & Huxel 2002). For example, Hewitt & Huxel (2002) manipulated the inoculation density of invaders in model food webs and found that low densities of invaders result in almost complete invasion resistance. Previous studies likewise set inoculation densities at arbitrarily low values relative to Lotka–Volterra equilibrium densities that mimic colonization by rare species (Post & Pimm 1983; Drake 1990). The assumption that invasive species are introduced at low densities does not conform to all empirical invasion scenarios (Carlton & Geller 1993). For example, empirical invasions where high densities of invaders are involved include the movement of entire assemblages with oyster culture transfers, hull fouling and ballast water release (Carlton 1985, 1987; Carlton & Geller 1993; Ruiz et al. 2000). When Hewitt & Huxel (2002) relaxed their assumption of low inoculation densities, they found that 50–60% of their single-species invasions were successful; closely reflecting our value (47%) for the proportion of successful invasions. In future work, we expect that invader biomass will play a stronger role in determining invasion success if initial invader biomass are distributed at and well below the equilibrium biomasses of the species in the webs.

Another aspect of our simulations affecting the generality of our results is the nature of the persistent niche model webs that are invaded. Our persistent webs are trophically shorter (low mean TL) and fatter (more basal and herbivore species) than niche webs not subjected to dynamics, which makes them more similar to food webs found in Nature (Williams & Martinez 2004a, 2008; Martinez et al. 2006; Thompson et al. 2007). However, the niche model tends to create food webs with less variability (Williams & Martinez 2000; Dunne et al. 2004) and slightly more intraspecific (Williams & Martinez 2000, 2008; Stouffer et al. 2006) than those that are found in Nature. In addition, due to computational considerations, we used a range of species richness that is lower than is found in empirical webs. Finally, our simulations ignore trophically important traits of plant nutrient consumption (Brose et al. 2005), variable body size (Brose et al. 2006; Otto et al. 2007) and non-trophic interactions such as facilitation. Such differences between our simulations and natural systems may limit the accuracy and applicability of our results. However, the many parallels between our results and many results of invasions found in Nature (see the electronic supplementary material for more examples) suggest that our simulations usefully inform invasion ecology.

5. CONCLUSION

Despite the common assertion that, other than propagule pressure and human affiliation (Williamson 1996; Jeschke & Strayer 2006), there are no strong taxa or biome independent determinants of invasion success, we identify potential generalizations describing successfully invasive species and more easily invaded communities by simulating species invasions within complex ecological networks. In general, an invader has a high probability of success if it is a generalist, a herbivore, an omnivore, a consumer with a feeding range that is low on a theoretical niche axis, and is relatively invulnerable to predation. Webs relatively easy to invade had low connectance, high mean biomass and a greater numbers of species. Integrating a complex-food web perspective into models of species invasion allows us to search a wide range of details that might affect invasibility and find potential systematic generalities among invasive species and habitats that promote successful invasions.

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REFERENCES

Allesina, S., Alonso, D. & Pascual, M. 2008 A general model for food web structure. Science 320, 658–661. (doi:10.1126/science.1156269)

Beddington, J. R. 1975 Mutual interference between parasites or predators and its effects on searching efficiency. J. Anim. Ecol. 14, 331–340. (doi:10.2307/3866)

Bright, C. 1998 Life out of bounds: bioinvasion in a borderless world. New York, NY: W.W. Norton & Company.

Briand, F. & Cohen, J. E. 1984 Community food webs have scale-invariant structure. Nature 307, 264–267. (doi:10.1038/307264a0)

Brose, U., Berlow, E. L. & Martinez, N. D. 2005 Scaling up keystone effects from simple to complex ecological
Species invasions in niche model webs

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networks. *Ecol. Lett.* 8, 1317–1325. (doi:10.1111/j.1461-0248.2005.00838.x)

Brose, U., Williams, R. J. & Martinez, N. D. 2006 Allometric scaling enhances stability in complex food webs. *Ecol. Lett.* 9, 1228–1236. (doi:10.1111/j.1461-0248.2006.00978.x)

Brown, J. H., Gillooly, J. F., Allen, A. P., Savage, V. M. & West, G. B. 2004 Toward a metabolic theory of ecology. *Ecology* 85, 1771–1789. (doi:10.1890/03-9000)

Camacho, J., Guimerà, R. & Amaral, L. A. N. 2002 Robust patterns in food web structure. *Phys. Rev. Lett.* 88, 228102. (doi:10.1103/PhysRevLett.88.228102)

Carlton, J. T. 1985 Transoceanic and interoceanic dispersal of coastal marine organisms: the biology of ballast water. *Oceanogr. Mar. Biol. Ann. Rev.* 23, 313–374.

Carlton, J. T. 1987 Patterns of transoceanic marine biological invasions in the Pacific Ocean. *Bull. Mar. Sci.* 41, 452–465.

Carlton, J. T. & Geller, J. B. 1993 Ecological roulette: the global transport and invasion of nonindigenous marine organisms. *Science* 261, 78–82. (doi:10.1126/science.261.5117.78)

Cattin, M. F., Bersier, L. F., Banasak-Richter, C., Baltensperger, R. & Gabriel, J. P. 2004 Phylogenetic constraints and adaptation explain food-web structure. *Nature* 427, 835–839. (doi:10.1038/nature02327)

Cohen, J. E. & Briand, F. 1984 Trophic links of community food webs. *Proc. Natl Acad. Sci. USA* 81, 4105–4109. (doi:10.1073/pnas.81.13.4105)

Cohen, J. E., Briand, F. & Newman, C. M. 1990 Community food webs: data and theory. Berlin, Germany: Springer.

DeAngelis, D. L., Goldstein, R. A. & O’Neill, R. V. 1975 A model for trophic interaction. *Ecology* 56, 881–892. (doi:10.2307/1936298)

Drapeau, P. A. 1990 The mechanics of community assembly and succession. *J. Theor. Biol.* 147, 213–233. (doi:10.1016/S0022-5193(05)80053-0)

Dunne, J. A. 2006 The network structure of food webs. In *Ecological networks: linking structure to dynamics in food webs* (eds M. Pascual & J. A. Dunne), pp. 27–86. Oxford, UK: Oxford University Press.

Dunne, J. A., Williams, R. J. & Martinez, N. D. 2002 Network structure and biodiversity loss in food webs: robustness increases with connectance. *Ecol. Lett.* 5, 558–567. (doi:10.1046/j.1461-0248.2002.00354.x)

Dunne, J. A., Williams, R. J. & Martinez, N. D. 2004 Network structure and robustness of marine food webs. *Mar. Ecol. Prog. Ser.* 273, 291–302. (doi:10.3354/meps273291)

Enserink, M. 1999 Biological invaders sweep in. *Science* 285, 1834–1843. (doi:10.1126/science.285.5435.1834)

Fox, J. W. 2006 Current food web models cannot explain the overall topological structure of observed food webs. *Oikos* 115, 97–109. (doi:10.1111/j.0030-0257.2006.1299.14868.x)

Fussmann, G. F., Ellner, S. P., Shertzer, K. W. & Hairston Jr, N. G. 2000 Crossing the Hopf bifurcation in a live predator–prey system. *Science* 290, 1358–1360. (doi:10.1126/science.290.5495.1358)

Hassell, M. P. & Varley, G. C. 1969 New inductive model for trophic interaction. *J. Theor. Biol.* 24, 213–233. (doi:10.1016/0022-5193(69)90025-K)

Martinez, N. D. 1992 Constant connectance in community food webs. *Am. Nat.* 139, 1208–1218. (doi:10.1086/253820)

Martinez, N. D. 1993a Effect of scale on food web structure. *Science* 260, 242–243. (doi:10.1126/science.260.5105.242)

Martinez, N. D. 1993b Effects of resolution on food web structure. *Oikos* 66, 403–412. (doi:10.2307/3544934)

Martinez, N. D. & Cushing, L. J. 2006 Additional model complexity reduces fit to complex food-web structure. In *Ecological networks: linking structure to dynamics in food webs* (eds M. Pascual & J. A. Dunne), pp. 87–89. Oxford, UK: Oxford University Press.

Martinez, N. D., Hawkins, B. A., Dawah, H. A. & Fei, P. 1999 Effects of sampling effort on characterization of food-web structure. *Ecology* 80, 1044–1055. (doi:10.2307/177037)

Martinez, N. D., Williams, R. J. & Dunne, J. A. 2006 Diversity, complexity, and persistence in large model ecosystems. In *Ecological networks: linking structure to dynamics in food webs* (eds M. Pascual & J. A. Dunne), pp. 167–185. Oxford, UK: Oxford University Press.

McCann, K. & Hastings, A. 1997 Re-evaluating the omnivory–stability relationship in food webs. *Proc. R. Soc. Lond. B* 264, 1249–1254. (doi:10.1098/rspb.1997.0172)

McCann, K. & Yodzis, P. 1995 Bifurcation structure of a three-species food chain model. *Theor. Popul. Biol.* 48, 93–125. (doi:10.1006/tpbi.1995.1023)

McCann, K., Hastings, A. & Huxel, G. R. 1998 Weak trophic interactions and the balance of nature. *Nature* 395, 794–798. (doi:10.1038/27427)

Montoya, J. M. & Solé, R. V. 2002 Small world patterns in food webs. *J. Theor. Biol.* 214, 405–412. (doi:10.1006/jtbi.2001.2460)

Otto, S. B., Rall, B. C. & Brose, U. 2007 Allometric degree distributions facilitate food web stability. *Nature* 450, 1226–1230. (doi:10.1038/nature06359)

Pascual, M. & Dunne, J. A. 2006 From small to large ecological networks in a dynamic world. In *Ecological networks: linking structure to dynamics in food webs* (eds M. Pascual & J. A. Dunne), pp. 3–24. Oxford, UK: Oxford University Press.

Post, W. M. & Pimm, S. L. 1983 Community assembly and food web stability. *Math. Biosci.* 64, 169–192. (doi:10.1016/0255-7805(83)90002-0)
