Density-dependent selection and the limits of relative

fitness

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1

Density-dependent selection and the limits of relative fitness

3 Abstract

Selection is commonly described in terms of relative fitness. Yet much of the ecological theory of selection in density-regulated populations seems to be incompatible with widelyused, constant-density relative fitness models such as the Wright-Fisher. Here we analyze the population ecological limits of relative fitness using a novel generalization of the Wright-Fisher model in which population density depends dynamically on the demographic rates of the types present. Our model clearly distinguishes between density-dependent selection and selection-dependent density. These two effects are confounded in many standard models of 10 density-regulated population growth. Both effects are necessary, in combination with strong 11 selection, for relative fitness to break down in populations close to demographic equilibrium. 12 Remarkably, both effects are not sufficient: we show that strong selection on a density-13 regulating trait that is subject to density-dependent selection can conform to the relative fitness description almost exactly. We reiterate the importance of reproductive excesses, 15 an aspect of selection omitted in many population growth models, which implies that even 16 strong selection need not affect density in many species. Our model also offers a possible 17 alternative to relative fitness when the latter is untenable, as is likely the case far from demographic equilibrium. (195 words)

21 Introduction

There are a variety of different measures of fitness. Some widely used examples in evolutionary ecology are expected lifetime reproductive ratio R_0 , intrinsic population growth rate r, saturation population density (often labeled "K") (Benton and Grant, 2000), and invasion fitness (Metz et al., 1992). In addition, "relative fitness" is the standard in much of evolutionary biology, particularly evolutionary genetics, where attention is generally restricted to relative genotypic proportions (Barton et al., 2007, pp. 468). The variety of fitness measures is not problematic in itself, because different measures may be more useful in different circumstances. But it should be clear how the measure being used is connected to the processes of birth and death which govern population biology (Metcalf and Pavard, 2007; Doebeli et al., 2017). While such a connection is fairly clear for absolute fitness measures like r, relative fitness seems largely divorced from population biology. It has even been proposed that relative fitness be justified from measure theory, abandoning population biology altogether (Wagner, 2010). Given the ubiquitous use of relative fitness, it is important that we understand its population ecological basis, both to clarify its domain of applicability, and 35 as part of the broader challenge of synthesizing ecology and evolution. 36 Constant relative fitness values can be justified as a linear approximation (Ewens, 2004, 37 277) (Charlesworth, 1994, Chap. 4) which is close to exact provided that selection 38 is sufficiently weak and stable over time. Yet strong, temporally-variable selection occurs 39 widely in nature and the lab, including in wild *Drosophila*, where population density varies by orders of magnitude each seasonal cycle (Messer et al., 2016; Bergland et al., 2014). 41 The question is whether relative fitness can be used when selection is not vanishingly weak. In age-structured populations that reproduce by outcrossing, selection cannot in general

be represented in terms of type-specific relative-fitness constants unless selection is weak

(Charlesworth, 1994, Chap. 4). We will therefore restrict our attention to asexual haploids

with little or no age structure, where the relative fitness description may be valid even if
selection is strong. This simplification also makes it easier to grasp the population ecological
underpinnings of selection, as opposed to the role of mating systems or complex changes in
age structure.

The basis of relative fitness is straightforward in the absence of crowding: it simply represents differences in intrinsic population growth rate. In discrete time, the change in frequency of type i is $\Delta p_i = \left(\frac{W_i}{\overline{W}} - 1\right) p_i$, where W_i is the intrinsic absolute growth factor of type i, and $\overline{W} = \sum_i W_i p_i$ is the population mean W. Here we can rescale W however we please and replace it with "relative fitness" w without affecting the ratio $\frac{W_i}{\overline{W}} = \frac{w_i}{\overline{w}}$. In continuous time, the canonical selection equation is $\frac{dp_i}{dt} = (r_i - \overline{r})p_i$, where W is replaced by the intrinsic exponential growth rate r (Crow et al., 1970, pp. 26). If there are two types present, a wildtype and a mutant for instance, then the continuous time canonical selection equation can be written as

$$\frac{dp_i}{dt} = sp_i(1 - p_i),\tag{1}$$

where the constant selection coefficient s is the difference in r between types. The corresponding adaptive sweeps follow a logistic curve.

The difficulty with Eq. (1) arises in crowded populations. Since crowded and uncrowded conditions are so different, we expect that s will often depend on density (Travis et al., 2013). Eq. (1) is then no longer a complete description of selection — we would also need to specify a model for how density is changing. Note that frequency-dependent selection does not raise similar problems; Eq. (1) is still a complete description of selection even if its behavior is more complicated due to s depending on frequency. Population genetics traditionally evades the issue of density-dependent selection by simply assuming that total population density N has reached its equilibrium value, which is assumed to be a fixed constant. The selection coefficient s now abstractly parameterizes the rate at which selection

changes relative frequencies, and no longer corresponds to differences in intrinsic growth rates r.

However, MacArthur famously argued that when population growth is density-regulated, 72 selection in crowded populations is intimately connected to the ability to keep growing at 73 higher densities than other types can tolerate (MacArthur and Wilson, 1967). The classic 74 example is the logistic model, where the type with the greatest saturation population den-75 sity "K" excludes the others (Fig. 1a). Similarly, the " R^* rule", a central tenet of resource 76 competition theory, states that when growth is limited by a single homogeneous consumable 77 resource, the type able to deplete the resource to the lowest equilibrium density R^* excludes the others (Grover, 1997). Differences in R^* will often entail differences in saturation density. The Lotka-Volterra competition model also couples selection in crowded populations 80 to density except in special cases (Smouse, 1976; Mallet, 2012). In these examples, both N81 and s change during, and as a result of, adaptive sweeps. It would therefore seem that the ubiquitous constant-N, relative fitness description of selection is incompatible with a huge 83 class of population ecological processes driving selection (Fig. 1b), even in the absence of age-structure and mating.

In light of this difficulty, the relative fitness description has been justified in broadly two different ways for crowded populations (we do not discuss Wagner's [2010] measuretheoretical justification, which is independent of population biology). The first is to simply assume that selection is density-independent but relax the assumption of constant N by allowing density to change as a result of selective sweeps (Barton et al., 2007, pp. 468)
(Prout, 1980). Obviously this does not address the problem that s can, in reality, depend on density. Type-specific responses to density are at the center of MacArthur's argument and the density-dependent selection literature that grew out of it (e.g. (Roughgarden, 1979)).

The second justification, which primarily grew out of a controversy over Haldane's "cost of selection", is to appeal to the existence of a "reproductive excess" of juveniles that are more

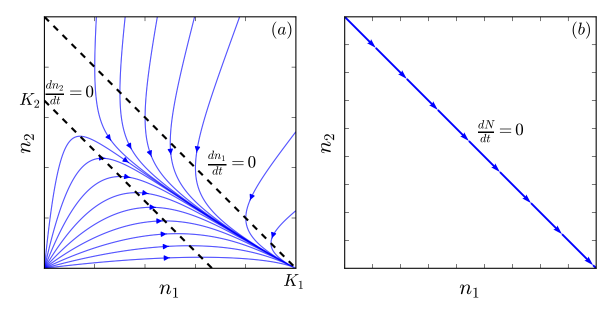


Figure 1: Selection in crowded environments shown as a phase diagram for the densities of two types n_1 and n_2 . (a) The logistic model $\frac{dn_1}{dt} = r_1(1 - \frac{n_1 + n_2}{K_1})n_1$ and $\frac{dn_2}{dt} = r_2(1 - \frac{n_1 + n_2}{K_2})n_1$ with $r_1 = r_2$ and $K_1 > K_2$. (b) The constant-N, relative fitness description of selection.

fragile than their adult counterparts (Turner and Williamson, 1968; Kimura and Crow, 1969; Nei, 1971). Selection can then be concentrated at the juvenile phase, uncoupling selection 97 from population density at the adult phase unless it is so strong that the reproductive excess 98 is depleted. This justifies Eq. (1) because, for a population in demographic equilibrium, 99 selective sweeps do not affect density, and so the density-dependence of selection does not 100 Unfortuantely this reproductive excess literature is also poorly integrated with 101 population ecology. Kimura and Crow (1969) took constant N as a requirement and then 102 derived some variants of the logistic model that satisfy this requirement. Nei (1971) proposed 103 a model with an explicit representation of reproductive excess, but used an unusual model of 104 competition based on pair-wise interactions which was only defined for at most two different 105 types. As a result, the role of reproductive excesses in justifying Eq. (1) is still largely verbal. 106 Here we study the population ecology of relative fitness using a novel model of density-107 dependent population growth based on territorial contests. Rather than attempting to make

sense of relative fitness in existing standard models of population growth (e.g. (Kimura and Crow, 1969; Mallet, 2012)), we instead do the reverse, and attempt to make population eco-110 logical sense of the widely-used Wright-Fisher relative-fitness model. Our starting point is 111 the classic lottery model of territorial contest (Sale, 1977; Chesson and Warner, 1981). Like 112 the Wright-Fisher model, the classic lottery assumes a saturated population with constant N, 113 and fitness involves a product of fertility and juvenile viability (Crow et al., 1970, pp. 185), 114 but unlike the Wright-Fisher model, generations can overlap. Our first task is to generalize 115 the lottery model to create a variable-density version of the Wright-Fisher model with over-116 lapping generations (sections "Model" and "Analytical approximation of the variable-density 117 lottery"). 118 Equipped with this new model, we turn to the evaluation of Eq. (1). We first discuss 119 selection on the ability to contest territories, which behaves like a pure constant-N, relative 120 fitness trait, and discuss how this fits with MacArthur's analysis of selection in crowded pop-121 ulations (section "K-selection and selection-dependent density"). We then consider selection 122 on density-regulating traits (section "Density-regulating traits and the threat of strong selec-123 tion"), and conclude by contrasting the classical density-dependent selection literature with 124 our results ("Discussion").

Model Model

Assumptions and definitions

We assume that reproductively mature individuals ("adults") each require their own territory to survive and reproduce. All territories are identical, and the total number of territories is T.

Time advances in discrete iterations, each representing the time from birth to reproductive maturity. In a given iteration, the number of adults of the i'th type will be denoted by n_i , the total number of adults by $N = \sum_i n_i$, and the number of unoccupied territories by

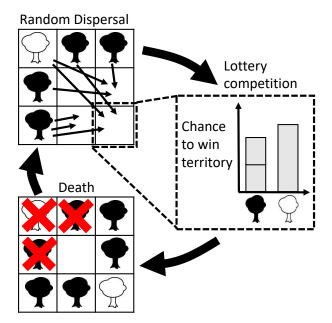


Figure 2: Each iteration of our model has three elements. First, propagules are produced by adults and dispersed at random (only propagules landing on unoccupied territories are shown). Some territories may receive zero propagules. Lottery competition then occurs in each unoccupied territory that receives a propagule (only illustrated in one territory). Each type has a probability proportional to $c_i x_i$ of securing a given territory, where c_i measures competitive ability and x_i is the number of propagules that disperse there. In the illustrated territory, the black type disperses more propagules but is a poorer competitor. Territories are then made available by deaths among those adults present at the start of the iteration (red crosses).

U = T - N. We assume that the n_i are large enough that stochastic fluctuations in the n_i 133 ("drift") can be ignored (with T also assumed large to allow for small type densities n_i/T). 134 Each iteration, adults produce new offspring ("propagules"). These disperse at random, 135 regardless of distance from their parents, and independently of each other (e.g. there is 136 no avoidance of territories crowded with propagules). We assume that adults cannot be 137 ousted by juveniles, so that propagules landing on occupied territories are doomed. We 138 assume that each adult from type i produces a constant number b_i of successfully dispersing 139 propagules; the number of propagules dispersing to unoccupied territories is then given by 140 $m_i = b_i n_i U/T$. The total number of these propagules will be denoted $M = \sum_i m_i$. Note that due to our assumption of uniform dispersal, the parameter b_i can be thought of as a measure of "colonization ability", which combines fecundity and dispersal ability (Levins and Culver, 1971; Tilman, 1994; Bolker and Pacala, 1999).

The number of propagules of the i'th type landing in any particular territory is denoted x_i . We assume that x_i follows a Poisson distribution $p_i(x_i) = l_i^{x_i} e^{-l_i}/x_i!$, where $l_i = m_i/U$ is the mean territorial propagule density for type i (the total propagule density will be denoted $L = \sum_i l_i$). This is strictly only an approximation of random dispersal, but it is an excellent approximation given our assumption that the n_i are large enough that drift can be ignored (Appendix A).

When multiple propagules land on the same territory, the victor is determined by lottery competition: type i wins a territory with probability $c_i x_i / \sum_j c_j x_j$, where c_i is a constant representing relative competitive ability (Fig. 2). We expect that a fraction $p_1(x_1) \dots p_G(x_G)$ of the U unoccupied territories will have the propagule composition x_1, \dots, x_G . Type i is expected to win a proportion $c_i x_i / \sum_j c_j x_j$ of these. Ignoring fluctuations about these two expectations (due to our no-drift, large n_i , large T approximation), type i's territorial acquisition is given by

$$\Delta_{+} n_{i} = U \sum_{x_{1}, \dots, x_{G}} \frac{c_{i} x_{i}}{\sum_{j} c_{j} x_{j}} p_{1}(x_{1}) \cdots p_{G}(x_{G}), \tag{2}$$

where the sum only includes territories with at least one propagule present.

Finally, we assume that adult mortality only occurs in adults present at the start of the iteration, and at a constant, type-specific per-capita rate $0 \le d_i \le 1$. Thus, the overall change in type abundances is

$$\Delta n_i = \Delta_+ n_i - d_i n_i. \tag{3}$$

Fig. 2 illustrates one iteration of the model.

63 Connection to the Wright-Fisher and classic lottery models

In the classic lottery model (Chesson and Warner, 1981), unoccupied territories are assumed to be saturated with propagules from every type $l_i \gg 1$. From the law of large numbers, the 165 composition of propagules in each territory will then not deviate appreciably from the mean 166 composition l_1, l_2, \ldots, l_G (G is the number of types present), and so the probability that type 167 i wins any particular unoccupied territory is approximately $c_i l_i / \sum_j c_j l_j$. Then the numbers 168 of territories won by each type $\Delta_+ n_1, \Delta_+ n_2, \dots, \Delta_+ n_G$ follow a multinomial distribution with 169 U trials and success probabilities $\frac{c_1 l_1}{\sum_j c_j l_j}, \frac{c_2 l_2}{\sum_j c_j l_j}, \dots, \frac{c_G l_G}{\sum_j c_j l_j}$, respectively. Type i is expected 170 to win a proportion $c_i l_i / \sum_j c_j l_j$ of the *U* available territories, and deviations from this 171 expected outcome are small (since T is large by assumption), giving 172

$$\Delta_{+}n_{i} = \frac{c_{i}l_{i}}{\sum_{j}c_{j}l_{j}}U = \frac{c_{i}l_{i}}{\overline{c}L}U,$$
(4)

where $\bar{c} = \sum_j c_j m_j / M$ is the mean competitive ability for a randomly selected propagule. In section "Analytical approximation of the density-dependent lottery", we derive a generalization of Eq. (4) that accommodates arbitrary propagule densities l_i .

There is a close connection between the classic lottery model and the Wright-Fisher 176 model of genetic drift (Svardal et al., 2015). In the Wright-Fisher model, type abun-177 dances are sampled each generation from a multinomial distribution with success proba-178 bilities $w_i n_i / \sum_j w_j n_j$, where w is relative fitness and the n_i are type abundances in the preceding generation. Population size N remains constant. This is equivalent to the classic 180 lottery model with non-overlapping generations ($d_i = 1$ for all i) and relative fitness given 181 by $w_i = b_i c_i$ i.e. a product of fertility and viability (Crow et al., 1970, pp. 185). Thus, the 182 classic lottery model is essentially the Wright-Fisher model extended to allow overlapping 183 generations, but ignoring drift. This means that our extension of the classic lottery model to 184 arbitrary densities represents a variable-density generalization of the Wright-Fisher model 186 (we also do not consider drift here).

Results

Analytical approximation of the variable-density lottery

Eq. (2) involves an expectation over the time-dependent dispersal distributions p_i , and is thus too complicated to give intuition about the dynamics of density-dependent lottery competition. We now evaluate this expectation.

Similarly to the high- l_i approximation of the classic lottery model, we replace the x_i with 192 appropriate mean values, although we cannot simply replace x_i with l_i as in Eq. (4). The 193 classic lottery model breaks down for types with low propagule density $(l_i \ll 1)$ because 194 territorial acquisition is then not correctly represented by a lottery in each territory with the 195 mean propagule density. For a type with low propagule density $l_i \ll 1$, we have $x_i = 1$ in the 196 territories where its propagules land, and so its growth comes entirely from territories which deviate appreciably from l_i . To account for this, we separate Eq. (2) into $x_i = 1$ and $x_i > 1$ 198 parts. Our more general approximation only requires that there are no large discrepancies in competitive ability (i.e. we do not have $c_i/c_j \gg 1$ for any two types). We obtain (details 200 in Appendix B) 201

$$\Delta_{+} n_{i} \approx \left[e^{-L} + (R_{i} + A_{i}) \frac{c_{i}}{\overline{c}} \right] l_{i} U, \tag{5}$$

202 where

$$R_i = \frac{\overline{c}e^{-l_i}(1 - e^{-(L-l_i)})}{c_i + \frac{\overline{c}L - c_i l_i}{L - l_i} \frac{L - 1 + e^{-L}}{1 - (1 + L)e^{-L}}},$$

203 and

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$$A_i = \frac{\overline{c}(1 - e^{-l_i})}{\frac{1 - e^{-l_i}}{1 - (1 + l_i)e^{-l_i}} c_i l_i + \frac{\overline{c}L - c_i l_i}{L - l_i} \left(L \frac{1 - e^{-L}}{1 - (1 + L)e^{-L}} - l_i \frac{1 - e^{-l_i}}{1 - (1 + l_i)e^{-l_i}} \right)}.$$

Comparing Eq. (5) to Eq. (4), the classic lottery per-propagate success rate $c_i/\bar{c}L$

has been replaced by three separate terms. The first, e^{-L} , accounts for propagules which land alone on unoccupied territories; these territories are won without contest. The second, $R_i c_i/\bar{c}$, represents competitive victories when the i type is a rare invader in a high density population (i.e. it determines invasion fitness (Metz et al., 1992)). The third term, $A_i c_i/\bar{c}$, represents competitive victories when the i type is abundant. The relative importance of these three terms varies with both the overall propagule density L and the relative propagule frequencies l_i/L . If $l_i \gg 1$ for all types, we recover the classic lottery model (only the $A_i c_i/\bar{c}$ term remains, and $A_i \to 1/L$).

Fig. 3 shows that Eq. (5) and its components closely approximate simulations of our variable-density lottery model over a wide range of propagule densities. Two types are present, one of which is at low frequency. The growth of the low-frequency type relies crucially on the low-density competition term $R_i c_i/\bar{c}$. On the other hand, $R_i c_i/\bar{c}$ is negligible for the high-frequency type, which depends instead on high density territorial victories. Fig. 3 also shows the breakdown of the classic lottery model at low propagule densities.

Eq. (5) takes a much simpler form if all types are competitively equivalent $(c_i = c)$,

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$$\Delta_{+}n_{i} = \frac{l_{i}}{L}(1 - e^{-L})U. \tag{6}$$

Here $1 - e^{-L}$ is the fraction of territories that receive at least one propagule under Poisson dispersal, $(1 - e^{-L})U$ is the total number of territories gained, and type i receives a fraction l_i/L of these. Total population density thus grows according to

$$\Delta N = (1 - e^{-L})U - \sum_{i} d_i n_i \tag{7}$$

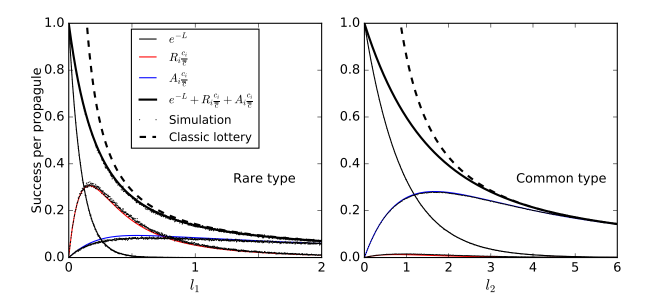


Figure 3: Comparison of the analytical approximation Eq. (5) with simulations. Perpropagule success probability $\Delta_+ n_i/l_i U$ from the classic lottery model, individual-based simulations of random dispersal and lottery competition, and Eq. (5) and its three components. Two types are present, a rare type with $c_1 = 1.5$, and a common type with $c_2 = 1$. Simulation points are almost invisible in for the common type due to near exact agreement with Eq. (5). Dashed lines in show the breakdown of the classic lottery model. Parameters: $m_1 = 10^4$ and $m_2 = 9 \times 10^4$ and U varies between 5×10^3 and 10^6 .

Selection-dependent density and K-selection

Equipped with our variable-density lottery model, we now start evaluating the validity of Eq. (1). In this section we explore whether we should expect population density to vary as a result of selection (Prout, 1980). Since the idea that density does vary with selection is closely connected to the notion of "K-selection", we start by revisiting MacArthur's analysis of selection in crowded environments (MacArthur and Wilson, 1967).

MacArthur considers a population with two types that have densities n_1 and n_2 subject to density-dependent growth described by

$$\frac{dn_1}{dt} = f_1(n_1, n_2) \qquad \frac{dn_2}{dt} = f_2(n_1, n_2). \tag{8}$$

The environment is assumed to remain constant apart from the type densities. The functions f_1 and f_2 must decline to zero if n_1 or n_2 are sufficiently large, because no population has unlimited resources. This defines the nullclines $f_1(n_1, n_2) = 0$ and $f_2(n_1, n_2) = 0$ in (n_1, n_2) space. The outcome of selection is then determined by the relationship between these nullclines. Specifically, a type will be excluded if its nullcline is completely contained in the region bounded by the other type's nullcline. In other words, for a type to have the possibility of persisting, it must be able to keep growing to higher densities than the other type can tolerate in some region of (n_1, n_2) space (Fig. 1a).

To formalize the relationship between nullclines, MacArthur used the symbol "K" to label the four intersection points of the nullclines with the n_1 and n_2 axes, specifically $f_1(K_{11},0) = 0$, $f_1(0,K_{12}) = 0$, $f_2(0,K_{22}) = 0$ and $f_2(K_{21},0) = 0$. These K values determine whether a region of higher-density growth exists for each type, provided that the nullclines are close to being straight lines. Note that only K_{11} and K_{22} are saturation densities akin to the K parameter in the logistic model; following widespread convention, we will refer to selection on these saturation densities as "K-selection" (Fig. 1a). The other intersection

points, K_{12} and K_{21} , are related to competition between types. For instance, in the Lotka-Volterra competition model we have

$$f_1(n_1, n_2) = r_1(1 - \alpha_{11}n_1 - \alpha_{12}n_2)n_1$$

$$f_2(n_1, n_2) = r_2(1 - \alpha_{22}n_1 - \alpha_{21}n_2)n_2$$
(9)

where $\alpha_{11} = 1/K_{11}$ and $\alpha_{22} = 1/K_{22}$ measure competitive effects within types, while $\alpha_{12} = 1/K_{12}$ and $\alpha_{21} = 1/K_{21}$ measure competitive effects between types (Fig. 4a).

Thus, when MacArthur concludes that "fitness is K" in crowded populations (MacArthur 241 and Wilson, 1967, pp. 149), the meaning is that selection either favors the ability to keep 242 growing at ever higher densities (moving a type's own nullcline outwards), or the ability to 243 suppress the growth of competitors at lower densities (moving the nullcline of competitors 244 inwards) (Gill, 1974). This general idea is much broader than "K-selection" in the sense 245 of selection for greater saturation density, and applies even if the nullclines are nonlinear 246 to such an extent that the "K" values themselves do not give much information about the 247 regions of high-density growth. 248

It is obvious from Eq. (9) that selection can favor a superior competitor in a crowded 249 population even if its saturation density is the same as, or lower than that of the other types 250 present. However, the Lotka-Volterra model still couples selection to population density 251 (Smouse, 1976). Fig. 4a shows Lotka-Volterra selection between two types with the same 252 saturation density ($\alpha_{11} = \alpha_{22}, \alpha_{21} > \alpha_{12}$). Even though the initial and final densities of a 253 sweep are the same, density is not constant over a sweep. Only a highly restricted subset 254 of r and α values will keep N constant over a selective sweep (further details in Appendix 255 C). Intuitively, for one type to exclude another with the same saturation density, competi-256 tive suppression of growth between types must be stronger than competitive suppression of 257 growth within types, causing a dip in N over the sweep. 258

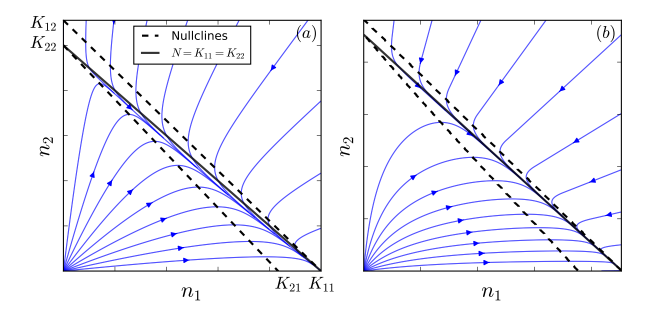


Figure 4: Selection between types with identical saturation density but different inter-type competitive ability. (a) Lotka-Volterra competition (Eq. 9) with $r_1 = r_2 = 1$, $\alpha_{11} = \alpha_{22} = 1$, $\alpha_{12} = 0.9$ and $\alpha_{21} = 1.2$. Trajectories do not follow the line $N = K_{11} = K_{22}$. (b) Lottery competition (Eq. 5) with $b_1 = b_2 = 5$, $d_1 = d_2 = 0.1$ and $c_1/c_2 = 5$. Trajectories converge on the line $N = K_{11} = K_{22}$.

By contrast, if one type in our density-dependent lottery model has a c advantage but 259 the types are otherwise identical (so that each type has the same saturation density), the 260 density trajectories converge on the line of constant density equal to the saturation density 261 (Fig. 4b). Selection then occurs purely along this line, similarly to Fig. 1b. This occurs 262 because c does not directly affect N: it only affects the relative likelihood for each type 263 to win a contested territory, not whether a territory is contested in the first place (this 264 can be seen formally in Eq. (7)). In other words, once the population reaches demographic 265 equilibrium, it behaves indistinguishably from a constant-N relative fitness model. While 266 quite different from classical growth models like the Lotka-Volterra, this is all perfectly 267 consistent with MacArthur's general argument. 268

The constant-N behavior of c-selection arises from the role of c as a trait determining relative competitive success in territorial contests. As such, this behavior is a result of a reproductive excesses Eq. (1). By contrast, previous models of selection-independent density either used unusual models of competition (Kimura and Crow, 1969; Nei, 1971), or made restrictive parameter choices in the Lotka-Volterra model (Appendix C; Smouse 1976; Mallet 2012).

²⁷⁵ Density-regulating traits and the threat of strong selection

In the previous section we showed that c-selection and the regulation of population density are independent even though population growth is density-regulated in our variable-density lottery. Nevertheless, selection and density regulation are intimately connected in widely used models of population growth, as well as for the lottery b and d traits.

To see why this connection potentially poses a threat to relative fitness, consider the simple birth-death model (Kostitzin, 1939, pp. 20) (Travis et al., 2013)

$$\frac{dn_i}{dt} = (b_i - \delta_i N)n_i \tag{10}$$

where δ_i is the per-capita increase in mortality rate due to crowding (for simplicity, there are no deaths when uncrowded), playing a similar role as K in the logistic model.

Starting from a monomorphic population, the frequency of a $\delta_i \to \delta_i(1-\epsilon)$ variant obeys

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$$\frac{dp_i}{dt} = \epsilon \delta_i N p_i (1 - p_i). \tag{11}$$

The selection coefficient $s = \epsilon \delta_i N$ thus depends on density (compare with Model III in Kimura and Crow (1969)). On the other hand, the frequency of a $b_i \to b_i (1 + \epsilon)$ variant will exactly obey Eq. (1) with $s = \epsilon b_i$, independent of density.

In practice the density dependence in Eq. (11) only matters if N changes substantially 288 during a sweep. This can easily occur if a population is far from demographic equilibrium 289 (we return to this scenario in the Discussion). However, even if N has reached equilibrium, 290 it will change substantially over a δ -sweep if selection on δ is sufficiently strong. To quantify 291 this effect, we need to account for how much N changes as a result of a δ -sweep beginning 292 and ending in equilibrium (Kimura and Crow, 1969); from Eq. (10) we have an increase 293 from $N_{\text{initial}} = b_i/\delta_i$ to $N_{\text{final}} = b_i/\delta_i(1-\epsilon) = N_{\text{initial}}/(1-\epsilon)$. The corresponding selection 294 coefficient increases from $s_{\text{initial}} = \epsilon b_i$ to $s_{\text{final}} = s_{\text{initial}}/(1-\epsilon)$. Consequently, noticeable 295 deviations from Eq. (1) occur with proportional changes to δ of order $\epsilon = 0.2$ and upwards i.e. selection must be quite strong (Fig. 5).

Let us now turn to selection on b and d in our lottery model. Recall that $m_i = b_i n_i U/T$, and so $L = M/U = \bar{b}N/T$ where \bar{b} is the population mean b. Thus, from Eq. (6) we have

$$\Delta n_i = \left(\frac{b_i}{\overline{b}} \frac{1 - e^{-\overline{b}N/T}}{N} (T - N) - d_i\right) n_i. \tag{12}$$

It can be seen that the mortality d is akin to the birth rate in Eq. (10), and so, while d does affect density, selection on d is density independent. Thus, d sweeps follow the canonical

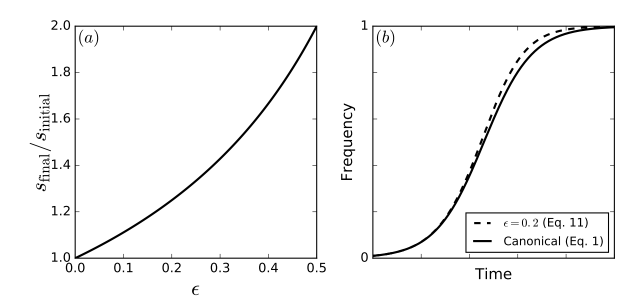


Figure 5: (a) Proportional change in the selection coefficient over a "K-like" sweep for a type that experiences proportionally $1-\epsilon$ fewer deaths induced by crowding. The population is in demographic equilibrium at the start and end of the sweep. (b) Example equilibrium-to-equilibrium δ -sweep (Eq. 11) for $\epsilon=0.2$ showing a noticeable deviation from the canonical selection equation.

relative fitness model exactly (Fig. 6).

At first glance, b in Eq. (12) appears to be analogous to the δ in Eq. (10) because it regulates density and is multiplied by the density-dependent term $f(\bar{b}, N) = \frac{1 - e^{-\bar{b}N/T}}{N}(T - N)$.

This term declines from \bar{b} at low density to zero at high density and as a result, selection on b is density-dependent. The source of this density-dependence is that the selective advantage from having greater b depends on the number of territories being contested; if almost all are occupied, then there is little advantage to having a greater b.

Nevertheless, the behavior of equilibrium-to-equilibrium b-sweeps are qualitatively differ-309 ent from the δ sweeps above. The reason is that b regulates density by controlling how many 310 unoccupied territories receive propagules. Thus, greater b means more propagules contesting 311 territories, but also more territories being contested. The net effect on $f(\bar{b}, N)$ is precisely 312 zero in equilibrium: in a single-type equilibrium we have $b_i/\bar{b}=1$ and so $f(\bar{b},N)=d_i$ exactly 313 at the beginning and end of a pure b sweep, even though the density N increases. Strictly 314 speaking there is some deviation in f(N) from d_i during the sweep, but this deviation is an 315 order of magnitude smaller than for a δ sweep (the deviation due to a sweep with propor-316 tional effect $b_i \to b_i(1+\epsilon)$ is only of order ϵ^2 , whereas the analogous effect in Fig. 5 is of order 317 ϵ ; see Appendix D for details). Since selection must already be quite strong for a δ sweep to 318 threaten Eq. (1), we conclude that b sweeps also obey the canonical selection equation (to a 319 close approximation).

Discussion

Summarizing the three traits in the variable-density lottery model: (i) c-selection is densitydependent, but c does not regulate density; (ii) d regulates density, but d-selection is densityindependent; (iii) b regulates density and b-selection is density-dependent. Despite these
differences, pure b, c and d sweeps starting and ending at equilibrium all obey the canonical

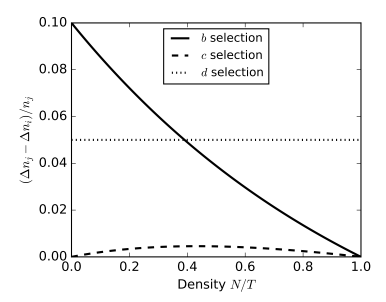


Figure 6: The density-dependence of selection in our variable-density lottery plotted as the selection coefficient $(\Delta n_j - \Delta n_i)/n_j$ experienced by an adaptive variant j present at the same frequency as a wildtype i. Here $b_i = 1$, $d_i = 0.5$ and $c_i = 1$. For b-selection we set $b_j = b_i(1 + \epsilon)$, and similarly for c and d, with $\epsilon = 0.1$. d-selection is density-independent, b-selection gets weaker with lower territorial availability, while c-selection initially increases with density as territorial contests become more important, but eventually also declines as available territories become scarce. The equilibrium density for the i type is ≈ 0.4 .

selection equation. This rich variety of behaviors in relation to density is quite different from that found in the classical density-dependent selection literature (Roughgarden, 1979; Christiansen, 2004).

To briefly review: based on a diploid, bi-allelic variant of the logistic model, the r/K329 scheme proposed a dichotomy between r-selection (uncrowded) and K-selection (crowded) 330 (MacArthur, 1962), with the latter taken to mean selection for greater saturation density 331 (Gill, 1974). A more general Lotka-Volterra model introduces the inter-type α_{ij} competition 332 coefficients, with selection on these termed " α -selection" (Gill, 1974; Joshi et al., 2001). 333 Setting aside r which confers no selective advantage at equilibrium, we are left with K and 334 α , which both behave like δ in that they are density-dependent and cause density to change 335 over a sweep (although N only dips temporarily during an α -sweep). Thus, strong selection 336 is sufficient for relative fitness to break down in the classical view of density-dependent 337 Indeed, in the defense of Eq. (1) given by Kimura and Crow (1969), it was 338 necessary to assume that s will be a few percent at most. 339

Our variable-density lottery model shows that it is not simply a lack of ecological realism that underlies the contrast between relative fitness with the classical view of densitydependent selection. Rather, in many population growth models, only one life-history stage
is represented, and the competitive effects resulting from crowding appear as a reduction in
absolute fitness that only depends on the type densities at this life-history stage (e.g. the n_i^2 and $n_i n_j$ terms in the Lotka Volterra equation). As noted in the introduction, this precludes selection concentrated at a fragile juvenile stage as a result of a reproductive excess
(Chesson, 1983; Turner and Williamson, 1968; Kimura and Crow, 1969; Nei, 1971).

Reproductive excesses appear in the variable-density lottery model when the number of propagules is greater than the number of available territories. Then only $\approx 1/L$ of the juveniles contesting available territories survive to adulthood. Unlike the role of adult density n_i in single-life-stage models, it is the propagule densities l_i that represent the crowding

that drives competition (a "critical age-group"; Charlesworth 1994, pp. 54). In general,
reproductive excesses will tend to produce strictly-relative lottery-type contests in which
fitter types grow at the expense of others by preferentially filling the available adult "slots".

The number of slots can remain fixed or change independently of selection at the juvenile
stage. By ignoring reproductive excesses, single life-stage models are biased to have total
population density be sensitive to ongoing selection. In this respect, the Wright-Fisher model
and similar viability selection heuristics actually capture an important ecological process.

We now turn to the breakdown of Eq. (1). We first discuss the problem shown in Fig. 5, 359 which occurs when strong selection changes population density and is also density-dependent. 360 In the variable-density lottery, this occurs if and only if types differ in more than one trait. 361 The c and d traits represent the two distinct directions in which density and selection inter-362 act: selection may depend on density, and density may depend on selection (Prout, 1980). 363 The combination is necessary to pose a threat to Eq. (1). However, the b trait remark-364 ably demonstrates that the combination is not sufficient, since the density-dependence of 365 b-selection disappears over equilibrium-to-equilibrium b-sweeps. Thus, the simple linear mod-366 els that have become standard in discussions of density-dependent selection (Roughgarden, 367 1979; Christiansen, 2004; Mallet, 2012; Travis et al., 2013) actually represent a complicated form of the interaction between density and selection, and their parameters confound the underlying issues. 370

While this is a conceptual reason to be wary of the classical density-dependent selection models, it is not clear how we should expect the trait variation in nature to align. For instance, should we expect mutations to generally affect b, c and d independently of each other, or pleiotropically such that δ -like selection is prevalent? In the case of well-mixed indirect exploitation competition for consumable resources, the R^* rule suggests that δ -like selection will be prevalent. However, for many populations consumable resources are not well-mixed. Spatial localization of consumable resources (e.g. due to restricted movement of nutrients through soils) will tend to create a territorial situation similar to the lottery model,
where resource competition only occurs locally and both it any interference competition are
subsumed into the competitive ability c, which does not affect N.

Relative fitness models truly break down when N is far from equilibrium and selection 381 is density-dependent (as seems likely; Travis et al. 2013). For example, wild Drosophila 382 experience large seasonal boom-bust cycles in population density coupled to strong selection 383 that drives large swings in allele frequency (Bergland et al., 2014). In this case there is no 384 choice but to abandon relative fitness, and our model provides one potentially suitable option. 385 Whether or not our density-dependent lottery model is a good description of *Drosophila* 386 ecology, the close connection between our model and Wright-Fisher is useful, because drift 387 in our model should behave broadly similarly. Thus, our model it should provide a useful 388 starting point for analyzing evolution in this and other far-from-equilibrium situations. 380

Another issue with the constant-N relative fitness description of selection is that it precludes consideration of longer-term aspects of the interplay between evolution and ecology
such as population extinction. A variety of approaches have been developed for dealing with
these issues in quantitative genetics (Burger and Lynch, 1995; Engen et al., 2013), population genetics (Bertram et al., 2017) and adaptive dynamics (Ferriere and Legendre, 2013;
Dieckmann and Ferrière, 2004). Although density-dependent selection is pertinent to these
longer-term issues (Travis et al., 2013), our focus here has been the description of the timedependent process by which selection changes allele frequencies. This is particularly critical
for making sense of evolution at the genetic level, for which we now have abundant data.

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484 Appendix A: Poisson approximation

- For simplicity of presentation, we have assumed a Poisson distribution for the x_i as our
- model of dispersal. Strictly speaking, the total number of i propagules $\sum x_i$ (summed over
- unoccupied territories) is then no longer a constant m_i , but fluctuates between generations

for a given mean m_i , which is more biologically realistic. Nevertheless, since we do not consider the random fluctuations in type abundances here, and for ease of comparison with the classic lottery model, we ignore the fluctuations in m_i . Instead we focus, on Poisson fluctuations in propagule composition in each territory.

In the exact model of random dispersal, the counts of a type's propagules across un-492 nocupied territories follows a multinomial distribution with dimension U, total number of 493 trials equal to m_i , and equal probabilities 1/U for a propagale to land in a given territory. 494 Thus, the x_i in different territories are not independent random variables. However, for suf-495 ficiently large U and m_i , this multinomial distribution for the x_i across territories is closely 496 approximated by a product of independent Poisson distributions for each territory, each with 497 rate parameter l_i (Arenbaev, 1977, Theorem 1). Since we are ignoring finite population size 498 effects, we effectively have $T \to \infty$, in which case U can only be small enough to violate 499 the Poisson approximation if there is vanishing population turnover, and then the dispersal 500 distribution is irrelevant anyway. Likewise, in ignoring stochastic finite population size for 501 the n_i , we have effectively already assumed that m_i is large enough to justify the Poisson 502 approximation (the error scales as $1/\sqrt{m_i}$; Arenbaev 1977). 503

Appendix B: Derivation of growth equation

We separate the right hand side of Eq. (2) into three components $\Delta_+ n_i = \Delta_u n_i + \Delta_r n_i + \Delta_a n_i$ which vary in relative magnitude depending on the propagule densities l_i . Following the notation in the main text, the Poisson distributions for the x_i (or some subset of the x_i) will be denoted p, and we use P as a general shorthand for the probability of particular outcomes.

510 Growth without competition

The first component, $\Delta_u n_i$, accounts for territories where only one focal propagule is present $x_i = 1$ and $x_j = 0$ for $j \neq i$ (u stands for "uncontested"). The proportion of territories where this occurs is $l_i e^{-L}$, and so

$$\Delta_u n_i = U l_i e^{-L} = m_i e^{-L}. \tag{13}$$

514 Competition when rare

The second component, $\Delta_r n_i$, accounts for territories where a single focal propagule is present along with at least one non-focal propagule (r stands for "rare") i.e. $x_i=1$ and $X_i\geq 1$ where $X_i=\sum_{j\neq i}x_j$ is the number of nonfocal propagules. The number of territories where this occurs is $Up_i(1)P(X_i\geq 1)=b_in_ie^{-l_i}(1-e^{-(L-l_i)})$. Thus

$$\Delta_r n_i = m_i e^{-l_i} \left(1 - e^{-(L-l_i)}\right) \left\langle \frac{c_i}{c_i + \sum_{j \neq i} c_j x_j} \right\rangle_{\tilde{p}}, \tag{14}$$

where $\langle \rangle_{\tilde{p}}$ denotes the expectation with respect to \tilde{p} , and \tilde{p} is the probability distribution of nonfocal propagule abundances x_j after dispersal (the significance of evaluating after dispersal should become clearer below), in those territories where exactly one focal propagule, and at least one non-focal propagule, landed.

Our approximation is to replace x_i with its mean in the last term in Eq. (14),

$$\left\langle \frac{c_i}{c_i + \sum_{j \neq i} c_j x_j} \right\rangle_{\tilde{p}} \approx \frac{c_i}{c_i + \sum_{j \neq i} c_j \langle x_j \rangle_{\tilde{p}}}.$$
 (15)

Below we justify this replacement by arguing that the standard deviation $\sigma_{\tilde{p}}(\sum_{j\neq i} c_j x_j)$ (with respect to \tilde{p}), is much smaller than $\langle \sum_{j\neq i} c_j x_j \rangle_{\tilde{p}}$. If this standard deviation is small as claimed, then the mean value gives an accurate representation of the values taken by $\sum_{j\neq i} c_j x_j$ across territories.

We first calculate $\langle x_j \rangle_{\tilde{p}}$. Let $X = \sum_j x_j$ denote the total number of propagules in a territory and $\mathbf{x}_i = (x_1, \dots, x_{i-1}, x_{i+1} \dots, x_G)$ denote the vector of non-focal abundances, so that $p(\mathbf{x}_i) = p_1(x_1) \cdots p_{i-1}(x_{i-1}) p_{i+1}(x_{i+1}) \cdots p_G(x_G)$. Then, \tilde{p} is the probability distribution for \mathbf{x}_i conditional on the territory having received $X \geq 2$ propagules, where $X_i = X - 1$ of them are nonfocal, giving

$$\tilde{p}(\mathbf{x}_i) = \sum_{X=2}^{\infty} P(X|X \ge 2)p(\mathbf{x}_i|X_i = X - 1)$$
(16)

Here the total number of propagules X follows a Poisson distribution with mean L, and we have $P(X|X \ge 2) = P(X)/P(X \ge 2) = P(X)/(1-(1+L)e^{-L})$. Thus,

$$\langle x_j \rangle_{\tilde{p}} = \sum_{\mathbf{x}_i} \tilde{p}(\mathbf{x}_i) x_j$$

$$= \frac{1}{1 - (1 + L)e^{-L}} \sum_{X=2}^{\infty} P(X) \sum_{\mathbf{x}_i} p(\mathbf{x}_i | X_i = X - 1) x_j. \tag{17}$$

The inner sum over \mathbf{x}_i is the mean number of propagules of a given nonfocal type j that will be found in a territory which received X-1 nonfocal propagules in total, which is equal to $\frac{l_j}{L-l_i}(X-1)$. Thus,

$$\langle x_j \rangle_{\tilde{p}} = \frac{l_j}{1 - (1 + L)e^{-L}} \frac{1}{L - l_i} \sum_{X=2}^{\infty} P(X)(X - 1)$$

$$= \frac{l_j}{1 - (1 + L)e^{-L}} \frac{L - 1 + e^{-L}}{L - l_i},$$
(18)

where the last line follows from $\sum_{X=2}^{\infty} P(X)(X-1) = \sum_{X=1}^{\infty} P(X)(X-1) = \sum_{X=1}^{\infty} P(X)X - \sum_{X=1}^{\infty} P(X)$.

Having evaluated the mean propagule numbers, we now evaluate the variance in propagule numbers to check that the mean value replacement in Eq. (15) is justified. This is complicated because the x_j are not independent with respect to \tilde{p} . Here we use the following approximation to give some insight into the magnitude of these fluctuations and also the nature of the correlations between the x_j . We replace \tilde{p} with \tilde{q} , defined as the \mathbf{x}_i Poisson dispersal probabilities conditional on $X_i \geq 1$ (which are independent). The distinction between \tilde{p} and \tilde{q} will be discussed further below. The \tilde{q} approximation gives $\langle x_j \rangle_{\tilde{q}} = \langle x_j \rangle_p / C = l_j / C$, where $C = 1 - e^{-(L-l_i)}$, with variances and covariances given by

$$\sigma_{\tilde{q}}^{2}(x_{j}) = \langle x_{j}^{2} \rangle_{\tilde{q}} - \langle x_{j} \rangle_{\tilde{q}}^{2}$$

$$= \frac{1}{C} \langle x_{j}^{2} \rangle_{p} - \frac{l_{j}^{2}}{C^{2}}$$

$$= \frac{1}{C} (l_{j}^{2} + l_{j}) - \frac{l_{j}^{2}}{C^{2}}$$

$$= \frac{l_{j}^{2}}{C} \left(1 - \frac{1}{C} \right) + \frac{l_{j}}{C}, \tag{19}$$

and

$$\sigma_{\tilde{q}}(x_{j}, x_{k}) = \langle x_{j} x_{k} \rangle_{\tilde{q}} - \langle x_{j} \rangle_{\tilde{q}} \langle x_{k} \rangle_{\tilde{q}}$$

$$= \frac{1}{C} \langle x_{j} x_{k} \rangle_{p} - \frac{l_{j} l_{k}}{C^{2}}$$

$$= \frac{l_{j} l_{k}}{C} \left(1 - \frac{1}{C} \right) \qquad j \neq k.$$
(20)

The distinction between \tilde{p} and \tilde{q} is closely connected to why the exact dispersal probability distribution \tilde{p} is evaluated after dispersal. \tilde{p} assumes that one of the propagules present in a given territory after dispersal belongs to the focal type (a statement about propagule identity), whereas \tilde{q} assumes that there is a focal propagule present before non-focal dispersal commences (a statement about propagule presence). As a result, if the focal type is rare, \tilde{q} predicts that the mean propagule density is L+1 in territories with one focal propagule. This is erroneous, because the mean number of propagules per territory when lottery competition commences is L, regardless of whether a focal propagule happens to be

present. The discrepancy causes rare invaders evaluated under \tilde{q} to have an intrinsic rarity disadvantage, which is absurd (i.e. even if b, c and d are identical, \tilde{q} predicts that a rare type 544 will decline in frequency). By contrast, Eq. (18) correctly predicts that there are on average 545 $\sum_{j\neq i} \langle x_j \rangle_{\tilde{p}} \approx L-1$ nonfocal propagules because \tilde{p} accounts for potentially large negative 546 covariances between the x_i "after dispersal". By neglecting the latter covariances, \tilde{q} overes-547 timates the fluctuations in $\sum_{j\neq i} c_j x_j$; thus \tilde{q} gives an upper bound on the fluctuations. The 548 discrepancy between \tilde{q} and \tilde{p} will be largest when L is of order 1 or smaller, because then 549 the propagule assumed to already be present under \tilde{q} is comparable to, or greater than, the 550 entire propagule density. 551

Decomposing the variance in $\sum_{j\neq i} c_j x_j$,

552

562

$$\sigma_{\tilde{q}}^{2}(\sum_{j \neq i} c_{j} x_{j}) = \sum_{j \neq i} \left[c_{j}^{2} \sigma_{\tilde{q}}^{2}(x_{j}) + 2 \sum_{k > j, k \neq i} c_{j} c_{k} \sigma_{\tilde{q}}(x_{j}, x_{k}) \right], \tag{21}$$

and using the fact that $\sigma_{\tilde{q}}(x_j, x_k)$ and the first term in Eq. (19) are negative because C < 1,
we obtain an upper bound on the relative fluctuations in $\sum_{j \neq i} c_j x_j$,

$$\frac{\sigma(\sum_{j\neq i} c_j x_j)}{\langle \sum_{j\neq i} c_j x_j \rangle} = C^{1/2} \frac{\left(\sum_{j\neq i} c_j^2 l_j + (1 - 1/C) \left(\sum_{j\neq i} c_j l_j\right)^2\right)^{1/2}}{\sum_{j\neq i} c_j l_j} < C^{1/2} \frac{\left(\sum_{j\neq i} c_j^2 l_j\right)^{1/2}}{\sum_{j\neq i} c_j l_j}. (22)$$

Suppose that the c_j are all of similar magnitude (their ratios are of order one). Then Eq. (22) is $\ll 1$ for the case when $L - l_i \ll 1$ (due to the factor of $C^{1/2}$), and also for the case when at least some of the nonfocal propagule densities are large $l_j \gg 1$ (since it is then of order $1/\sqrt{L-l_i}$). The worst case scenario occurs when $L-l_i$ is of order one. Then Eq. (22) gives a relative error of approximately 50%, which from our earlier discussion we know to be a substantial overestimate when L is of order 1. Our numerical results (Fig. 3) confirm that the relative errors are indeed small.

However, the relative fluctuations in $\sum_{j\neq i} c_j x_j$ can be large if some of the c_j are much

larger than the others. Specifically, in the presence of a rare, extremely strong competitor $(c_j l_j \gg c_{j'} l_{j'})$ for all other nonfocal types j', and $l_j \ll 1$, then the RHS of Eq. (22) can be large and we cannot make the replacement Eq. (15).

Substituting Eqs. (15) and (18) into Eq. (14), we obtain

$$\Delta_r n_i \approx m_i R_i \frac{c_i}{\overline{c}},\tag{23}$$

where R_i is defined in Eq. (6).

568 Competition when abundant

The final contribution, $\Delta_a n_i$, accounts for territories where two or more focal propagules are present (a stands for "abundant"). Similarly to Eq. (14), we have

$$\Delta_a n_i = U(1 - (1 + l_i)e^{l_i}) \left\langle \frac{c_i x_i}{\sum_j c_j x_j} \right\rangle_{\hat{p}}$$
(24)

where \hat{p} is the probability distribution of both focal and nonfocal propagule abundances *after* dispersal in those territories where at least two focal propagules landed.

Again, we argue that the relative fluctuations in $\sum c_j x_j$ are much smaller than 1 (with respect to \hat{p}), so that,

$$\left\langle \frac{c_i x_i}{\sum_j c_j x_j} \right\rangle_{\hat{p}} \approx \frac{c_i \langle x_i \rangle_{\hat{p}}}{\sum_j c_j \langle x_j \rangle_{\hat{p}}}.$$
 (25)

Following a similar procedure as for $\Delta_r n_i$, where the vector of propagule abundances is

denoted \mathbf{x} , the mean focal type abundance is,

$$\langle x_i \rangle_{\hat{p}} = \sum_{\mathbf{x}} x_i p(\mathbf{x} | x_i \ge 2)$$

$$= \sum_{x_i} x_i p(x_i | x_i \ge 2)$$

$$= \frac{1}{1 - (1 + l_i)e^{-l_i}} \sum_{x_i \ge 2} p(x_i) x_i$$

$$= l_i \frac{1 - e^{-l_i}}{1 - (1 + l_i)e^{-l_i}}.$$
(26)

For nonfocal types $j \neq i$, we have

575

$$\langle x_{j} \rangle_{\hat{p}} = \sum_{\mathbf{x}} x_{j} p(\mathbf{x} | x_{i} \geq 2)$$

$$= \sum_{X} P(X | x_{i} \geq 2) \sum_{\mathbf{x}} x_{j} p(\mathbf{x} | x_{i} \geq 2, X)$$

$$= \sum_{X} P(X | x_{i} \geq 2) \sum_{\mathbf{x}_{i}} p(x_{i} | x_{i} \geq 2, X) \sum_{\mathbf{x}_{i}} x_{j} p(\mathbf{x}_{i} | X_{i} = X - x_{i})$$

$$= \sum_{X} P(X | x_{i} \geq 2) \sum_{x_{i}} p(x_{i} | x_{i} \geq 2, X) \frac{l_{j}(X - x_{i})}{L - l_{i}}$$

$$= \frac{l_{j}}{L - l_{i}} \left[\sum_{X} P(X | x_{i} \geq 2) X - \sum_{x_{i}} p(x_{i} | x_{i} \geq 2) x_{i} \right]$$

$$= \frac{l_{j}}{L - l_{i}} \left(L \frac{1 - e^{-L}}{1 - (1 + L)e^{-L}} - l_{i} \frac{1 - e^{-l_{i}}}{1 - (1 + l_{i})e^{-l_{i}}} \right). \tag{27}$$

and in going from 4 to 5 we have separately evaluated the contributions from the X and x_i terms in the numerator.

To calculate the standard deviation in $\sum_{j\neq i} c_j x_j$, we use a similar approximation as for $\Delta_r n_i$: \hat{p} is approximated by \hat{q} , defined as the \mathbf{x} dispersal probabilities in a territory

In going from line 3 to 4, we used the same logic used to evaluate the inner sum in Eq. (17),

conditional on $x_i > 2$ (that is, treating the x_j as independent). All covariances between

nonfocal types are now zero, so that $\sigma_{\hat{q}}^2(\sum c_j x_j) = \sum c_j^2 \sigma_{\hat{q}}^2(x_j)$, where $\sigma_{\hat{q}}^2(x_j) = l_j$ for $j \neq i$, and

$$\sigma_{\hat{q}}^{2}(x_{i}) = \frac{l_{i}}{D} \left(l_{i} + 1 - e^{-l_{i}} - \frac{l_{i}}{D} \left(1 - e^{-l_{i}} \right)^{2} \right), \tag{28}$$

where $D = 1 - (1 + l_i)e^{-l_i}$, and

$$\frac{\sigma_{\hat{q}}(\sum c_j x_j)}{\langle \sum c_j x_j \rangle} = \frac{\left(\sum_{j \neq i} c_j^2 l_j + c_i^2 \sigma_{\hat{q}}^2(x_i)\right)^{1/2}}{\sum_{j \neq i} c_j l_j + c_i l_i (1 - e^{-l_i})/D}.$$
 (29)

Similarly to Eq. (22), the RHS of Eq. (29) is $\ll 1$ for the case that $L \ll 1$ (due to a factor of $D^{1/2}$), and also for the case when at least some of the propagule densities (focal or nonfocal) are large — provided that c_i and the c_j are all of similar magnitude. Again, the worst case scenario occurs when l_i and $L - l_i$ are of order 1, in which case Eq. (29) is around 35%, which is again where the \hat{q} approximation produces the biggest overestimate of the fluctuations in \mathbf{x} . Similarly to Eq. (22), the RHS of (29) will not be $\ll 1$ in the presence of a rare, extremely strong competitor.

Combining Eqs. (24) and (25), we obtain

$$\Delta_a n_i = m_i A_i \frac{c_i}{\overline{c}},\tag{30}$$

where A_i is defined in Eq. (6).

Appendix C: Total density in the Lotka-Volterra competition model

Here we show that under the Lotka-Volterra model of competition, total density N does not in general remain constant over a selective sweep in a crowded population even if the types

have the same saturation density (for a related discussion on the density- and frequencydependence of selection in the Lotka-Volterra model, see (Smouse, 1976; Mallet, 2012)).

We assume equal effects of crowding within types $\alpha_{11} = \alpha_{22} = \alpha_{\text{intra}}$ and $N = 1/\alpha_{\text{intra}}$ and check whether it is then possible for $\frac{dN}{dt}$ to be zero in the sweep $(n_1, n_2 \neq 0)$. Substituting these conditions into Eq. (9), we obtain

$$\frac{dn_1}{dt} = r_1(\alpha_{11} - \alpha_{12})n_1n_2
\frac{dn_2}{dt} = r_2(\alpha_{22} - \alpha_{21})n_1n_2$$
(31)

Adding these together, $\frac{dN}{dt}$ can only be zero if

$$r_1(\alpha_{\text{intra}} - \alpha_{12}) + r_2(\alpha_{\text{intra}} - \alpha_{21}) = 0.$$
 (32)

To get some intuition for Eq. (32), suppose that a mutant arises with improved competitive ability but identical intrinsic growth rate and saturation density $(r_1 = r_2 \text{ and } \alpha_{11} = \alpha_{22})$. This could represent a mutation to an interference competition trait, for example (Gill, 1974). Then, according the above condition, for N to remain constant over the sweep, the mutant must find the wildtype more tolerable than itself by exactly the same amount that the wildtype finds the mutant less tolerable than itself.

Even if we persuaded ourselves that this balance of inter-type interactions is plausible in some circumstances, when multiple types are present the requirement for constant N becomes

$$\sum_{ij} r_i (\alpha_{\text{intra}} - \alpha_{ij}) p_i p_j = 0, \tag{33}$$

which depends on frequency and thus cannot be satisfied in general for constant inter-type coefficients α_{ij} . We conclude that selection in the Lotka-Volterra competition model will generally involve non-constant N.

Appendix D: Density-dependence of b-selection

In section "Density-regulating traits and the threat of strong selection" we argued that the density-dependent factor $f(\bar{b}, N)$ is unchanged at the beginning and end points of an equilibrium-to-equilibrium b. Here we estimate the magnitude of the deviation in $f(\bar{b}, N)$ during the sweep.

For simplicity, we introduce the notation D=N/T and assume that D is small. We can thus make the approximation $1-e^{-\bar{b}D}\approx \bar{b}D$ and $f(\bar{b},N)\approx \bar{b}(1-D)$. We expect this to be a conservative approximate based on the worst case scenario, because N is most sensitive to an increase in b in this low-density linear regime. We first calculate the value of $f(\bar{b},N)$ at the halfway point in a sweep, where the halfway point is estimated with simple linear averages for b and b. The sweep is driven by a b variant with $b_j=b_i(1+\epsilon)$, and we denote the corresponding initial and final densities by b and b respectively, where we have b and b in this low-density linear regime.

$$f_{\text{half}} = f(\frac{b_i + b_j}{2}, \frac{N_i + N_j}{2}) = \frac{b_i + b_j}{2} \left(1 - \frac{D_i + D_j}{2} \right)$$

$$= \frac{1}{4} (b_i + b_j)(2 - D_i - D_j)$$

$$= \frac{1}{4} (2d_i + b_i(1 - D_j) + b_j(1 - N_i)). \tag{34}$$

Dividing by d_i , the proportional deviation in f(N) at the midpoint of the sweep is

$$\frac{f_{\text{half}}}{d_i} = \frac{1}{4} \left(2 + \frac{b_i}{b_j} + \frac{b_j}{b_i} \right)
= \frac{1}{4} \left(2 + \frac{1}{1+\epsilon} + 1 + \epsilon \right)
= 1 + \frac{1}{4} (\epsilon^2 - \epsilon^3 + \dots),$$
(35)

where we have used the Taylor expansion $\frac{1}{1+\epsilon} = 1 - \epsilon + \epsilon^2 - \epsilon^3 + \dots$

By contrast, for a δ sweep in Eq. (10), the density-dependent term N increases by a factor of $\frac{1}{1-\epsilon} = 1 + \epsilon + \epsilon^2 + \ldots$ Thus, the deviations in f(N) are an order of magnitude smaller than those shown in Fig. (5), and even proportional changes of order $\epsilon = 0.1$ will cause a negligible deviation from the canonical selection equation.