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

[I'm going to revise this after your next round of comments] Selection is commonly described by assigning relative fitness values to genotypes. Yet when selection is strong, the ecological view of selection in density-regulated populations seems to be incompatible with constant-density relative fitnesses. Here we analyze the population ecological limits of relative fitness using a novel of density-dependent selection which contains a "reproductive excess. Our model clearly distinguishes between density-dependent selection and changes in density driven by selection. These two effects are confounded in standard models of 10 density-regulated population growth, but both are necessary, in combination with strong 11 selection, for relative fitness to break down in populations close to demographic equilibrium. Remarkably, both effects are not sufficient: we give an example of strong selection 13 on a density-regulating trait subject to density-dependent selection that conforms to the density-independent relative fitness description almost exactly. We reiterate the importance 15 of reproductive excesses in many species, which allows even strong selection to have no effect 16 on density. Our model also offers a possible alternative to relative fitness when the latter is 17 untenable, as is likely the case far from demographic equilibrium. (191 words)

# 20 Introduction

There are a variety of different measures of fitness, such as expected lifetime reproductive 21 ratio  $R_0$ , intrinsic population growth rate r, equilibrium population density/carrying capac-22 ity (often labeled "K") (Benton and Grant, 2000), and invasion fitness (Metz et al., 1992). 23 In addition, "relative fitness" is widely used in evolutionary genetics, where the focus is on relative genotypic frequencies (Barton et al., 2007, pp. 468). The variety of fitness mea-25 sures is not problematic in itself, but it should be clear how these measures are connected to the processes of birth and death which ultimately drive selection (Metcalf and Pavard 2007; Doebeli et al. 2017; Charlesworth 1994, pp. 178). While such a connection is clear 28 for absolute fitness measures like r or  $R_0$ , relative fitness has only weak justification from 29 population ecology. It has even been proposed that relative fitness be justified from measure 30 theory, abandoning population biology altogether (Wagner, 2010). Given the widespread use 31 of relative fitness in evolutionary genetics, it is important to understand its population eco-32 logical basis, both to clarify its domain of applicability, and as part of the broader challenge 33 of synthesizing ecology and evolution. 34 For haploids tracked in discrete time, the change in the abundance  $n_i$  of type i over a 35 time step can be expressed as  $\Delta n_i = (W_i - 1)n_i$  where  $W_i$  is "absolute fitness" (i.e. the 36 abundance after one time step is  $n'_i = W_i n_i$ ). The corresponding change in frequency is 37  $\Delta p_i = \left(\frac{W_i}{\overline{W}} - 1\right) p_i$ , where  $\overline{W} = \sum_i W_i p_i$ . In continuous time, the Malthusian parameter  $r_i$ replaces  $W_i$  and we have  $\frac{dn_i}{dt} = r_i n_i$  and  $\frac{dp_i}{dt} = (r_i - \overline{r})p_i$  (Crow et al., 1970). Note that we can replace the  $W_i$  with any set of values proportional to the  $W_i$  without affecting the ratio  $W_i/\overline{W}$  or  $\Delta p_i$ . These "relative fitness" values tell us how type frequencies change, but give no information about the dynamics of total population density  $N = \sum_{i} n_{i}$  (Barton et al., 2007, pp. 468). Similarly in the continuous case, adding an arbitrary constant to the Malthusian parameters  $r_i$  has no effect on  $\frac{dp_i}{dt}$  (these would then be relative log fitnesses).

Relative fitness is often parameterized in terms of selection coefficients which represent the advantages of different types relative to each other. For instance, in continuous time  $s = r_i - r_j$  is the selection coefficient of type i relative to type j. Assuming that only i and j are present, the change in frequency can be written as

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

Thus, if  $r_i$  and  $r_j$  are constant, the frequency of the i type will grow logistically with a constant rate parameter s. We then say that selection is independent of frequency and density. The discrete time case is more complicated. Defining the selection coefficient by  $W_i = (1+s)W_j$ , and again assuming i and j are the only types present, we have

Hence, even in the simplest case that  $W_i$  and  $W_j$  are constant, selection is frequency-

dependent in discrete time (note that this frequency dependence is negligible when s is

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$$\Delta p_i = \frac{W_i - W_j}{\overline{W}} p_i (1 - p_i) = \frac{s}{1 + sp_i} p_i (1 - p_i). \tag{2}$$

small compared to 1; see Frank 2011). We will refer to both the continuous and discrete time selection equations (1) and (2) throughout this paper, but the simpler continuous time case will be our point of comparison for the rest of this section.

In a constant environment, and in the absence of crowding,  $r_i$  is a constant "intrinsic" population growth rate. The interpretation of Eq. (1) is then simple: the selection coefficient s is simply the difference in intrinsic growth rates. However, growth cannot continue at a non-zero constant rate indefinitely: the population is not viable if  $r_i < 0$ , whereas  $r_i > 0$  implies endlessly increasing population density. Thus, setting aside unviable populations, the increase in population density must be checked by crowding. This implies that the Malthusian parameters  $r_i$  eventually decline to zero (e.g. Begon et al. 1990, pp. 203). Selection can then be density-dependent, and indeed this is probably not uncommon, because

crowded and uncrowded conditions can favor very different traits (Travis et al., 2013). Eq. (1) is then not a complete description of selection — it lacks an additional coupled equation 67 describing the dynamics of N, on which s in Eq. (1) now depends. In general we cannot simply specify the dynamics of N independently, because those ecological dynamics are 69 coupled with the evolutionary dynamics of type frequency (Travis et al., 2013). Thus, in the 70 presence of density-dependent selection, the simple procedure of assigning constant relative 71 fitness values to different types has to be replaced with an ecological description of absolute 72 growth rates. Note that frequency-dependent selection does not raise a similar problem, 73 because a complete description of selection still only requires us to model the type frequencies, not the ecological variable N as well. 75

In practice, many population genetic models simply ignore density dependence and assign
a constant relative fitness to each type. Selection is typically interpreted as operating through
viability, but the ecological processes underlying the regulation of population density are
frequently left unspecified (e.g. Gillespie 2010; Nagylaki et al. 1992; Ewens 2004). Density
either does not enter the model at all, or if finite-population size effects ("random genetic
drift") are important, then N is assumed to have reached some fixed equilibrium value
(Fig. 1b).

A rather different picture emerges in more ecologically explicit studies of selection in density-regulated populations. Following Fisher's suggestion that evolution tends to increase density in the long term (Fisher, 1930; Leon and Charlesworth, 1978; Lande et al., 2009), as well as the influential concept of K-selection (specifically, the idea that selection in crowded conditions favors greater equilibrium density; MacArthur 1962), many studies of density-regulated growth have focused on the response of density to selection (Kostitzin, 1939; MacArthur and Wilson, 1967; Roughgarden, 1979; Christiansen, 2004). Indeed, both N and s change during, and as a result of, adaptive sweeps in many of the most widely used models of density-regulated population growth. The latter includes simple birth-death

(Kostitzin, 1939) and logistic models (Fig. 1a; MacArthur 1962; Roughgarden 1979; Boyce 1984), variants of these models using other functional forms for the absolute fitness penalties of crowding (Kimura, 1978; Charlesworth, 1971; Lande et al., 2009; Nagylaki, 1979; Lande et al., 2009), and the "R\* rule" of resource competition theory (which states that the type able to deplete a shared limiting consumable resource to the lowest equilibrium density R\* excludes the others; Grover 1997). Density also changes in response to selection in the Lotka-Volterra competition model, at least during a sweep (except in special cases; Gill 1974; Smouse 1976; Mallet 2012).

The constant-N, constant-s description of selection also precludes consideration of longer-100 term aspects of the interplay between evolution and ecology such as population extinction. 101 A variety of approaches have been developed to address this in quantitative genetics (Burger 102 and Lynch, 1995; Engen et al., 2013), population genetics (Bertram et al., 2017) and adaptive 103 dynamics (Ferriere and Legendre, 2013; Dieckmann and Ferrière, 2004). Although density-104 dependent selection is pertinent to this longer-term issue, our focus here is the description of 105 the time-dependent process by which selection changes allele frequencies. This is particularly 106 critical for making sense of evolution at the genetic level, for which we now have abundant 107 data. 108

In light of the complications arising from density-dependence, the assignment of densityindependent relative fitnesses has been justified as an approximation that holds when selection is weak and N changes slowly (Kimura and Crow 1969; Ewens 2004, pp. 277;
Charlesworth 1994, Chap. 4). Under these conditions, s is approximately constant in Eq. (1),
at least for some number of generations. If s depends only on density, not frequency, this
approximate constancy can hold over entire selective sweeps (Otto and Day, 2011).

However, the preceding arguments do not imply that the constant relative fitness idealization of population genetics *only* applies when selection is weak and N is stable (or when
selection is actually density-independent). The idealization of assigning relative fitness val-

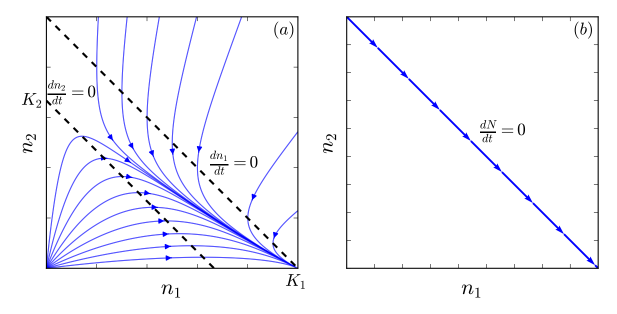


Figure 1: Phase diagram for the densities of two types  $n_1$  and  $n_2$  undergoing selection. (a) The logistic model  $\frac{dn_1}{dt} = r_1(1 - \frac{n_1 + n_2}{K_1})n_1$  and  $\frac{dn_2}{dt} = r_2(1 - \frac{n_1 + n_2}{K_2})n_1$  with  $r_1 = r_2$  and  $K_1 > K_2$ . (b) The constant-N, relative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

ues to genotypes is powerful, and so it is important to understand the specifics of when and how it succeeds or fails when selection is not weak, or N is not stable. For instance, in wild Drosophila, strong seasonally-alternating selection happens concurrently with large "boom-bust" density cycles (Messer et al., 2016; Bergland et al., 2014). Are we compelled to switch to a more ecologically-detailed model of selection based on Malthusian parameters or birth/death rates in this important model system? And if we make this switch, how much ecological detail do we need?

Here we argue that the simplified models of density-regulated growth mentioned above are misleading in their representation of the interplay between selection and density. This ultimately derives from their failure to account for "reproductive excess", that is, an excess of juveniles that experience stronger selection than their adult counterparts (Turner and Williamson, 1968). By allowing selection to be concentrated at a juvenile "bottleneck", reproductive excess makes it possible for the density of adults to remain constant even

under strong selection. Reproductive excess featured prominently in early debates about the regulation of population density (e.g. Nicholson 1954), and also has a long history in 132 evolutionary theory, particularly related to Haldane's "cost of selection" (Haldane, 1957; 133 Turner and Williamson, 1968). Additionally, reproductive excess is implicit in foundational 134 evolutionary-genetic models like the Wright-Fisher, where each generation involves the pro-135 duction of an infinite number of zygotes, of which a constant number N are sampled to form 136 the next generation of adults. Likewise in the Moran model, a juvenile is always available to 137 replace a dead adult every iteration no matter how rapidly adults are dying, and as a result 138 N remains constant. 139

Nevertheless, studies of density-dependent selection rarely incorporate reproductive excess. This requires that we model a finite, density-dependent excess, which is substantially
more complicated than modeling either zero (e.g. logistic) or infinite (e.g. Wright-Fisher)
reproductive excess. Nei's "competitive selection" model incorporated a finite reproductive
excess to help clarify the "cost of selection" (Nei, 1971; Nagylaki et al., 1992), but used an
unusual representation of competition based on pairwise interactions defined for at most two
different genotypes, and was also restricted to equal fertilities for each genotype.

In models with detailed age structure, it is usually assumed that the density of a "critical age group" mediates the population's response to crowding (Charlesworth, 1994, pp. 54). Reproductive excess is a special case corresponding to a critical pre-reproductive age 149 group. A central result of the theory of density-regulated age-structured populations is that 150 selection proceeds in the direction of increasing equilibrium density in the critical age group 151 (Charlesworth, 1994, pp. 148). This is a form of the classical K-selection ideas discussed 152 above, but restricted to the critical age group (juveniles, in this case). The interdepen-153 dence of pre-reproductive selection and reproductive density is thus overlooked as a result 154 of focusing on density in the critical age group. 155

We re-evaluate the validity of the constant relative fitness description of selection in a

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novel model of density-regulated population growth that has a finite reproductive excess. Our model is inspired by the classic discrete-time lottery model, which was developed by 158 ecologists to study competition driven by territorial contests in reef fishes and plants (Sale, 159 1977; Chesson and Warner, 1981), and which has some similarities to the Wright-Fisher 160 model (Svardal et al., 2015). Each type is assumed to have three traits: fecundity b, mortality 161 d, and competitive ability c. In each iteration of the classic lottery model, each type produces 162 a large number of juveniles, such that N remains constant (infinite reproductive excess). 163 Competitive ability c affects the probability of winning a territory, and behaves like a pure 164 relative fitness trait. Thus, fitness involves a product of fertility and juvenile viability akin 165 to standard population genetic models of selection (e.g. Crow et al. 1970, pp. 185). We 166 relax the large-juvenile-number assumption of the lottery model to derive a variable-density 167 lottery with a finite, density-dependent reproductive excess. 168

The properties of density-dependent selection in our model are strikingly different from the classical literature discussed above. The strong connection between crowding and selection for greater equilibrium density is broken: selection need not affect density at all. And when it does, the density-independent discrete-time selection equation (2) is almost exact even for strong selection, provided that any changes in density are driven only by selection (as opposed to large deviations from demographic equilibrium), and that selection occurs on only one of the traits b, c, or d. On the flip side, the constant relative fitness approximation fails when strong selection acts concurrently on two or more of these traits, or when the population is far from demographic equilibrium.

# $_{^{178}}$ Model

### 179 Assumptions and definitions

We restrict our attention to asexual haploids, since it is then clearer how the properties 180 of selection are tied to the underlying population ecological assumptions. We assume that 181 reproductively mature individuals ("adults") require their own territory to survive and re-182 produce. All territories are identical, and the total number of territories is T. Time advances 183 in discrete iterations, each representing the time from birth to reproductive maturity. In a 184 given iteration, the number of adults of the i'th type will be denoted by  $n_i$ , the total number 185 of adults by  $N = \sum_{i} n_{i}$ , and the number of unoccupied territories by U = T - N. We assume 186 that the  $n_i$  are large enough that stochastic fluctuations in the  $n_i$  (drift) can be ignored (with 187 T also assumed large to allow for low type densities  $n_i/T \ll 1$ ). 188

Each iteration, adults produce propagules which disperse at random, independently of 189 distance from their parents, and independently of each other. We assume that each adult 190 from type i produces  $b_i$  propagules on average, so that the mean number of i propagules 191 dispersing to unoccupied territories is  $m_i = b_i n_i U/T$ . The parameter  $b_i$  can be thought of as 192 a measure of "colonization ability", which combines fertility and dispersal ability (Levins and 193 Culver, 1971; Tilman, 1994). Random dispersal is then modeled using a Poisson distribution 194  $p_i(x_i) = l_i^{x_i} e^{-l_i}/x_i!$  for the number  $x_i$  of i propagates dispersing to any particular unoccupied 195 territory, where  $l_i = m_i/U$  is the mean propagule density in unoccupied territories. The 196 total propagule density will be denoted  $L = \sum_{i} l_{i}$ . 197

We assume that adults cannot be ousted by juveniles, so that recruitment to adulthood occurs exclusively in unoccupied territories. When multiple propagules land on the same unoccupied territory, the winner is determined by lottery competition: type i wins a territory with probability  $c_i x_i / \sum_i c_i x_i$ , where  $c_i$  is a constant representing relative competitive ability (Fig. 2). Since the expected fraction of unoccupied territories with propagule composition

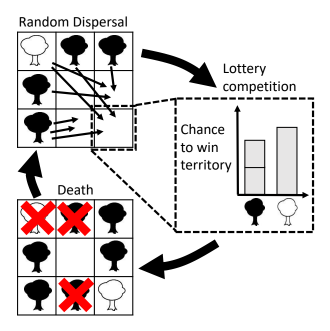


Figure 2: One iteration of our model. Propagules are dispersed by adults at random (only those propagules landing on unoccupied territories are shown). Some territories may receive zero propagules. Lottery competition then occurs in each territory that receives more than one propagule (only illustrated in one territory). In a given territory, type i has probability proportional to  $c_i x_i$  of winning the territory, where  $c_i$  measures competitive ability and  $x_i$  is the number of i propagules present. In the illustrated territory, more black propagules are present, but white is a stronger competitor and has a higher probability of winning. Adult deaths make new territories available for the next iteration (red crosses).

 $x_1, \ldots, x_G$  is  $p_1(x_1) \cdots p_G(x_G)$  where G is the number of types present, and type i is expected to win a proportion  $c_i x_i / \sum_i c_i x_i$  of these, type i's expected territorial acquisition is given by

$$\Delta_{+} n_{i} = U \sum_{x_{1}, \dots, x_{G}} \frac{c_{i} x_{i}}{\sum_{i} c_{i} x_{i}} p_{1}(x_{1}) \cdots p_{G}(x_{G}).$$
(3)

Here the sum only includes territories with at least one propagule present. Since we do not consider random genetic drift here, we will not analyze the fluctuations around these two expectations.

Adult mortality occurs after lottery recruitment at a constant, type-specific per-capita rate  $d_i \geq 1$ , and can affect adults recruited in the current iteration, such that the new abundance at the end of the iteration is  $(n_i + \Delta_+ n_i)/d_i$  (Fig. 2). In terms of absolute fitness, this can be written as

$$W_i = \frac{1}{d_i} \left( 1 + \frac{\Delta_+ n_i}{n_i} \right). \tag{4}$$

Here  $\frac{\Delta_+ n_i}{n_i}$  is the per-capita rate of territorial acquisition, and  $1/d_i$  is the fraction of type i adults surviving to the next iteration.

# 214 Connection to the classic lottery model

In the classic lottery model (Chesson and Warner, 1981), unoccupied territories are assumed to be saturated with propagules from every type  $(l_i \to \infty \text{ for all } i)$ . From the law of large numbers, the composition of propagules in each territory will not deviate appreciably from the mean composition  $l_1, l_2, \ldots, l_G$ . Type i is thus expected to win a proportion  $c_i l_i / \sum_i c_i l_i$ of the U available territories,

$$\Delta_{+} n_{i} = \frac{c_{i} l_{i}}{\sum_{i} c_{i} l_{i}} U = \frac{c_{i} l_{i}}{\overline{c} L} U, \tag{5}$$

where  $\bar{c} = \sum_i c_i m_i / \sum_i m_i$  is the mean competitive ability for a randomly selected propagule.

Note that all unoccupied territories are filled in a single iteration of the classic lottery model,

whereas our more general model Eq. (3) allows for territories to be left unoccupied and hence

also accommodates low propagule densities.

# ${f Results}$

### $_{\scriptscriptstyle{225}}$ Analytical approximation of the variable-density lottery

Here we evaluate the expectation in Eq. (3) to better understand the dynamics of density-226 dependent lottery competition. Similarly to the classic lottery model, we replace the  $x_i$ , 227 which take different values in different territories, with "effective" mean values. However, 228 since we want to allow for low propagule densities, we cannot simply replace the  $x_i$  with 229 the means  $l_i$  as in the classic lottery. For a low density type, growth comes almost entirely 230 from territories with  $x_i = 1$ , for which its mean density  $l_i \ll 1$  is not representative. We 231 therefore separate Eq. (3) into  $x_i = 1$  and  $x_i > 1$  components, taking care to ensure that the 232 effective mean approximations for these components are consistent with each other (details 233 in Appendix B). The resulting variable-density 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 235 types). We obtain 236

$$\Delta_{+} n_{i} \approx \left[ e^{-L} + (R_{i} + A_{i}) \frac{c_{i}}{\overline{c}} \right] l_{i} U, \tag{6}$$

237 where

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

238 and

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

Comparing Eq. (6) to Eq. (5), the classic lottery per-propagate success rate  $c_i/\bar{c}L$  has 239 been replaced by three separate terms. The first,  $e^{-L}$ , accounts for propagules which land 240 alone on unoccupied territories; these propagules secure the territories without contest. The 241 second,  $R_i c_i/\bar{c}$ , represents competitive victories on territories where only a single i propagule 242 lands, together with at least one other propagule from a different type (this term dominates 243 the growth of a rare invader in a high density population and determines invasion fitness). 244 The third term,  $A_i c_i/\bar{c}$ , represents competitive victories in territories where two or more i 245 type propagules are present. The relative importance of these three terms varies with both 246 the overall propagule density L and the relative propagule frequencies  $l_i/L$ . If  $l_i \gg 1$  for all 247 types, we recover the classic lottery model (only the  $A_i c_i/\bar{c}$  term remains, and  $A_i \to 1/L$ ). 248 Fig. 3 shows that Eq. (6) and its components closely approximate simulations of our 249 variable-density lottery model over a wide range of propagule densities. Two types are 250 present, one of which is at low frequency. The growth of the low-frequency type relies 251 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 252 for the high-frequency type, which depends instead on high density territorial victories. Fig. 3 253 also shows the breakdown of the classic lottery model at low propagule densities. 254

In the special case that all types are competitively equivalent (identical  $c_i$ ), Eq. (6) takes a simpler form,

$$\Delta_{+}n_{i} = \frac{l_{i}}{L}(1 - e^{-L})U. \tag{7}$$

This formula can also be deduced directly from Eq. (3):  $1 - e^{-L}$  is the fraction of territories that receive at least one propagule under Poisson dispersal,  $(1 - e^{-L})U$  is the total number of such territories, and type i is expected to receive a fraction  $l_i/L$  of these. Total population density thus grows according to

$$\Delta N = (1 - e^{-L})U - \sum_{i} d_i n_i \tag{8}$$

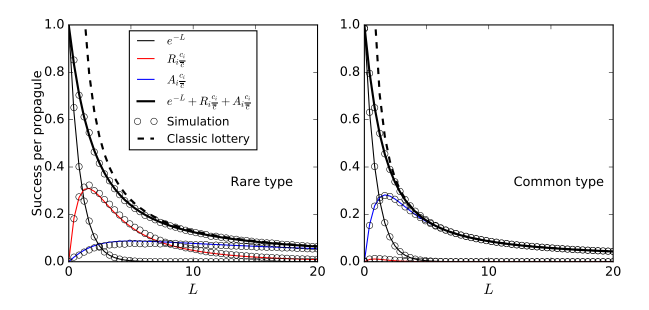


Figure 3: Comparison of Eq. (6), the classic lottery model, and simulations. The vertical axis is per-propagule success rate for all propagules  $\Delta_+ n_i/m_i$ , and for the three separate components in Eq. (6). Two types are present with  $c_1 = 1$ ,  $c_2 = 1.5$  and  $l_2/l_1 = 0.1$ . Simulations are conducted as follows:  $x_1, x_2$  values are sampled  $U = 10^5$  times from Poisson distributions with respective means  $l_1, l_2$ , and the victorious type in each territory is then decided by random sampling weighted by the lottery win probabilities  $c_i x_i/(c_1 x_1 + c_2 x_2)$ . Dashed lines show the failure of the classic lottery model at low density.

#### Density-dependent selection in the variable-density lottery

We now outline the basic properties of selection on b, c and d. The birth and mortality rates b and d are the traits which regulate density; b controls the fraction of unoccupied territories
that are contested, while d controls adult mortality. Competitive ability c does not regulate
density since it only affects the relative likelihood for each type to win a contested territory.
Thus, selection between types which only differ in c occurs without causing N to change
(Eq. (8) shows this formally).

Selection in our variable density lottery model is density-dependent. Since this a discrete 268 time model, this means that when two types are present, the selection factor  $(W_i - W_j)/\overline{W}$ 269 in Eq. (2) depends on N. Note that density-dependent selection is sometimes taken to mean 270 a qualitative change in which types are fitter than others at different densities (Travis et al., 271 2013). While reversal in the order of fitnesses and co-existence driven by density-regulation 272 are possible in our variable-density lottery (a special case of the competition-colonization 273 trade-off; Levins and Culver 1971; Tilman 1994; Bolker and Pacala 1999), questions related 274 to co-existence are tangential to our aims and will not be pursued further here. 275

Selection on c is density-dependent, with the strength of selection peaking at an intermediate density (Fig. 4). This intermediate peak occurs because at low density most territories
are claimed without contest, whereas at high density few unoccupied territories are available
to be contested. To see how selection on b and d depend on density, we write Eq. (7) in the
alternative form

$$\frac{\Delta_{+}n_{i}}{n_{i}} = \frac{b_{i}}{\overline{b}} \frac{1 - e^{-\overline{b}N/T}}{N} (T - N), \tag{9}$$

where we have used that fact that  $L = \bar{b}N/T$ , and  $\bar{b}$  is the population mean b.

It is clear *d*-selection is independent of density. On the other hand, the strength of *b*selection declines with density because the advantage of having greater *b* gets smaller the
fewer territories there are to be claimed (Fig. 4).

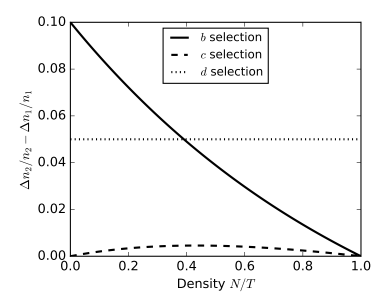


Figure 4: The density-dependence of selection in our variable-density lottery between an adaptive variant 2 and a wildtype 1 with equal frequencies. Here  $b_1 = 1$ ,  $d_1 = 0.5$  and  $c_1 = 1$ . For b-selection we set  $b_2 = b_1(1 + \epsilon)$ , and similarly for c and d, with  $\epsilon = 0.1$ . 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 wildtype equilibrium density is  $n_1/T \approx 0.4$ , suggesting that this population would rarely experience declining strength of c-selection with increasing N.

#### The response of density to selection; c-selection versus K-selection

We now turn to the issue of how density responds to selection, comparing our variable-density lottery to previous studies of density-regulated populations (Prout, 1980). As we saw in the previous section, c-selection has no effect on population density in our variable-density lottery. To make sense of how c-selection fits with previous population growth models, we now revisit MacArthur's general treatment of K-selection (MacArthur and Wilson, 1967).

MacArthur considered a population with two types that have densities  $n_1$  and  $n_2$  subject to density-dependent growth,

$$\frac{dn_1}{dt} = f_1(n_1, n_2) \qquad \frac{dn_2}{dt} = f_2(n_1, n_2). \tag{10}$$

The environment is assumed to remain constant apart from changing type densities. The functions  $f_1$  and  $f_2$  must decline to zero if  $n_1$  or  $n_2$  are sufficiently large, because the resources required for growth are limited. This defines 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. Thus, 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).

MacArthur used "K" to label the four intersection points of the nullclines with the axes, specifically  $f_1(K_{11}, 0) = 0$ ,  $f_1(0, K_{12}) = 0$ ,  $f_2(K_{21}, 0) = 0$  and  $f_2(0, K_{22}) = 0$ . 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 equilibrium densities akin to the K parameter in the logistic model (Fig. 1a). The other intersection points,  $K_{12}$  and  $K_{21}$ , are related to competition between types. To be more concrete, in the

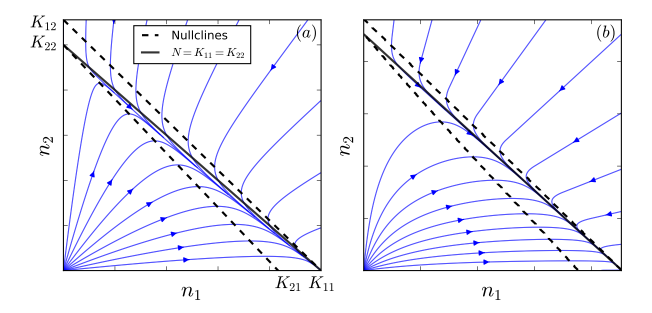


Figure 5: Selection between types with identical equilibrium density but different inter-type competitive ability. (a) Lotka-Volterra competition (Eq. 11) with  $r_1 = r_2 = 1$ ,  $\alpha_{11} = \alpha_{22} = 1$ ,  $\alpha_{12} = 0.9$  and  $\alpha_{21} = 1.2$ . Trajectories do not follow the line  $N = K_{11} = K_{22}$ . (b) Lottery competition (Eq. 6) with  $b_1 = b_2 = 5$ ,  $d_1 = d_2 = 0.1$  and  $c_1/c_2 = 5$ . Trajectories converge on the line  $N = K_{11} = K_{22}$ .

Lotka-Volterra competition model we have

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

$$f_2(n_1, n_2) = r_2(1 - \alpha_{22}n_1 - \alpha_{21}n_2)n_2$$
(11)

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. Hence, "fitness is K" in crowded populations (MacArthur and Wilson, 1967, pp. 149) in the sense that selection either favors the ability to keep growing at ever higher densities (moving a type's own nullcline outwards), or the ability to suppress the growth of competitors at lower densities (moving the nullcline of competitors inwards). This general idea is much broader than selection for greater equilibrium density (Gill, 1974).

Compared to simple birth-death models (Kostitzin, 1939) or variants of the logistic 308 (Roughgarden, 1979), the Lotka-Volterra model clearly distinguishes between intra- and 309 inter-type competitive effects. Thus, when selection acts on inter-type competitive effects, 310 one type can displace another without having a greater equilibrium density (Fig. 5a). This 311 has been termed " $\alpha$ -selection" to distinguish it from K-selection, which involves intra-type 312 competitive effects and changes in equilibrium density Gill (1974); Joshi et al. (2001). Al-313 though the initial and final densities of an  $\alpha$ -selection sweep are the same, density neverthe-314 less does change transiently in the Lotka-Volterra model (constant density only occurs for 315 a highly restricted subset of r and  $\alpha$  values; further details in Appendix C; also see Mallet 316 2012; Smouse 1976). Intuitively, for one type to exclude the other, competitive suppression 317 of growth between types must be stronger than competitive suppression of growth within 318 types, causing N to dip over a sweep (Fig. 5a). 319

By contrast, density trajectories for c-selection in our variable-density lottery converge
on a line of constant equilibrium density (Fig. 5b). This means that once the population
reaches demographic equilibrium, it behaves indistinguishably from a constant-N relative
fitness model (Fig. 1b). This complete uncoupling of density from c-selection arises due to
the presence of an excess of propagules which pay the cost of selection without affecting adult
density (Nei, 1971). As a result, Eq. (??) holds in demographic equilibrium even though
c-selection is density-dependent.

# 327 Density-regulating traits and the threat of strong selection

For Eq. (??) to break down, the selection coefficient s must depend on density. As shown in Fig. 4, this is not the case for d; the selection coefficient for a d-variant with  $d_2 = d_1(1-\epsilon)$  is a constant  $s = \epsilon d_1$ . Eq. (??) also holds when the population is at demographic equilibrium and density is unaffected by the outcome of selection; as discussed in the previous section, the latter is the case for c-selection in our model. Thus, to threaten Eq. (??), we require

selection to be density-dependent, and also density to be changing. This can obviously occur if density-dependent selection is occurring in a population far from demographic equilibrium. In this case the validity of Eq. (??) depends on the specifics of the rate and magnitude of demographic change (we return to this in the Discussion). However, Eq. (??) can be threatened even in demographically-stable populations if a density-regulating trait is subject to density-dependent selection, as is the case for b in our variable-density lottery.

Before we discuss the b trait, it is helpful to summarize the threat to Eq. (??) in simpler models of density-regulated growth, as exemplified by the birth-death model (Kostitzin, 1939)

$$\frac{dn_i}{dt} = (b_i - \delta_i N)n_i. \tag{12}$$

Here  $\delta_i$  is per-capita mortality due to crowding (for simplicity, there are no deaths when uncrowded). Starting from a type 1 population in equilibrium, a variant with  $\delta_2 = \delta_1(1 - \epsilon)$  has density-dependent selection coefficient  $s = \epsilon \delta_1 N$  in Eq. (??), which will change over the course of the sweep as N shifts from its initial type 1 equilibrium to a type 2 equilibrium. From Eq. (12), the equilibrium densities at the beginning and end of the sweep are  $N_{\text{initial}} = b_1/\delta_1$  and  $N_{\text{final}} = b_1/(\delta_1(1 - \epsilon)) = N_{\text{initial}}/(1 - \epsilon)$  respectively, and so  $s_{\text{initial}} = \epsilon b_1$  and  $s_{\text{final}} = s_{\text{initial}}/(1 - \epsilon)$ . Consequently, substantial deviations from Eq. (??) occur if there is sufficiently strong selection on  $\delta$  (Fig. 6; Kimura and Crow 1969; Crow et al. 1970).

Equilibrium-to-equilibrium b-sweeps in our variable-density lottery are qualitatively different from  $\delta$  sweeps in this simpler birth-death model, because greater b not only means more propagules contesting territories, but also more territories being contested. Together, the net density-dependent effect on b-selection coefficients is zero; in Eq. (), since  $b_i/\bar{b}=1$ in a single-type equilibrium, the density-dependence factor  $f(\bar{b},N)=\frac{1-e^{-\bar{b}N/T}}{N}(T-N)$  is exactly equal to the constant mortality rate at the beginning and end of a b-sweep, even though b and density change. During the sweep there is some deviation in  $f(\bar{b},N)$ , but

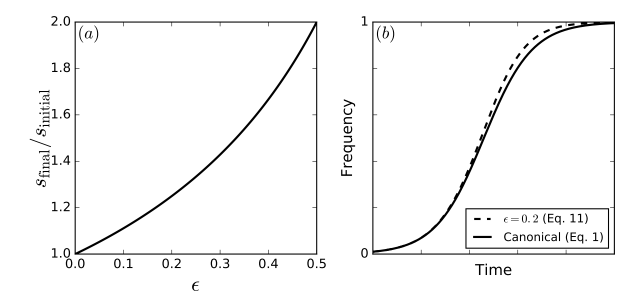


Figure 6: (a) Change in the selection coefficient between the beginning and end of a sweep of a type that experiences proportionally  $1 - \epsilon$  fold fewer crowding-induced deaths. The population is in demographic equilibrium at the start and end of the sweep. (b) Example equilibrium-to-equilibrium sweep.

this deviation is an order of magnitude smaller than for a  $\delta$  sweep (the density-dependent deviation constant s in Fig. 6 is of order  $\epsilon$ , whereas the analogous effect for b sweep in our variable-density lottery is only of order  $\epsilon^2$ ; see Appendix D for details). Since selection must already be strong for a  $\delta$ -sweep to threaten Eq. (??), the density-independent model applies effectively exactly for equilibrium b-sweeps. Note, however, that the selection coefficient for b-sweeps (as defined by differences in  $\Delta n_i/n_i$ ) does depend on frequency because of the  $1/\bar{b}$  factor.

If selection acts simultaneously on more than one trait in our variable-density lottery, then evolution in a density-regulating trait can drive changes in the strength of selection on a trait subject to density-dependent selection (Fig. 7). This can produce behavior analogous to selection on  $\delta$  in Fig. 6.

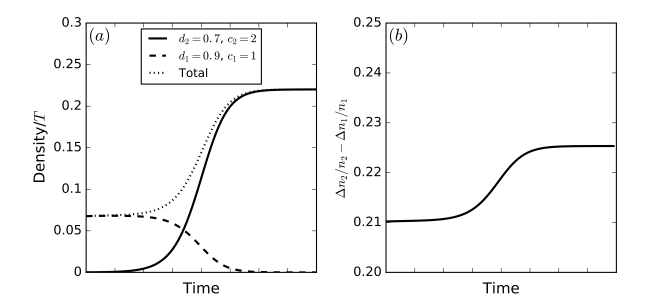


Figure 7: Simultaneous selection on d and c in our variable-density lottery model as predicted by Eq. (6). Selection is not constant over the sweep because d is density-regulating and c is density-dependent.

#### Discussion

Summarizing the properties of selection in our variable-density lottery model: (i) c-selection 360 is density-dependent, but c does not regulate density; (ii) d regulates density, but d-selection 370 is density-independent; (iii) b regulates density and b-selection is density-dependent. Yet, 371 despite the differences between b, c and d, in a constant environment selection that only 372 involves one of these traits obeys the density-independent relative fitness description of se-373 lection almost exactly. When strong selection acts on more than one of b, c and d (Fig. 7), 374 the constant-s approximation breaks down. The c and d traits exemplify the two distinct 375 directions in which density and selection can interact: selection may depend on density, and density may change in response to ongoing selection (Prout, 1980). The combination of 377 both is necessary to threaten the constant-s approximation. Remarkably, the b trait demon-378 strates that the combination is not sufficient; the density-dependence of b-selection effectively 379 disappears over equilibrium-to-equilibrium b-sweeps.

Selection in the variable-density lottery is quite different from classical density-dependent 381 selection (see "Introduction" and "The response of density to selection; c-selection versus 382 K-selection"). In the latter, only one life-history stage is represented, and the effects of 383 crowding appear as a reduction in absolute fitness that only depends on the type densities at 384 this life-history stage (e.g. the  $n_i^2$  and  $n_i n_j$  terms in the Lotka-Volterra equation). Selection 385 in crowded populations takes broadly one of two forms: selection for greater carrying capacity 386 (K-selection) or selection on competition coefficients ( $\alpha$ -selection). These are both " $\delta$ -like" 387 in the sense that selection depends on density and also causes density to change ( $\delta$  is defined 388 in Eq. (12)). Strong selection is therefore sufficient for Eq. (??) to break down (Fig. 6), and 389 no distinction is made between density-regulating and density-dependent traits. 390

The distinctive properties of selection in the variable-density lottery arise from a repro-391 ductive excess which appears when the number of propagules is greater than the number 392 of available territories. Then only  $\approx 1/L$  of the juveniles contesting unoccupied territories 393 survive to adulthood. Unlike the role of adult density  $n_i$  in single-life-stage models, it is 394 the propagule densities  $l_i$  that represent the crowding that drives competition. Reproduc-395 tive excess produces relative contests in which fitter types grow at the expense of others by preferentially filling the available adult "slots". The number of available slots can remain fixed or change independently of selection at the juvenile stage. By ignoring reproductive 398 excess, single life-stage models are biased to have total population density be more sensi-399 tive to ongoing selection. In this respect, the viability selection heuristics that are common 400 in population genetics (Gillespie, 2010, pp. 61) actually capture an important ecological 401 process without making the full leap to complex age-structured models. 402

Looking beyond the variable-density lottery, it is not clear which forms of crowdinginduced selection are more likely to occur in nature. Even if reproductive excesses are ubquitous, strictly relative c-like traits could pleiotropically interact with density-regulating traits so often that  $\delta$ -like behavior is prevalent. For instance, in the case in the case of well-

mixed indirect exploitation competition for consumable resources, the  $R^*$  rule suggests that competitive ability is intimately linked to equilibrium resource density, and hence that  $\delta$ -like 408 behavior would be prevalent. However, this conclusion is sensititive to the assumptions of 409 well-mixed resource competition models. Spatial localization of consumable resources (e.g. 410 for plants due to restricted movement of nutrients through soils) will tend to create territorial 411 contests similar to the lottery model, where resource competition only occurs locally and can 412 be sensitive to contingencies such as the timing of propagule arrival Bolker and Pacala (1999). 413 In this case, resource competition is effectively subsumed into a territorial competitive ability 414 trait akin to c, which would likely affect N much more weakly than suggested by the  $R^*$  rule 415 (assuming no pleiotropic interactions with b or d). 416

Moreover, even in well mixed populations, competition does not only involve indirect ex-417 ploitation of shared resources, but also direct interference. Interference competition can dra-418 matically alter the dynamics of resource exploitation Case and Gilpin (1974); Amarasekare 419 (2002), and is more likely than the exploitation of shared resource pools to involve rela-420 tive contests akin to c-selection. For instance, sexual selection can be viewed as a form of 421 relative interference competition between genotypes. Thus, a priori we should not expect 422 crowding in nature to only involve selection that is  $\delta$ -like. Other forms of selection like cselection (that is, strictly relative traits in density-regulated populations) are also likely to be important. Note that in the classical density-dependent selection literature, interference 425 competition is closely associated with  $\alpha$ -selection and the idea that selection need not affect 426 equilibrium density Gill (1974). However, as explained above,  $\alpha$ -selection does transiently 427 affect population density and can therefore be regarded as  $\delta$ -like. 428

The above findings underscore that the most serious threat to the constant-s approximation arises due to deviations from demographic equilibrium as a result of changes in the demographic rates of the types already present i.e. as a result of a temporally-variable environment. While transient deviations from demographic equilibrium driven by the appearance of new types can also threaten the constant-s approximation, they require strong selection that is both density-dependent and affects a density-regulating trait (and even then the constant-s approximation may hold). In contrast, temporally-variable environments can dramatically alter frequency trajectories for individual sweeps (e.g. Fig. 9.5 in Otto and Day (2011); Fig. 5 in Mallet (2012)), as well as the long-term outcomes of selection (Lande et al., 2009).

This suggests that in systems like the wild Drosophila example mentioned in the Introduction, there is indeed no choice but to abandon relative fitness. Our variable-density lottery could provide a useful starting point for analyzing evolution in this and other farfrom-equilibrium situations for two reasons: 1) the b, c, d trait scheme neatly distinguishes between different aspects of the interplay between density and selection; 2) lottery models in general are mathematically similar to the Wright-Fisher model, which should facilitate the analysis of genetic drift when N is unstable.

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# Appendix A: Growth equation derivation

- In this appendix we derive Eq. (6). 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.
- We start by separating the right hand side of Eq. (3) into three components

$$\Delta_{+}n_{i} = \Delta_{u}n_{i} + \Delta_{r}n_{i} + \Delta_{a}n_{i}, \tag{13}$$

which vary in relative magnitude depending on the propagule densities  $l_i$ . 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

$$\Delta_u n_i = U l_i e^{-L} = m_i e^{-L}. \tag{14}$$

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  $(x_i = 1 \text{ and } X_i \ge 1 \text{ where } X_i = \sum_{j \ne i} x_j$ is the number of nonfocal propagules; r stands for "rare"). The number of territories where this occurs is  $Up_i(1)P(X_i \ge 1) = m_i e^{-l_i}(1 - e^{-(L-l_i)})$ . Thus

$$\Delta_r n_i = m_i e^{-l_i} \left(1 - e^{-(L-l_i)}\right) \left\langle \frac{c_i}{c_i + \sum_{j \neq i} c_j x_j} \right\rangle_{\tilde{p}}, \tag{15}$$

where  $\langle \rangle_{\tilde{p}}$  denotes the expectation with respect to the probability distribution  $\tilde{p}$  of nonfocal propagule abundances  $x_j$ , in those territories where exactly one focal propagule, and at least one non-focal propagule, landed.

The final contribution,  $\Delta_a n_i$ , accounts for territories where two or more focal propagules are present  $(x_i \ge 2; a \text{ stands for "abundant"})$ . Similar to Eq. (15), we have

$$\Delta_a n_i = U(1 - (1 + l_i)e^{-l_i}) \left\langle \frac{c_i x_i}{\sum_j c_j x_j} \right\rangle_{\hat{p}}$$
(16)

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

To derive Eq. (6) we approximate the expectations in Eq. (15) and Eq. (16) by replacing

 $x_i$  and the  $x_j$  with "effective" mean values as follows

$$\left\langle \frac{c_i}{c_i + \sum_{j \neq i} c_j x_j} \right\rangle_{\tilde{p}} \approx \frac{c_i}{c_i + \sum_{j \neq i} c_j \langle x_j \rangle_{\tilde{q}}}.$$
 (17)

$$\left\langle \frac{c_i x_i}{\sum_j c_j x_j} \right\rangle_{\hat{p}} \approx \frac{c_i \langle x_i \rangle_{\hat{q}}}{\sum_j c_j \langle x_j \rangle_{\hat{q}}}.$$
 (18)

Here the effective means  $\langle \rangle_{\tilde{q}}$  and  $\langle \rangle_{\hat{q}}$  are taken with respect to new distributions  $\tilde{q}$  and  $\hat{q}$ , respectively. In the following subsection we define  $\tilde{q}$  and  $\hat{q}$  and explain our reasoning for using these distributions to take the effective means.

### The effective distributions $ilde{q}$ and $\hat{q}$

579

The approximations (17) and (18) must be consistent between rare and common types. To 584 illustrate, suppose that two identical types (same b, c and d) are present, with low  $l_1 \ll 1$ 585 and high density  $l_2 \approx L \gg 1$  respectively. Since L is large, uncontested territories make 586 up a negligible fraction of the total. The rare type grows almost entirely due to  $\Delta_r n_1$ , 587 while the common type grows almost entirely due to  $\Delta_a n_2$ . To ensure consistency, the ap-588 proximate per-capita growth rates implied by the approximations (17) and (18) must be 589 equal  $\Delta_r n_1/m_1 = \Delta_a n_2/m_2$ . Even small violations of this consistency condition would mean 590 exponential growth of one type relative to the other. This behavior is clearly pathological, 591 because any single-type population can be arbitrarily partitioned into identical rare and com-592 mon subtypes. Thus, predicted growth or decline would depend on an arbitrary assignment 593 of rarity. 594 For example, suppose that we use  $\tilde{p}$  and  $\hat{p}$  to calculate the effective means. The right hand side of Eq. (17) is then approximately 1/(L+1), and since  $l_1 \ll 1$  and  $L \gg 1$  we have  $\Delta_r n_1 \approx 1/(L+1)$  in Eq. (15). Similarly, for the common type,  $\sum_j \langle x_j \rangle_{\hat{p}} = L$  in Eq. (18), and so  $\Delta_a n_2 \approx 1/L$ . Thus, the identical rare type is pathologically predicted to decline in 599 frequency.

The effective distributions  $\tilde{q}$  and  $\hat{q}$  are devised to avoid this pathology. The idea is to make the approximation that the distribution for the total number of propagules per territory is the same in all territories. This is only an approximation because conditioning on focal propagules being present does change the distribution of X in the corresponding subset of territories (in the above example, the mean propagule density across all territories is L, but in the territories responsible for the growth of the rare type we have  $\langle X \rangle_{\tilde{p}} = L + 1$ ).

More formally, let  $\mathbf{x}$  denote the vector of propagule abundances  $(x_1, \dots, x_G)$  in a given territory, and  $\mathbf{x}_i = (x_1, \dots, x_{i-1}, x_{i+1} \dots, x_G)$  similarly denote the vector of non-focal abundances, so that  $p(\mathbf{x}_i) = p_1(x_1) \cdots p_{i-1}(x_{i-1}) p_{i+1}(x_{i+1}) \cdots p_G(x_G)$ . The corresponding total propagule numbers are denoted  $X = \sum_j x_j$  and  $X_i = X - x_i$ . Then, in territories where one focal propagule and at least one non-focal propagule are present, the effective distribution is defined by

$$\tilde{q}(\mathbf{x}_i) = \sum_{X=2}^{\infty} P(X|X \ge 2) p(\mathbf{x}_i|X_i = X - 1), \tag{19}$$

where the total number of propagules X follows a Poisson distribution with mean L, and  $P(X|X \ge 2) = P(X)/P(X \ge 2) = P(X)/(1 - (1 + L)e^{-L})$ . Similarly, in territories where more than one focal propagule is present, the effective distribution is defined by

$$\hat{q}(\mathbf{x}) = \sum_{X=2}^{\infty} P(X|X \ge 2)p(\mathbf{x}|x_i \ge 2, X). \tag{20}$$

### Calculating the effective means

Here we calculate the effective means, starting with the  $\Delta_r n_i$  component. We have

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

$$= \frac{1}{1 - (1 + L)e^{-L}} \sum_{X=2}^{\infty} P(X) \sum_{\mathbf{x}_i} p(\mathbf{x}_i | X_i = X - 1) x_j. \tag{21}$$

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,

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

$$= \frac{l_j}{1 - (1 + L)e^{-L}} \frac{L - 1 + e^{-L}}{L - l_i},$$
(22)

where the last line follows from  $\sum_{X=2}^{\infty} P(X)(X-1) = \sum_{X=1}^{\infty} P(X)(X-1) = \sum_{X=1}^{\infty} P(X)X - \sum_{X=1}^{\infty} P(X)$ . Substituting Eqs. (17) and (22) into Eq. (15), 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. (7).

Turning now to the  $\Delta_a n_i$  component, the mean focal abundance is

$$\langle x_i \rangle_{\hat{q}} = \sum_{\mathbf{x}} \hat{q}(\mathbf{x}) x_i$$

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

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

$$= l_i \frac{1 - e^{-l_i}}{1 - (1 + l_i)e^{-l_i}}.$$
(24)

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

$$\langle x_{j} \rangle_{\hat{q}} = \sum_{X=2}^{\infty} P(X|X \ge 2) \sum_{\mathbf{x}} p(\mathbf{x}|x_{i} \ge 2, X) x_{j}$$

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

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

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

$$= \frac{l_{j}}{L - l_{i}} \left( L \frac{1 - e^{-L}}{1 - (1 + L)e^{-L}} - l_{i} \frac{1 - e^{-l_{i}}}{1 - (1 + l_{i})e^{-l_{i}}} \right). \tag{25}$$

In going from line 2 to 3, we used the same logic used to evaluate the inner sum in Eq. (21), and in going from 3 to 4 we have separately evaluated the contributions from the X and  $x_i$ terms in the numerator. Combining these results with Eqs. (16) and (18), we obtain

$$\Delta_a n_i = m_i A_i \frac{c_i}{\overline{c}},\tag{26}$$

where  $A_i$  is defined in Eq. (7).

#### Approximation limits

Eq. (17) and (18) must not only be consistent with each other, they must also be individually good approximations. Here we evaluate these approximations.

The fundamental requirement for making the replacement in Eqs. (17) and (18) is that we can ignore the fluctuations in the  $x_i$  and hence replace them with a constant effective mean value. Mathematically, we require that the standard deviations  $\sigma_{\tilde{q}}(\sum_{j\neq i}c_jx_j)$  and  $\sigma_{\tilde{q}}(\sum_{j}c_jx_j)$  must be sufficiently small compared to the corresponding means  $\langle\sum_{j\neq i}c_jx_j\rangle_{\tilde{q}}$ and  $\langle\sum_{j}c_jx_j\rangle_{\hat{q}}$  in Eqs. (17) and (18) respectively.

To evaluate these standard deviations, we will work with  $\tilde{p}$  and  $\hat{p}$  distributions instead of  $\tilde{q}$  and  $\hat{q}$ . This is mathematically much simpler because the  $x_i$  are independent under  $\tilde{p}$  and  $\hat{p}$ , and is justified by the fact that  $\tilde{p}$  and  $\hat{p}$  are closely related to  $\tilde{q}$  and  $\hat{q}$  respectively, and so we expect the relevant means and standard deviations will be similar.

Starting with Eq. (17), we have  $\langle x_j \rangle_{\tilde{p}} = l_j/C$ , where  $C = 1 - e^{-(L-l_i)}$ , and the corresponding variances and covariances are given by

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

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

$$= \left(1 - \frac{1}{C}\right) \frac{l_{j}^{2}}{C} + \frac{l_{j}}{C}, \tag{27}$$

and

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

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

$$= \left(1 - \frac{1}{C}\right) \frac{l_{j} l_{k}}{C} \qquad j \neq k. \tag{28}$$

Note that 1-1/C is negative because C<1. Decomposing the variance in  $\sum_{j\neq i} c_j x_j$ ,

$$\sigma_{\tilde{p}}^2(\sum_{j\neq i} c_j x_j) = \sum_{j\neq i} \left[ c_j^2 \sigma_{\tilde{p}}^2(x_j) + 2 \sum_{k>j, k\neq i} c_j c_k \sigma_{\tilde{p}}(x_j, x_k) \right], \tag{29}$$

636 we obtain

$$\frac{\sigma(\sum_{j\neq i} c_j x_j)}{\langle \sum_{j\neq i} c_j x_j \rangle} = C^{1/2} \frac{\left(\sum_{j\neq i} c_j^2 l_j + (1 - \frac{1}{C}) \left(\sum_{j\neq i} c_j l_j\right)^2\right)^{1/2}}{\sum_{j\neq i} c_j l_j}.$$
 (30)

Eq. (30) reveals two key points. First, when the  $c_i$  have similar magnitudes (their ratios 637 are of order one), Eq. (17) is an excellent approximation. In this case, the right hand side of 638 Eq. (30) is approximately equal to  $C^{1/2} \left( \frac{1}{L-l_i} + 1 - \frac{1}{C} \right)^{1/2}$ , which is small for both low and 639 high nonfocal densities. The worst case scenario occurs when  $L - l_i$  is of order one, and it 640 can be directly verified that Eq. (17) is then still a good approximation (see Fig. 8). Second, 641 if some of the  $c_j$  are much larger than the others, the relative fluctuations in  $\sum_{j\neq i} c_j x_j$  can 642 be large. Specifically, in the presence of a rare, strong competitor  $(c_j l_j \gg c_{j'} l_{j'})$  for all other nonfocal types j', and  $l_j \ll 1$ ), then the right hand side of Eq. (30) can be large and we cannot make the replacement Eq. (17). Fig. 8 shows the breakdown of the effective mean approximation when the are large differences in c. Turning now to Eq. (18), all covariances between nonfocal types are now zero, so that

10 Turning now to Eq. (18), all covariances between nonlocal types are now zero, so that  $\sigma_{\hat{p}}^2(\sum c_j x_j) = \sum c_j^2 \sigma_{\hat{p}}^2(x_j)$ , where  $\sigma_{\hat{p}}^2(x_j) = l_j$  for  $j \neq i$ . Here

$$\sigma_{\hat{p}}^{2}(x_{i}) = \frac{l_{i}}{D} \left( l_{i} + 1 - e^{-l_{i}} - \frac{l_{i}}{D} \left( 1 - e^{-l_{i}} \right)^{2} \right), \tag{31}$$

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

$$\frac{\sigma_{\hat{p}}(\sum c_j x_j)}{\langle \sum c_j x_j \rangle} = \frac{\left(\sum_{j \neq i} c_j^2 l_j + c_i^2 \sigma_{\hat{p}}^2(x_i)\right)^{1/2}}{\sum_{j \neq i} c_j l_j + c_i l_i (1 - e^{-l_i})/D}.$$
(32)

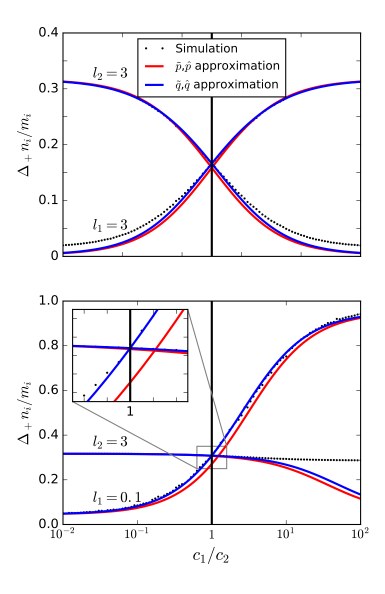


Figure 8: Comparison of our  $\tilde{q},\hat{q}$  approximation with simulations, and also with the naive  $\tilde{p},\hat{p}$  approximation, as a function of the relative c difference between two types. Our approximation breaks down in the presence of large c differences. The inset shows the pathology of the  $\tilde{p},\hat{p}$  approximation — growth rates are not equal in the neutral case c=1. Simulation procedure is the same as in Fig. 3, with  $U=10^5$ .

Similarly to Eq. (30), the right hand side of Eq. (32) is small for both low and high nonfocal densities. Again, the worst case scenario occurs when  $l_i$  and  $L - l_i$  are of order 1, but Eq. (18) is still a good approximation in this case. Again, the approximation breaks down in the presence of a rare, strong competitor (Fig. 8).

# Appendix B: 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 equilibrium 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. (11), we obtain

$$\frac{dn_1}{dt} = r_1(\alpha_{11} - \alpha_{12})n_1n_2 
\frac{dn_2}{dt} = r_2(\alpha_{22} - \alpha_{21})n_1n_2$$
(33)

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

$$r_1(\alpha_{\text{intra}} - \alpha_{12}) + r_2(\alpha_{\text{intra}} - \alpha_{21}) = 0.$$
(34)

To get some intuition for Eq. (34), suppose that a mutant arises with improved competitive ability but identical intrinsic growth rate and equilibrium density  $(r_1 = r_2 \text{ and } \alpha_{11} = \alpha_{22})$ . This could represent a mutation to an interference competition trait, for example (Gill,  $^{664}$  1974). Then, according the above condition, for N to remain constant over the sweep, the  $^{665}$  mutant must find the wildtype more tolerable than itself by exactly the same amount that  $^{666}$  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

$$\sum_{ij} r_i (\alpha_{\text{intra}} - \alpha_{ij}) p_i p_j = 0, \tag{35}$$

which depends on frequency and thus cannot be satisfied in general for constant inter-type coefficients  $\alpha_{ij}$ . Therefore, Lotka-Volterra selection will generally involve non-constant N.

# Appendix C: 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 N. 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  $D_i$  and  $D_j$  respectively, where we

have  $d_i = b_i(1 - D_i) = b_j(1 - D_j)$ . We obtain

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

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

$$= \frac{1}{4} (2d_i + b_i(1 - D_j) + b_j(1 - N_i)). \tag{36}$$

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

$$\frac{f_{\text{half}}}{d_i} = \frac{1}{4} \left( 2 + \frac{b_i}{b_j} + \frac{b_j}{b_i} \right) 
= \frac{1}{4} \left( 2 + \frac{1}{1+\epsilon} + 1 + \epsilon \right) 
= 1 + \frac{1}{4} (\epsilon^2 - \epsilon^3 + \dots),$$
(37)

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

By contrast, for a  $\delta$  sweep in Eq. (12), 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. (6).