A preliminary rcK derivation

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"...the concept of fitness is probably too complex to allow of a useful mathematical development. Since it enters fundamentally into many population genetics considerations, it is remarkable how little attention has been paid to it." — Warren J. Ewens, Mathematical Population Genetics I, 2004

1 Introduction

Evolutionary models differ greatly in their treatment of fitness. In models of genetic evolution, genotypes are typically assigned constant (or frequency-dependent) selection coefficients describing the change in their relative frequencies over time due to differences in viability. This considerably simplifies the mathematics of selection, facilitating greater genetic realism, and can be justified over sufficiently short time intervals [1, p. 276]. However, selection can have very different effects when operating on different types of traits, and evolutionary changes in one population can lead to complicated ecological responses.

By contrast models of phenotypic trait evolution represent the change in phenotypic abundances over time using absolute fitness functions which describe how those traits affect survival and reproduction in particular ecological scenarios. This approach is powerful enough to model eco-evolutionary feedbacks between co-evolving traits, but is generally problem-specific and restricted to only a few traits at a time.

Far less work has been done to model fitness in more general terms than particular traits or ecological scenarios, while still capturing key distinctions between different forms of selection. Perhaps this is not surprising given that fitness is such a complex quantity, dependent on all of a phenotype's

functional traits [2] as well as its biotic and abiotic environment. In most cases, a detailed, trait-based, predictive model of fitness would be enormously complicated and have narrow applicability. It is therefore easy to doubt the feasibility of a simplified, general mathematical treatment of fitness [1, p. 276]. Even MacArthur's famous r/K selection scheme is now almost exclusively known as a framework for understanding life-history traits, and judged on its failure in that role [3, 4, 5, 6]. In spite of the r/K scheme's original purpose as an extension of the existing population-genetic treatment of selection to account for population density [7], comparatively few attempts have been made to develop it further as a mathematical analysis of the major different forms of selection.

Nevertheless, there are strong indications there are broader principles governing the operation of selection. In many groups of organisms (including corals [8], insects [9], fishes [10] and plants [11]), species can be partitioned into a small number of distinct trait clusters corresponding to fundamentally distinct "primary strategies" [12]. The most famous example is Grime's plant trait classification scheme [13, 14, 11]. Grime considered two broad determinants of population density: stress (persistent hardship e.g. due to resource scarcity, unfavorable temperatures or toxins) and disturbance (intermittent destruction of vegetation e.g. due to trampling, herbivory, pathogens, extreme weather or fire). The extremes of these two factors define three primary strategies denoted by C/S/R respectively: competitors "C" excel in low stress, low disturbance environments; stress tolerators "S" excel in high stress, low disturbance environments; and ruderals "R" excel in low stress, high disturbance environments. Survival is not possible in high-stress, highdisturbance environments. Grime showed that measures of C, S and R across a wide range of plant species are anti-correlated, so that strong C-strategists are weak S and R strategists, and so on. Thus, plant species can be classified on a triangular C/S/R ternary plot [13]. Trait classification schemes for other organisms closely parallel Grime's scheme [12].

Trait classification schemes show empirically that, beneath the complicated details of trait variation, even among closely-related species, fitness is predominantly determined by a few key factors such as intrinsic reproducive rate or stress-tolerance. However, while trait classification schemes are firmly grounded in trait data, they are verbal and descriptive rather than mathematical, a recognized hinderance to their broader applicability (e.g. [15]).

The aim of this paper is explore the interplay between some major di-

mensions of fitness in a simplified, spatially-homogeneous model of genotype growth, dispersal and competition. Building on the earlier r/K and C/S/R schemes, a central question is how fitness depends on the interaction between population density, intrinsic birth/death rates and competitive ability.

We broadly follow the spirit of MacArthur's r/K selection scheme in that our model is intended to account for fundamentally different forms of selection without getting entangled in the intricacies of particular ecological scenarios. However, rather than building directly on MacArthur's formalism and its later extensions [cite alpha stuff] (e.g. using Lotka-Volterra equations), our model is devised primarily with Grime's C/S/R scheme in mind, and represents a quantitative formalization of how C/S/R manifests at the level of genotype evolution (as opposed to divergence between species). This choice is motivated in part by the substantial empirical support for the C/S/R scheme, and in part by the failings of the r/K low/high density dichotomy—many growth ability traits will confer advantages at both low and high densities, in which case r- and K- selection will effectively coincide (empirically, positive correlations between measures of r and K are common, both between species and strains [16, 17, 18, 19], and as a result of experimental evolution [16, 20]).

In section

2 Model

We assume that each individual in a population requires its own territory to survive and reproduce (a site-occupancy model). All territories are identical, and the total number of territories is K. Time t advances in discrete steps, each representing one generation. In generation t, the number of mature individuals (henceforth called "adults") of the i'th genotype is $n_i(t)$, the total number of adults is $N(t) = \sum_i n_i(t)$, and the number of unoccupied territories U(t) = K - N(t).

Each generation, adults produce m_i new offspring (henceforth called "propagules") which disperse at random over the U unnocupied territories. We ignore propagules that land on occupied territories — these are assumed to have no chance of survival and are not included in m_i . For simplicity, we assume $m_i = b_i n_i$, where b_i is a constant, genotype-specific birth rate.

The number of individuals of the *i*'th genotype landing in any particular territory is denoted x_i . Random dispersal implies that in the limit $K \to \infty$

 ∞ , with n_i/K held fixed, x_i is Poisson distributed with mean territorial propagule density $l_i = m_i/U$. Although K is finite in our model, we assume that K and the n_i are large enough that x_i is Poisson-distributed to a good approximation (details in Appendix A). This Poisson distribution is denoted $p_i(x_i)$. In particular, we do not consider the stochastic behavior that occurs at very low n_i due to the lottery nature of territorial acquisition (this will be discussed further when we consider the behavior of mutant lineages).

When multiple propagules land on the same territory, they compete to secure the territory as they develop. This territorial contest is modeled as a weighted lottery: the probability that genotype i wins a given territory by the next generation is $c_i x_i / \sum_j c_j x_j$ where c_i is a constant representing relative competitive ability.

The increase in n_i over one generation due to territorial acquisition, $\Delta_+ n_i$, is the sum of genotype i's victories over all U unoccupied territories. Since $p_1(x_1) \dots p_G(x_G)$ is equal to the proportion of unoccupied territories with x_1, \dots, x_G of the respective propagules (again, we assume that K is large enough that fluctuations in this proportion are negligible), this sum can be replaced by an expectation over the p_i . This gives

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

In addition to propagule birth and competition, occupied territories become unoccupied when adults die. We assume that these adult deaths occur at a constant, genotype-specific rate d_i . Thus, the overall change in genotype abundances is

$$\Delta n_i(t) = \Delta_+ n_i(t) - d_i n_i(t). \tag{2}$$

The change in total population size N, obtained by summing Eq. (2) over genotypes, is

$$\Delta N = U(1 - e^{-L}) - \sum_{i} d_{i} n_{i}, \tag{3}$$

where $L = \sum_{j} l_{j}$ is the overall propagule density. ΔN is independent of the competitive ability coefficients c_{i} ; thus, the c_{i} represent a strictly relative aspect of fitness in the sense that they can only influence N indirectly by changing genotype frequencies (which may change the mean birth and death rates in the population).

3 Results

3.1 Mean Field Approximation

Eq. (2) gives little intuition about the dynamics of density-dependent lottery competition, since (1) involves an expectation over the random dispersal distributions p_i , which depend on how the n_i change over time. We now evaluate this expectation using a "mean field" approximation; the intuition behind this approximation is as follows.

If the unoccupied territories are saturated with propagules from every genotype $(l_i \gg 1 \text{ for all genotypes})$, the fluctuations in the x_i are small compared to their means l_i (since the x_i are Poisson distributed), and so the composition of propagules in a territory will only rarely differ appreciably from the mean composition l_1, l_2, \ldots, l_G . Consequently, we can replace x_i with l_i in Eq. (1). This gives the classic lottery model [21],

$$\Delta_{+}n_{i}(t) = U(t)\frac{c_{i}m_{i}}{\sum_{j}c_{j}m_{j}} = b_{i}n_{i}\frac{1}{L}\frac{c_{i}}{\bar{c}},$$
(4)

where $\bar{c} = \sum_j c_j m_j / M$ is the mean propagule competitive ability for a randomly selected propagule $(M = \sum_j m_j)$ is the total number of propagules).

However, in general the l_i are not all large, and the x_i cannot simply be replaced by their means in Eq. (1). Indeed, Eq. (4) is nonsensical if l_i is sufficiently small: genotype i can win at most m_i territories, yet Eq. (4) demands a fraction $c_i m_i / \sum_j c_j m_j$ of the unoccupied territories U, no matter how large U is. The source of this pathological behavior when $l_i \ll 1$ is that $x_i = 1$ in the few territories where i propagules do land, and so i's growth comes entirely from territories which deviate appreciably from the mean.

Our mean field approximation is similar to the high- l_i approximation leading to Eq. (4) in that we replace the x_i with appropriate mean values. The key distinction is that territories with a single propagule from the focal genotype, which are critical at low densities, are handled separately. In place of the requirement of $l_i \gg 1$ for all i, our approximation only requires that there are no large discrepancies in competitive ability (discussed further below). We obtain (details in Appendix B)

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

where

$$R_{i} = \frac{\bar{c}e^{-l_{i}}(1 - e^{-(L-l_{i})})}{c_{i} + \frac{L-1 + e^{-L}}{1 - (1 + L)e^{-L}} \frac{\bar{c}L - c_{i}l_{i}}{L - l_{i}}},$$
(6)

and

$$A_{i} = \frac{\overline{c}(1 - e^{-l_{i}})}{c_{i}l_{i}\frac{1 - e^{-l_{i}}}{1 - (1 + l_{i})e^{-l_{i}}} + \sum_{j \neq i} \frac{c_{j}l_{j}}{L - l_{j}} \left(L\frac{1 - e^{-L}}{1 - (1 + L)e^{-L}} - l_{j}\frac{1 - e^{-l_{j}}}{1 - (1 + l_{j})e^{-l_{j}}}\right)}.$$
 (7)

Comparing Eq. (5) to Eq. (4), the classic lottery per-propagule 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 term, $R_i c_i/\bar{c}$ represents competitive victories when the i genotype is a rare invader in a high density population: from Eq. (6), $R_i \to 0$ when the i genotype is abundant ($l_i \gg 1$), or other genotypes are collectively rare ($L-l_i \ll 1$). The third term, $A_i c_i/\bar{c}$, represents competitive victories when the i genotype is abundant: $A_i \to 0$ if $l_i \ll 1$. 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 genotypes, we recover the classic lottery model (only the $A_i c_i/\bar{c}$ term remains, and $A_i \to 1/L$). Thus, Eq. (5) generalizes the classic lottery model to account for arbitrary propagule densities for each genotype.

Fig. 1 shows that Eq. (5) (and its components) closely approximate direct simulations of random dispersal and lottery competition over a wide range of propagule densities (obtained by varying U). Two genotypes are present, one of which has a c-advantage and is at low frequency. The growth of the low-frequency genotype relies crucially on the low-density competition term $R_i c_i/\bar{c}$, and also to a lesser extent on the high density competition term $A_i c_i/\bar{c}$ if l_1 is large enough (Fig. 1b). On the other hand, $R_i c_i/\bar{c}$ is negligible for the high-frequency genotype, which depends instead on high density territorial victories (Fig. 1d).

3.2 Invasion of rare genotypes and coexistence

To determine how b, c and d will evolve in a population where those traits are being modified by mutations, we need to know whether mutant lineages will grow (or decline) starting from low densities. In this section we discuss the behavior of rare genotypes predicted by Eq. (5).

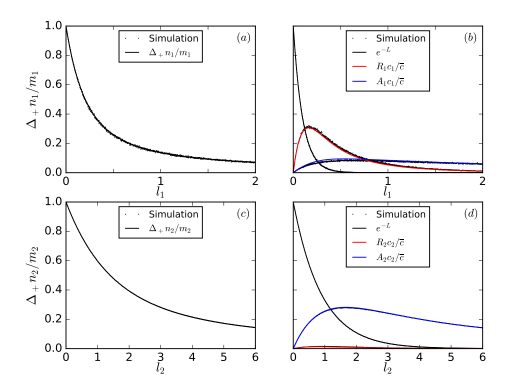


Figure 1: The change in genotype abundances in a density dependent lottery model is closely approximated by Eq. (5). $\Delta_+ n_i/m_i$ from Eq. (5) (and its separate components) are shown, along with direct simulations of random dispersal and lottery competition over one generation over a range of propagule densities (varied by changing U with the m_i fixed). Two genotypes are present. (a) and (b) show low-frequency genotype with c-advantage $(m_1/M=0.1, c_1=1.5)$, (c) and (d) show the high-frequency predominant genotype $(m_2/M=0.9, c_2=1)$.

Suppose that a population with a single genotype i is in equilibrium. Then $R_i = 0$, $\bar{c} = c_i$ and $\Delta n_i = 0$, and so Eq. (5) gives

$$b_i \left(e^{-L} + A_i \right) - d_i = 0. (8)$$

Now suppose that a new genotype j, which is initially rare, appears in the population. Then $A_j \ll 1$, $l_j \ll L$ and $\bar{c} \approx c_i$, and so, from Eq. (5), n_j will increase if

$$b_j \left(e^{-L} + R_j \frac{c_j}{c_i} \right) - d_j > 0. \tag{9}$$

Combining Eqs. (8) and (9), it is easily verified that if j is superior in one trait, but otherwise identical to i, it will invade. Moreover, j will eventually exclude i, since it is strictly superior. However, stable coexistence is possible between genotypes that are superior in different traits. To illustrate, suppose that j is better at securing territories $(c_j > c_i)$, that i is better at producing propagules $(b_i > b_j)$, and that $d_i = d_j$. Coexistence occurs if j will invade an i-dominated population, but i will also invade a j-dominated population ("mutual invasion"). It is not hard to show that this is possible, since if b_i is so large that $L \gg 1$ when i is dominant, and b_j is so small that $L \ll 1$ when j is dominant, then, combining Eqs. (8) and (9), we find that j invades i if

$$b_j c_j R_j - b_i c_i A_i > 0, (10)$$

while i invades j if

$$b_i - b_i > 0. (11)$$

Eq. (11) is satisfied by assumption, while Eq. (10) can always be satisfied with large enough c_j . The mechanism for coexistence is that territorial contests are important in *i*-dominated populations (high L), ensuring that the c-specialist j is not excluded, yet territorial contests are irrelevant in j-dominated populations (low L), ensuring that the b-specialist i is not excluded. Fig. 2 shows an example of this coexistence between b and c specialists.

A similar argument applies for coexistence between high-c and low-d specialists; again coexistence occurs because the importance of territorial contests declines along with propagule density L as the c-specialist increases in frequency. Coexistence is technically possible between b- and d-specialists which exactly satisfy $b_i/d_i = b_j/d_j$ (this follows from the fact that all propagules have the same success rate when $c_i = c_j$ i.e. $A_i + R_i = A_j + R_j$). However,

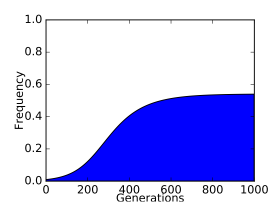


Figure 2: Coexistence between b ($c_i = 1$, $b_i = 1$) and c ($c_j = 2$, $b_j = 0.7$) specialists, where $d_i = d_j = 0.3$. Vertical axis shows frequency of the c-specialist predicted by Eq. (5).

this coexistence scenario is not biologically relevant, since the tiniest deviation from $b_i/d_i = b_j/d_j$ will lead to the eventual exclusion of the genotype with greater b_i/d_i .

If the rare genotype j arises due to mutation, then it's initial low-density behavior is more complicated than the above invasion analysis suggests. The mutant lineage starts with one individual $n_j = 1$, and remains at low abundance for many generations after its initial appearance. During this period, the mutant abundance n_j will behave stochastically, and the deterministic equations (1) and (5) do not apply (Section 2). However, if n_j becomes large enough, its behavior will become effectively deterministic, and governed by Eq. (5). For mutants with fitness greater than the population mean fitness, this process is known as "establishment", and occurs when n_j is of order 1/s, where s is the mutant's fitness advantage relative to the mean [23]. Here we do not consider the initial stochastic behavior of novel mutants, and have restricted our attention to the earliest deterministic behavior of rare genotypes. In particular, for beneficial mutations we have only considered the case where s is large enough that deterministic behavior starts when $n_j \ll N$.

3.3 Environmental archetypes and Grime's triangle

We can then determine which mutational changes in the traits b_i , c_i and d_i will be favored under different environmental conditions. Of particular interest

are Grime's "disturbance", "stress" and "ideal" environmental archetypes.

To proceed, we map these archetypes to different parameter regimes in our model. The ideal archetype is defined by low extrinsic mortality $d_i \ll 1$ and high birth rates, such that b_i significantly exceeds d_i , population density N/K is high, and thus so is the overall propagule density $L \gg 1$. In this case, competition between propagules is important.

The disturbance archetype is defined by unavoidably high extrinsic mortality, usually due to physical destruction of individual organisms, to the extent that the population is kept at low densities $N/K \ll 1$. This means that, even though b_i may be quite high, it cannot greatly exceed d_i , and that propagule densities are low $l_i \ll 1$. The terms proportional to c_i/\bar{c} in Eq. (5) are then negligible, and $\Delta_+ n_i$ depends primarily on b_i .

The stress archetype is more ambiguous, and has been the subject of an extensive debate in the plant ecology literature. In Grime's view, the archetypal stressful environmental imposes such severe challenges that success depends primarily on the ability to tolerate the stressors. Population density N/K is suppressed to such low levels that competition between individuals is not important. In our model, this again corresponds to b_i only slightly exceeding d_i and $l_i \ll 1$, similar to the disturbance archetype, except that stress now constrains b_i to be small.

The alternative view is that the stress archetype should rather be interpreted as a large reduction in the maximum number of individuals that can be supported [22]. For example, in the case that the stress is induced by a scarcity of consumable resources, competition for resources would likely be intense, and the stressed population should actually be regarded as having a high population density. In our model, this would imply a large reduction in K (greater per-individual territorial requirement), such that even though fewer individuals are supported, density N/K is still high.

The mapping of environmental achetypes to our model parameters is summarized in the first three rows of Fig. 3. Also shown is the approximate dependence of $\Delta_+ n_i$ on b_i and c_i for each archetype (fourth row), which can be used infer the expected direction of evolution for the traits b_i , c_i and d_i (fifth row).

The latter is inferred as follows. As noted in the previous section, if beneficial mutants can survive the low-abundance stochastic regime, their behavior is governed deterministically by Eq. (5). If their advantage in b, c or d is large enough to overcome the lottery rarity disadvantage, they will proceed to grow deterministically (establishment). The probability of estab-

	Ideal	Disturbance	Stress (G)	Stress (K)
Parameter-	$d_i \ll 1$	$d_i \gg 1$	$b_i \ll 1$	$b_i \ll 1$
regime	$b_i > d_i$	$b_i \approx d_i$	$b_i \approx d_i$	$b_i > d_i$
Density	High	Low	Low	High
$\Delta_+ n_i \propto$	$b_i c_i$	b_i	b_i	$b_i c_i$
Evolution	$\uparrow b, \uparrow c$	$\uparrow b$	$\downarrow d$	$\uparrow c$

Figure 3: The realization of Grime's environmental archetypes in our model, as well as the low-K variant of the stress archetype. Shown are the mapping to our parameters of each archetype, the approximate dependence of $\Delta_+ n_i$ on b_i and c_i , as well as the corresponding expected evolutionary changes in b_i , c_i and d_i .

lishment increases with the mutant fitness advantage, and is of order 1/N for neutral mutations. Consequently, the direction of evolutionary change is determined by which trait changes confer an appreciable benefit, subject to the constraints imposed by the environmental.

For example, in Grime's version of the stress archetype, population density is low, so competition is not important, and so only mutants with greater b or lower d will have an appreciably greater Δn_i . Mutations in c are effectively neutral, and will rarely establish. However, by definition of the stress archetype, b is constrained to be small. Thus, while some rare mutations may produce small improvements in b, it is much more likely that mutations will arise that lower d, making this the expected direction of evolutionary change for Grime's stress archetype.

Following Grime's original argument for a triangular scheme [14], Fig. 4 represents each environmental archetype schematically as a vertex on a triangular space defined by perpendicular stress and disturbance axes. The ideal archetype lies at the origin (no stress or disturbance), while the stress and disturbance archetypes lie at the limits of survival on their respective axes. The hypotenuse connecting the stress and disturbance endpoints represents the limits of survival in the presence of a combination of stress and disturbance. The direction of evolutionary change is different at each vertex, leading to the emergence of different trait clusters or "primary strategies".

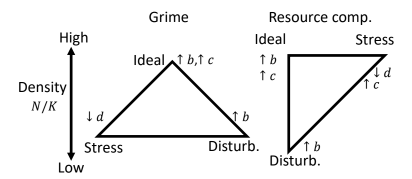


Figure 4: The realization of Grime's triangle in our model. Schematic representation of the triangular space bounded by the low/high extremes of stress/disturbance. The low-K interpretation of stress is also shown. The vertices of the triangles correspond to environmental archetypes. Selection favors different traits at each vertex, leading to different trait clusters.

4 Discussion

Grime's triangle without constraints r-K correlation, meaning of K selection Actual K selection Significance of stage structure caveats: large c discrepancy

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Appendix A: Poisson approximation

The propagule numbers x_i in different territories are not independent random variables. To determine the dispersal outcomes in all unoccupied territories

exactly, we would need to proceed territory-by-territory as follows. In the first territory we evaluate, x_i drawn from a binomial distribution with m_i trials and success probability 1/U. In the second, x_i is drawn from a binomial distribution with $m_i - x$ trials and success probability 1/(U - 1), where x is the number of propagules that landed in the first territory. And so on.

For sufficiently large K, holding n_i/K fixed, the Poisson limit theorem implies that the binomial distributions for x_i at each successive stage of this procedure are all closely approximated by a Poisson distribution with mean l_i , where we have used the fact that large K implies large U except in the biologically uninteresting case that there is vanishing population turnover $d_i \sim 1/K$.

Under the Poisson approximation, the total number of genotype i propagules $\sum x_i$ (summed over unoccupied territories) will deviate about its mean value m_i . Since the coefficient of variation of $\sum x_i$ is proportional to $1/\sqrt{m_i}$, these deviations are negligible unless m_i is very small (say of order 100 or less).

Appendix B: Derivation of growth equation

We separate the right hand side of Eq. (1) 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; for instance $p(x_1, \ldots, x_G) = p_1(x_1) \ldots p_G(x_G)$ and $p(x_1, \ldots, x_{i-1}, x_{i+1}, \ldots, x_G) = p_1(x_1) \ldots p_{i-1}(x_{i-1})p_{i+1}(x_{i+1}) \ldots p_G(x_G)$. We use P as a general shorthand for the probability of particular outcomes.

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

$$\Delta_u n_i = U l_i e^{-L} = m_i e^{-L}. \tag{12}$$

Competition when rare

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

$$\Delta_r n_i = m_i e^{-l_i} P \left\langle \frac{c_i}{c_i + \sum_{j \neq i} c_j x_j} \right\rangle_{\tilde{p}}, \tag{13}$$

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

We now show that, with respect to \tilde{p} , the standard deviation in $\sum_{j\neq i} c_j x_j$, $\sigma(\sum_{j\neq i} c_j x_j)$, is much smaller than its mean $\langle \sum_{j\neq i} c_j x_j \rangle_{\tilde{p}}$. Then x_j can be replaced by its mean in the last term in Eq. (13),

$$\left\langle \frac{c_i}{c_i + \sum_{j \neq i} c_j x_j} \right\rangle_{\tilde{p}} \approx \frac{c_i}{c_i + \sum_{j \neq i} c_j \langle x_j \rangle_{\tilde{p}}},\tag{14}$$

which will give us Eq. (6).

The exact expression for $\langle x_j \rangle_{\tilde{p}}$ is somewhat complicated. Letting k 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, \tilde{p} can

be written as

$$\tilde{p}(\mathbf{x}_{i}) = p(\mathbf{x}_{i}|k \geq 2, x_{i} = 1),
= \frac{P(k \geq 2|\mathbf{x}_{i}, x_{i} = 1)p(\mathbf{x}_{i}|x_{i} = 1)}{P(k \geq 2)},
= \frac{p(\mathbf{x}_{i}|x_{i} = 1)}{1 - (1 + L)e^{-L}},
= \frac{1}{1 - (1 + L)e^{-L}} \sum_{k=2}^{\infty} P(k)p(\mathbf{x}_{i}|\sum_{j \neq i} x_{j} = k - 1),
= \frac{1}{1 - (1 + L)e^{-L}} \sum_{k=2}^{\infty} \frac{P(k)\delta_{k-1}^{\sum_{j \neq i} x_{j}}}{P(\sum_{j \neq i} x_{j} = k - 1)} p(\mathbf{x}_{i}),
= \frac{Le^{-l_{i}}}{1 - (1 + L)e^{-L}} \sum_{k=1}^{\infty} \left(\frac{L}{L - l_{i}}\right)^{k} \frac{\delta_{k}^{\sum_{j \neq i} x_{j}}}{k + 1} p(\mathbf{x}_{i}), \tag{15}$$

where $\delta_k^{\sum_{j\neq i} x_j} = 1$ if $\sum_{j\neq i} x_j = k$, and equals zero otherwise. Then, since

$$\sum_{\mathbf{x}_i} \delta_k^{\sum_{j \neq i} x_j} p(\mathbf{x}_i) x_j = \frac{l_j}{L - L_j} k P(\sum_{j \neq i} x_j = k)$$
$$= l_j P(\sum_{j \neq i} x_j = k - 1), \tag{16}$$

after some algebra we obtain,

$$\langle x_j \rangle_{\tilde{p}} = \frac{l_j}{1 - (1 + L)e^{-L}} \frac{L - 1 + e^{-L}}{L - l_i}.$$
 (17)

To calculate the relative fluctuations in $\sum_{j\neq i} c_j x_j$, we use the following approximation, which gives considerably simpler expressions for the means, variances and covariances of the x_j compared with the exact expressions using \tilde{p} . Rather than evaluating the situation in each territory after dispersal as above, we let \tilde{p} instead be the \mathbf{x}_i dispersal probabilities in a territory where one focal propagule is assumed to be present, conditional on $\sum_{j\neq i} x_j > 1$. This gives $\langle x_j \rangle_{\tilde{p}} = l_j/C$,

$$\sigma^2(x_j) = \frac{l_j^2}{C} \left(1 - \frac{1}{C} \right) + \frac{l_j}{C},\tag{18}$$

and

$$\sigma(x_j, x_k) = \frac{l_j l_k}{C} \left(1 - \frac{1}{C} \right), \tag{19}$$

where $C = 1 - e^{-(L-l_i)}$ (note the difference from Eq. (17) for $\langle x_j \rangle_{\tilde{p}}$). Then, since

$$\sigma^{2}(\sum_{j\neq i} c_{j}x_{j}) = \sum_{j\neq i} \left[c_{j}^{2}\sigma^{2}(x_{j}) + 2\sum_{k>j} c_{j}c_{k}\sigma(x_{j}, x_{k}) \right], \tag{20}$$

and 1/C > 1, we have

$$\frac{\sigma(\sum_{j\neq i} c_j x_j)}{\langle \sum_{j\neq i} c_j x_j \rangle} < C^{1/2} \frac{\left(\sum_{j\neq i} c_j^2 l_j\right)^{1/2}}{\sum_{j\neq i} c_j l_j}.$$
 (21)

Without loss of generality, we restrict attention to the case that the total nonfocal density $L - l_i$ is of order 1 or larger (otherwise $\Delta_r n_i$ does not contribute significantly to $\Delta_+ n_i$ because $\Delta_r n_i$ is proportional to $C = 1 - e^{-(L-l_i)}$).

When at least some of the nonfocal propagule densities are large $l_j \gg 1$, then the RHS of Eq. (21) is $\ll 1$, as desired. This is also the case if none of the nonfocal genotype densities are large and the c_j are all of similar magnitude (their ratios are of order one); the worst case scenario occurs when $(L - l_i) \sim O(1)$, in which case the negative covariances (Eq. (19)) which were neglected in the RHS of Eq. (21) significantly reduce the overall variance $\sigma^2(\sum_{j\neq i} c_j x_j)$.

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, if $c_j l_j \gg c_k l_k$ $(j, k \neq i, j \neq k)$ and $l_j \ll 1$ (i.e. in the presence of a rare, extremely strong competitor), then we cannot make the replacement Eq. (14).

Substituting Eqs. (14) and (17) into Eq. (13), we obtain

$$\Delta_r n_i \approx m_i R_i \frac{c_i}{\overline{c}},\tag{22}$$

where R_i is defined in Eq. (6).

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. (13), we

have

$$\Delta_a n_i = U(1 - (1 + l_i)e^{l_i}) \left\langle \frac{c_i x_i}{\sum_j c_j x_j} \right\rangle_{\hat{p}}$$
(23)

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

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

$$\left\langle \frac{c_i x_i}{\sum_j c_j x_j} \right\rangle_{\hat{p}} \approx \frac{c_i \langle x_i \rangle_{\hat{p}}}{\sum_j c_j \langle x_j \rangle_{\hat{p}}}.$$
 (24)

Following a similar procedure as for $\Delta_r n_i$, where the vector of propagule abundances is denoted \mathbf{x} , we have

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

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

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

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

for $j \neq i$, and

$$\langle x_i \rangle_{\hat{p}} = l_i \frac{1 - e^{-l_i}}{1 - (1 + l_i)e^{-l_i}}.$$
 (26)

To calculate the relative fluctuations in $\sum_{j\neq i} c_j x_j$, we use a similar approximation as for $\Delta_r n_i$: \tilde{p} is approximated by the \mathbf{x} dispersal probabilities in a territory where at least two focal propagule is assumed to be present. All covariances are now zero, so that $\sigma^2(\sum c_j x_j) = \sum c_j^2 \sigma^2(x_j)$, where $\sigma^2(x_j) = l_j$ for $j \neq i$. The expression for $\sigma^2(x_i)$ is more complicated. We assume $p(x_i = 0) \approx 0$ without loss of generality (since otherwise $D \gg 1$ and Δn_a is negligible). Then

$$\sigma^2(x_i) = \frac{l_i^2}{D} \left(1 - \frac{1}{D} \right) + \frac{l_i}{D},\tag{27}$$

where $D = 1 - (1 + l_i)e^{-l_i}$, analogous to Eq. (18), and

$$\frac{\sigma(\sum c_j x_j)}{\langle \sum c_j x_j \rangle} \approx \frac{\left(\sum_{j \neq i} c_j^2 l_j + c_i^2 \sigma^2(x_i)\right)^{1/2}}{\sum_{j \neq i} c_j l_j + c_i l_i / D}.$$
 (28)

Similarly to Eq. (21), the RHS of (28) will not be $\ll 1$ if there is a nonfocal genotype j with $l_j \ll 1$ and $c_j l_j \gg c_k l_k$ for $j, k \neq i, j \neq k$. When this is not the case, then since l_i must be of order 1 or larger for $\Delta_a n$ to make an appreciable contribution to $\Delta_+ n_i$, the RHS of Eq. (28) is $\ll 1$ as desired.

Combining Eqs. (23) and (24), we obtain

$$\Delta_a n_i = m_i A_i \frac{c_i}{\overline{c}},\tag{29}$$

where A_i is defined in Eq. (7).