Local adaptation and dispersal evolution interact to drive population response to climate change

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1 Introduction

Climate change is expected to dramatically reshape global biogeographic patterns as some species shift their ranges to track changing environmental conditions (Gonzalez et al., 2010; Peñuelas and Boada, 2003; Hansen et al., 2001; Scholze et al., 2006). These range shifts are generally expected to proceed upwards in latitude, elevation, or both as average global temperatures continue to rise (Loarie et al., 2009). In fact, contemporary range shifts have already been observed in a wide range of taxa (Chen et al., 2011; Walther et al., 2002; Parmesan and Yohe, 2003; Parmesan, 2006; Parmesan et al., 1999; Perry et al., 2005). Such range shifts present significant challenges to current and future conservation efforts as they can result in the extinction of populations failing to track a changing climate (Davis and Shaw, 2001; Parmesan, 2006; Zhu et al., 2012; Sekercioglu et al., 10 2008), the creation of novel species assemblages as not all species shift their ranges at the same rate or at all (Hobbs et al., 2009; Gilman et al., 2010; Williams and Jackson, 2007), or both. It is 12 therefore crucial to understand the dynamics of such climate induced range shifts to better inform 13 current and future conservation work.

While contemporary range shifts are a relatively new phenomenon, important insights can be gained from the related but distinct process of range expansion. Range expansions have been studied for decades, leading to a robust understanding of both the ecological (Skellam, 1951; Hastings et al., 2005) and evolutionary (Shine et al., 2011; Burton et al., 2010; Excoffier et al., 2009; Kubisch et al., 2014) mechanisms responsible for shaping such expansions. For example, while the speed of a

range expansion can be well approximated by a combination of the species' intrinsic growth rate and dispersal ability (Hastings et al., 2005), recent research demonstrates that evolution in both of these traits can have important implications for both the mean and variance of expansion speed through time (Weiss-Lehman et al., 2017; Ochocki and Miller, 2017; Szűcs et al., 2017; Phillips, 2015). As range shifts also involve a leading edge of population advance, they are likely to be affected by similar ecological and evolutionary mechanisms as have been shown to drive dynamics in range expansions (Hargreaves and Eckert, 2014). However, while such insights from range expansions are valuable for understanding and predicting dynamics of range shifts, it is important to recognize that these two processes have significant differences as well.

In particular, range expansions and range shifts due to climate change involve fundamentally distinct initial conditions. Range expansions typically begin from the successful establishment and subsequent spread of a small, founding population as typically occurs with the range expansions of invasive and reintroduced species (Hastings et al., 2005). Such founding populations represent samples from some larger source population and as such lack any initial spatial population structure. In contrast, range shifts involve entire populations with existing spatial structure within the initially stable range (Hargreaves and Eckert, 2014). Such spatial structure can take the form of local adaptation within the range, degree of transition in population size from range core to edge, or some combination of the two (Hargreaves and Eckert, 2014; Hargreaves et al., 2015; Henry et al., 2013).

These different aspects of spatial population structure have been shown to dramatically impact
the response of populations to climate change. For example, the nature of the gradient forming
the range edge (e.g. declines in birth rates vs. increases in extinction risk) can directly change the
probability of extinction a species faces during a climate driven range shift (Henry et al., 2013). A
population's risk of extinction during a range shift has also been related to local adaptation within
the range. Specifically, a broad environmental niche (i.e. little local adaptation) can decrease
a population's ability to track a changing climate if dispersal occurs in a stepping stone manner,
allowing some individuals to block dispersal of better adapted phenotypes (Atkins and Travis, 2010).
Local adaptation also has the potential to interact with dispersal evolution during climate change,

driving increased dispersal evolution in an asexual species (Hargreaves et al., 2015), though it is unclear if this result will hold for sexually reproducing species in which locally adapted genotypes interact via gene flow. As the gradient forming the range edge can influence the distribution of dispersal phenotypes within the range (Henry et al., 2013), it is likely that this would also result in an altered interaction between dispersal evolution and local adaptation in sexually reproducing species. In fact, pollen dispersal in flowering plants has been shown to dramatically reduce the likelihood of a range shift in response to climate change (Aguilée et al., 2016). Evolution during range shifts due to local adaptation and dispersal evolution are both potentially important drivers of range shift dynamics (Van Petegem et al., 2016) and it is therefore important to consider both when predicting the dynamics of populations responding to climate change.

Here, we use a complex, individual-based model incorporating evolution in both fitness and dispersal distance to determine the interaction between local adaptation and the starkness of the gradient at the range edge in driving population dynamics during climate change for a sexually reproducing species. Using this model, we simultaneously vary the strength of local adaptation, the starkness of the range boundary, and the speed of climate change to ascertain the relationship among them and test how dispersal evolution interacts with local adaptation to help or hinder a population's ability to track a changing climate. We additionally compare the dynamics of successful versus doomed simulated populations to understand the factors contributing to population extinction and identify warning signs for populations in danger of extinction as a result of a range shift.

68 2 Methods

$_{59}$ 2.1 Overview

70 **2.1.1** Purpose

This model tests an evolving population's ability to track a changing climate under a variety of conditions. Specifically, populations are simulated with (1) stark or gradual range boundaries, (2) strong or weak local selection pressures, and (3) varying speeds of climate change. In all simulations,

individual dispersal and relative fitness are defined by an explicit set of quantitative diploid loci subject to mutation, thus allowing both traits to evolve over time. All simulations begin with stable climate conditions for 2000 generations to allow the populations to reach a spatial equilibrium before the onset of climate change. Climate change is then modeled as a constant, directional shift in the location of environmentally suitable habitat (see the *Submodels* section below). Finally, simulations end with another short period of climate stability to assess the population's ability to persist and recover after shifting its range.

81 2.1.2 State variables and scales

The model simulates a population of male and female individuals characterized by spatial coordinates for their location and diploid loci for both fitness and dispersal. Space is modeled as a lattice of discrete patches overlaying a continuous Cartesian coordinate system with a fixed width along the y axis and without bounds on the x axis. To avoid edge effects along the y axis, the model employs wrapping boundaries such that if an individual disperses out of the landscape on one side, it appears at the opposite end of the same row of the landscape. Patches are defined by the location of the patch center in x and y coordinates and a patch width parameter defining the relationship between continuous Cartesian space and the discrete patches used for population dynamics (see the Submodels section below).

The abiotic environment is defined by a parameter denoting the center of the range, the slope

The abiotic environment is defined by a parameter denoting the center of the range, the slope of the range boundary, and the width of the range (See Figure 1). The range can then shift in space by altering the parameter defining the range center, which is how climate change is implemented in the model. Further, a gradient of phenotypic optimum values for fitness is imposed within the range to allow for local adaptation within the range boundaries. The severity of this gradient is defined by a separate parameter independent of the other environmental parameters (greater values of this parameter will lead to steeper gradients in phenotypic optima within the range).

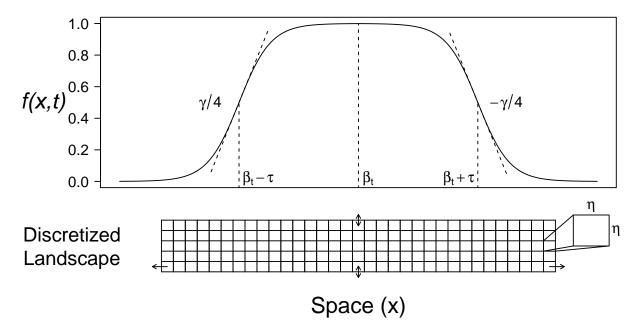


Figure 1: Environmental quality, as defined by f(x,t), over a spatial gradient. The parameters of f(x,t) are shown on the figure at significant points along the x axis. Additionally, the lattice of discrete patches in which population dynamics occur is shown beneath. This lattice of $\eta \times \eta$ patches are mapped to environmental conditions along the x axis using f(x,t) and are environmentally constant along the y axis. Dispersal is unbounded in the x direction and implemented with wrapping boundaries in the y direction.

98 2.1.3 Process overview and scheduling

Time is also modeled in discrete intervals defining single generations of the population. Within gg each generation, individuals first reproduce within their natal patch. After reproduction, all adults 100 die and the offspring disperse according to their dispersal kernels (Submodels), resulting in dis-101 crete, non-overlapping generations (See Figure 2). Reproduction occurs according to a stochastic 102 implementation of the classic Ricker model (Ricker, 1954) taking into account the mean fitness 103 of individuals within the patch (Submodels). Parental pairs form via random sampling of the lo-104 cal population (with replacement) weighted by individual fitness such that individuals with high 105 fitness are likely to produce multiple offspring while individuals with no fitness may not produce 106 any. Individuals inherit one allele from each parent at each loci, assuming independent segregation 107 and a mutation process (Submodels). After reproduction, all individuals in the parental generation 108 perish, forming discrete non-overlapping generations. The offspring then disperse according to their 109 dispersal traits to start the next generation in their post dispersal locations. 110

111 2.2 Design concepts

112 **2.2.1** Emergence

Emergent phenomena in this model include the spatial equilibrium of population abundances and trait values within the stable range, the demographic dynamics of the shifting population during climate change, and the evolutionary trajectories of both fitness and dispersal traits during climate change. These are all examined in the context of their impact on the population's ability to keep pace with the changing climate.

118 2.2.2 Stochasticity

All biological processes in this model are stochastic including realized population growth in each patch, dispersal distances of each individual, and inheritance of loci. Environmental parameters are fixed, however, and the process of climate change (i.e. the movement of environmentally suitable habitat through time) is deterministic. Thus, the model removes the confounding influence of environmental stochasticity to focus on demographic and evolutionary dynamics of range shifts.

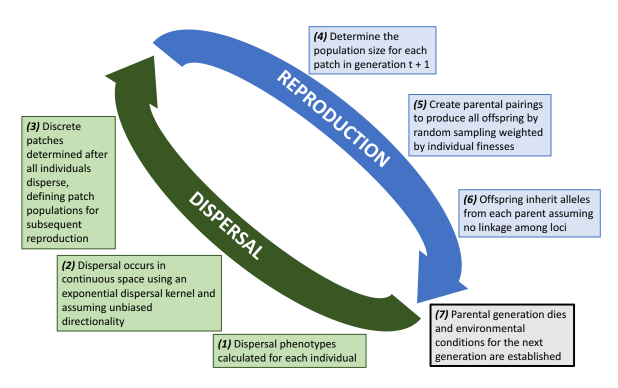


Figure 2: The life cycle of simulated populations is shown divided between events contributing to reproduction and dispersal. Each generation begins with new offspring dispersing according to their phenotype, after which reproduction occurs in local populations defined by the discrete lattice. After reproduction, all parental individuals perish, resulting in discrete, non-overlapping generations.

24 2.2.3 Interactions

Individuals in the model interact via mating and density-dependent competition within patches.

Other important interactions are the relationship between dispersal evolution and local adaption,
particularly in edge populations, and how this relationship impacts a population's ability to avoid
extinction and track a changing climate. These interactions are examined in the context of different
configurations of the initial, stable range and different speeds of climate change.

130 2.2.4 Desired output

After each model run, full details of all surviving individuals at the last time point are recorded (spatial coordinates and loci values for both traits). If a population went extinct during the model run, the time of extinction is recorded. Throughout the simulation, certain aggregated values are calculated and recorded for each occupied patch including the population size and the mean and variance of allele values for each trait.

136 2.3 Details

137 2.3.1 Initialization

The following parameters are set at the beginning of each simulation and form the initial conditions 138 of the model: the mean and variance for allele values of each trait, population size, location of 139 environmentally suitable habitat, number of generations for the pre-, post-, and during climate 140 change periods of the simulation, and all other necessary parameters for the submodels defined 141 below. Simulated populations are initialized in the center of the range and allowed to spread and 142 equilibrate throughout the range during the period of stable climate conditions. This ensures that the populations reacting to a changing climate truly represent the expected spatial distribution for 144 a given range, rather than the initial parameter values used in the simulation. See Table 1 for a 145 full list of all parameter values used in the simulations described here.

147 2.3.2 Submodels

Environmentally suitable habitat Environmentally suitable habitat is determined by the population's carrying capacity as it ranges in space (K_x) . The maximum achievable carrying capacity (K_{max}) occurs in the center of the species' range and declines with increasing distance from the center. Specifically, the carrying capacity at a location x is defined as a combination of K_{max} and a function f(x,t), where f(x,t) ranges from 1 in the range center to 0 far away from the center and is defined as follows

$$f(x,t) = \begin{cases} \frac{e^{\gamma(x-\beta_t+\tau)}}{1+e^{\gamma(x-\beta_t+\tau)}} & x \le \beta_t \\ \frac{e^{-\gamma(x-\beta_t-\tau)}}{1+e^{-\gamma(x-\beta_t-\tau)}} & x > \beta_t \end{cases}$$
 (1)

where $beta_t$ defines the center of the area of suitable habitat at time t, τ sets the width of the range, and γ affects the slope of the function at the range boundaries (See Figure 1). To understand the relationship between γ and the slope of f(x,t) at the range boundary, the partial derivative of f(x,t) over the spatial dimension can be shown to be

$$f(x,t) = \begin{cases} \frac{e^{\gamma(x-\beta_t+\tau)}}{1+e^{\gamma(x-\beta_t+\tau)}} & x \le \beta_t\\ \frac{e^{-\gamma(x-\beta_t-\tau)}}{1+e^{-\gamma(x-\beta_t-\tau)}} & x > \beta_t \end{cases}$$
 (2)

yielding a derivative of $\pm \frac{\gamma}{4}$ at the inflection points on either side of the range center $(x = \beta_t \pm \tau)$.

Population dynamics occur within discrete patches, so to calculate a K_x value for a discrete patch from the continuous function f(x,t), we use another parameter defining the spatial scale of each patch $(\eta;$ See Figure 1). The local carrying capacity of a patch centered on x (K_x) is then calculated as the mean of f(x,t) over the interval of the patch multiplied by K_{max} .

$$K_x = \frac{K_{max}}{\eta} \int_{x-\frac{\eta}{2}}^{x+\frac{\eta}{2}} f(x,t) dx \tag{3}$$

By varying the parameters defining f(x,t), we can change both the total achievable carrying capacity of the population throughout the range (by altering both τ and γ) and the slope at which K_x declines to 0 (by altering γ). We are primarily interested in the effect of altering the slope at which K_x declines at the range boundaries, however as doing so can also alter the total achievable carrying capacity of the population, we control for this confounding factor by fixing the total area under the curve f(x,t). The indefinite integral of f(x,t) can be shown to be

$$\int_{-\infty}^{\infty} f(x,t)dx = \frac{2ln(e^{\gamma\tau} + 1)}{\gamma} \tag{4}$$

which can be solved for τ . Thus, if the total area under the curve is fixed, an appropriate value of τ can be calculated for each value of γ .

Thus, γ and τ are both fixed within a given simulation and $beta_t$ (the location of the center of suitable habitat) is used to simulate climate change. During the periods before and after climate change β_t is constant, but to simulate climate change it varies with time as follows

$$\beta_t = \nu \eta(t - \hat{t}) \tag{5}$$

where ν is the velocity of climate change per generation in terms of discrete patches, η is the spatial scale of each patch, t is the current generation, and \hat{t} is the last generation of stable climatic conditions before the onset of climate change.

Local adaptation To allow an arbitrary degree of local adaptation within the range, the local phenotypic optima $(z_{opt,x})$ is set as follows

$$z_{opt,x} = \lambda(x - \beta_t) \tag{6}$$

where λ defines the strength of local selection with values close to 0 resulting in little to no change in phenotypic optimum across the range and values of greater magnitude resulting in large differences in phenotypic optima across the range. Individual fitness $(w_{i,x})$ values are then calculated according to the following equation assuming stabilizing selection

$$w_{i,x} = e^{\frac{-(z_i - z_{opt,x})^2}{2\omega^2}} \tag{7}$$

where ω defines the strength of stabilizing selection and z_i is an individual's fitness phenotype (Lande, 1976). All loci are assumed to contribute additively to the phenotype with no dominance or epistasis, meaning an individual's phenotype is simply the sum of the individual's allele values for the fitness trait.

Population dynamics Population growth within each patch is modeled with a stochastic implementation of the classic Ricker model (Ricker, 1954; Melbourne and Hastings, 2008). To account for fitness effects on population growth, expected population growth is scaled by the mean relative fitness of individuals within the patch (\bar{w}_x) . The expected number of new offspring in patch x at time t+1 is then given by

$$\hat{N}_{t+1,x} = \bar{w}_x F_{t,x} \frac{R}{\psi} e^{\frac{-RN_{t,x}}{K_x}}$$
(8)

where $F_{t,x}$ is the number of females in patch x at time t, R is the intrinsic growth rate for the population, ψ is the expected sex ratio of the population, $N_{t,x}$ is the number of individuals (males and females) in patch x at time t, and K_x is the local carrying capacity based on the environmental conditions. To incorporate demographic stochasticity, the realized number of offspring for each patch is then drawn from a Poisson distribution.

$$N_{t+1,x} \sim Poisson(\hat{N}_{t+1,x}) \tag{9}$$

Parentage of the offspring is then assigned by random sampling of the local male and female population. The sampling is weighted by individual fitness and occurs with replacement so highly fit individuals are likely to have multiple offspring while low fitness individuals may not have any. Each offspring inherits one allele per locus from each parent, assuming no linkage among loci. After reproduction, all members of the previous generation die and the offspring disperse to begin the next generation.

Mutation Inherited alleles are subject to mutation such that some offspring might not inherit identical copies of certain alleles from their parents. The mutation process is defined by two parameters for each trait T: the diploid mutation rate (U^T) and the mutational variance (V_m^T) .

Using these parameters along with the number of loci defining trait $T(L^T)$, the per locus probability of a mutation is

$$\frac{U^T}{2L^T} \tag{10}$$

208 Mutational effects are drawn from a normal distribution with mean 0 and a standard deviation of

$$\sqrt{V_m^T U^T} \tag{11}$$

By defining the mutation process in this manner rather than setting a probability of mutation and mutational effect directly, similar mutational dynamics can be imposed regardless of the number of loci used in the simulation.

Dispersal Finally, individuals disperse according to an exponential dispersal kernel defined by each individual's dispersal phenotype. An individual's dispersal phenotype is the expected dispersal distance and is given by

$$d_i = \frac{D\eta e^{\rho \Sigma L^D}}{1 + e^{\rho \Sigma L^D}} \tag{12}$$

where D is the maximum expected dispersal distance in terms of discrete patches, η is the spatial 215 scale of discrete patches, ρ is a constant determining the slope of the transition between 0 and D, 216 and the summation is the sum of all alleles contributing to dispersal. Thus, as with fitness, loci are 217 assumed to contribute additively with no dominance or epistasis. The expected dispersal distance, 218 d_i is then used to draw a realized distance from an exponential dispersal kernel. The direction 219 of dispersal is drawn from a uniform distribution bounded by 0 and 2π . If a dispersal trajectory 220 takes an individual outside the bounds of the landscape in the y axis, the individual reappears 221 at the same x coordinate but the opposite end of the y axis, thus wrapping the top and bottom 222 edges of the landscape to avoid edge effects. Dispersal occurs from the center of each patch and the individual's new patch is then determined according to its location in the overlaid grid of η x 224 η patches (see Figure 1).

2.4 Simulation descriptions

To explore the interacting role of local adaptation and the slope of the decline in population abundance at the range edge on trait evolution and the dynamics of range shifts, we simulated 100 populations each over a range of parameter values varying both the degree of local adaptation and the decline in population abundance at the range edge (See Table 1). Additionally, for each parameter combination we simulated three different speeds of climate change. All simulations were performed in R version 3.4.4 (Team, 2000) and the code is available at https://github.com/tpweiss06/ShiftingSlopes.

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