Genetic paths to evolutionary rescue and the distribution of fitness effects along them

Matthew M Osmond*,1, Sarah P Otto* and Guillaume Martin[†]

*Biodiversity Centre & Department of Zoology, University of British Columbia, †Institut des Sciences de l'Evolution de Montpellier, Université Montpellier II

ABSTRACT

17

23

24

25

26

28

- The past century has seen substantial theoretical and empirical progress on the genetic basis of adaptation. Over this same period a pressing need to prevent the evolution of drug resistance has uncovered much about the potential genetic basis of persistence in declining populations. However, we have little theory to predict and generalize how persistence by sufficiently rapid adaptation might be realized in this explicitly demographic scenario. Here we use Fisher's geometric model with absolute fitness to begin a line of theoretical inquiry into the genetic basis of evolutionary rescue, focusing here on asexual populations that adapt through *de novo* mutations. We show how the dominant genetic path to rescue switches from a single mutation to multiple as mutation rates and the severity of the environmental change increase. In multi-step rescue, intermediate genotypes that themselves go extinct provide a 'springboard' to rescue genotypes. Comparing to a scenario where persistence is assured, our approach allows us to quantify how a race between evolution and extinction leads to a genetic basis of adaptation that is composed of fewer loci of larger effect. We hope this work brings awareness to the impact of demography on the genetic basis of adaptation.
- KEYWORDS Antimicrobial drug resistance; Evolutionary escape; Fisher's geometric model; Genetic basis of adaptation; Mathematical theory

our understanding of the genetic basis of adaptation is rapidly improving due to the now widespread use of experimental evolution and genomic sequencing (see examples in Bell 2009; Stapley *et al.* 2010; Dettman *et al.* 2012; Schlötterer *et al.* 2015). A recurrent observation, especially in asexual microbes, is that the more novel the environment and the stronger the selection pressure, the more likely it is that adaptation primarily proceeds by fewer mutations of larger effect (i.e., that adaptation is oligogenic *sensu* Bell 2009). An extreme case is the evolution of drug resistance, which is often achieved by just one or two mutations (e.g., Bataillon *et al.* 2011; Pennings *et al.* 2014).

However, drugs, and other sufficiently novel environments, will often induce not only strong selection but also population decline. Such declines hinder both the production and maintenance of genetic variation (Otto and Whitlock 1997), thus impeding evolution and threatening extinction. In this scenario, drug resistance evolution is a particular instance of the more general phenomenon of evolutionary rescue (Gomulkiewicz and

Holt 1995; Bell 2017), where persistence requires sufficiently fast adaptive evolution.

Most theory on the genetics of adaptation (reviewed in Orr 2005) assumes constant population size and therefore does not capture the characteristic 'race' between adaptation and extinction that occurs during evolutionary rescue. Many models have been created to describe this race (reviewed in Alexander et al. 2014) but so far largely focus on two extreme genetic bases, both already introduced in Gomulkiewicz and Holt (1995): rescue is either caused by minute changes in allele frequencies across many loci in sexuals (i.e., the infinitesimal model; Fisher 1918) or by the substitution of a single large effect 'resistance' mutation (e.g., one locus, two allele models). We therefore lack a theoretical framework for the genetic basis of evolutionary rescue that captures the arguably more realistic situation where an intermediate number of mutations are at play (but see exceptions below). The absence of such a framework prevents us from predicting the number of mutations that evolutionary rescue will take and the distribution of their effect sizes. The existence of a more complete framework could therefore provide valuable information to those investigating the genetic basis of drug resistance (e.g., the number and effect sizes of mutations to look for) and would

extend our understanding on the genetic basis of adaptation to cases of non-equilibrial demography (i.e., rapid evolution and eco-evo dynamics). 119

57

70

71

82

83

100

101

102

104

105

106

112

113

114

115

116

Despite these gaps in the theory on the genetic basis of evolutionary rescue, there is a wealth of data. For example, the genetic basis of resistance to a variety of drugs is known in many species of bacteria (reviewed in MacLean et al. 2010), fungi (reviewed in Robbins et al. 2017), and viruses (reviewed in Yilmaz et al. 2016). This abundance of data reflects both the applied need to prevent drug resistance and the relative ease of isolating the genotypes that survive (hereafter "rescue genotypes"), e.g., in a Luria-Delbrück fluctuation assay (reviewed in Bataillon and Bailey 2014). Assaying fitness in the environment used to isolate mutants (e.g., in the drug) then provides the distribution of fitness effects of potential rescue genotypes. Additional data on the genetic basis of drug resistance arise from the construction of mutant libraries (e.g., Weinreich et al. 2006) and the sequencing of natural populations (e.g., Pennings et al. 2014). Together, the data show that resistance often appears to arise by a single mutation (e.g., MacLean and Buckling 2009; Lindsey et al. 2013; Gerstein et al. 2012) but not always (e.g., Bataillon et al. 2011; Pennings et al. 2014; Gerstein et al. 2015; ?). The data also indicate that the fitness effect of rescue genotypes is more often large than small, creating a hump-shaped distribution of selection coefficients (e.g., Kassen and Bataillon 2006; MacLean and Buckling 2009; Gerstein et al. 2012; Lindsey et al. 2013; Gerstein et al. 2015) that is similar in shape to that proposed by Kimura (1983) but with a lower bound that is often much greater than zero.

Theory on evolutionary rescue (reviewed in Alexander et al. 2014) has primarily focused on the probability of rescue rather than its genetic basis. However, a few studies have varied the potential genetic basis enough to make some inference about how evolutionary rescue is likely to happen. For instance, in the 151 context of pathogen host-switching, Antia et al. (2003) numerically explored the probability of rescue starting from a single 153 ancestral individual when n intermediate mutations (each occurring from the previous with the same probability) are required. 155 They focused on the case of rescue by two mutations (n = 1), ₁₅₆ when the intermediate genotype had a growth rate that was either the same as the ancestor, below that of the ancestor, or intermediate between the ancestor and the rescue genotype. In all cases they found that rescue became less likely as the number of intermediate mutations increased, suggesting that rescue will generally proceed by the fewest possible mutations. This framework was expanded greatly by Iwasa et al. (2004a) who derived approximations for the probability of rescue with an arbitrary number of mutational steps, considering arbitrary mutational networks (i.e., allowing mutations between any two genotypes) and standing genetic variation in an ancestral population. Assuming the probability of mutation between any two genotypes is of the same order, they showed that genetic paths with fewer mutational steps contributed more to the probability of rescue, again suggesting rescue will occur by the fewest possible mutations. Iwasa et al. (2004a) also found that multiple simultaneous mutations (i.e., arising in the same meiosis) can contribute more $\,^{169}$ to rescue than paths that gain these same mutations sequen- 170 tially (i.e., over multiple generations) when the growth rates of 171 the intermediate mutations are small enough, suggesting that 172 rare large mutations can be the most likely path to rescue when 173 the population is very maladapted or the rescue phenotype is 174 complex (i.e., there is a fitness valley separating the wildtype 175 and rescue genotype). This point was previously demonstrated by Alexander and Day (2010), who emphasized that multiple simultaneous mutations become the dominant path to rescue in the most challenging environments. As a counterpoint, Uecker and Hermisson (2016) explored a greater range of fitness values in a two-locus two-allele model, showing that, with standing genetic variation, rescue by sequential mutations at two loci (two mutational steps) can be more likely than rescue by mutation at a single locus (one simultaneous mutational step) when the wildtype is very maladapted, as, in the former case, the single mutants can act as a buffer in the face of environmental change. In summary, current theory indicates that the genetic basis of rescue hinges on the chosen set of genotypes, their fitnesses, and the mutation rates between them. So far these choices have been in large part arbitrary or chosen for mathematical convenience.

Here we follow the lead of Anciaux et al. (2018) by implicitly choosing the range of genotypes, fitnesses, and mutation rates from an empirically-justified fitness-landscape model (Tenaillon 2014). In particular, we use Fisher's geometric model to describe adaptation following an abrupt environmental change that instigates population decline. There are two key differences between this approach and earlier models using Fisher's geometric model (e.g., Orr 1998): here 1) the dynamics of each genotype depends only on their absolute fitness (instead of only on their relative fitness) and 2) multiple mutations can segregate simultaneously (instead of assuming only sequential fixation), allowing multiple mutations to fix – in our case, rescue the population – together as a single haplotype (i.e., stochastic tunnelling, Iwasa et al. 2004b). In this non-equilibrium scenario, variation in absolute fitness, which allows population size to vary, can create feedbacks between demography and evolution, which could strongly impact the genetic basis of adaptation relative to the constant population size case. In contrast to Anciaux et al. (2018), our focus here is on the genetic basis of evolutionary rescue and we also explore the possibility of rescue by mutant haplotypes containing more than one mutation. In particular, we ask: (1) How many mutational steps is evolutionary rescue likely to take? and (2) What is the expected distribution of fitness effects of the surviving genotypes and their component mutations?

We first introduce the modelling framework before summarizing our main results. We then present the mathematical analyses we have used to understand these results and end with a discussion of our key findings.

Data availability

Code used to derive analytical and numerical results and produce figures (referred to here as File S1; Mathematica, version 9.0; Wolfram Research Inc. 2012) and code used to run individual-based simulations (Python, version 3.5; Python Software Foundation), as well as simulation data and freely accessible versions of File S1 (CDF and PDF), are available at https://github.com/mmosmond/GeneticBasisOfRescue.

Model

Fisher's geometric model

We map genotype to phenotype to fitness using Fisher's geometric model, originally introduced by Fisher (1930, p. 38-41) and reviewed by Tenaillon (2014). In this model each genotype is characterized by a point in n-dimensional phenotypic space, \vec{z} . We ignore environmental effects, and thus the phenotype is the breeding value. Some phenotype, \vec{o} , has maximum

fitness and fitness declines monotonically as phenotypes de- 234 part from \vec{o} . Without loss of generality, the *n* phenotypic axes are chosen and scaled such that fitness can be described by a multivariate normal distribution with variance 1 in each dimension, no covariance, and height W_{max} . Thus the fitness of phenotype \vec{z} is $W(\vec{z}) = W_{max} \exp(-||\vec{z} - \vec{o}||^2/2)$, where $||ec{z}-ec{o}||=\sqrt{\sum_{i=1}^n(z_i-o_i)^2}$ is the Euclidean distance of $ec{z}$ from ²⁴⁰ the optimum, \vec{o} . Here we are interested in absolute fitness; we take $\ln[W(\vec{z})] = m(\vec{z}) = m_{max} - ||\vec{z} - \vec{o}||^2/2$ to be the continuous-time growth rate (m is for Malthusian fitness) of 242 phenotype \vec{z} . We ignore density- and frequency-dependence ₂₄₃ in $m(\vec{z})$ for simplicity. The fitness effect, i.e., selection coefficient, of phenotype z' relative to z in a continuous-time model 245 is exactly $s = \log[W(z')/W(z)] = m(z') - m(z)$ (Martin and 246 Lenormand 2015). This is approximately equal to the selection coefficient in discrete time (W(z')/W(z) - 1) when selection is weak (W(z') - W(z) << 1).

178

179

180

18

184

185

186

187

188

189

190

192

193

194

195

197

200

20

202

203

20

208

209

210

215

216

217

218

219

22

222

223

224

231

232

To make analytical progress we use the isotropic version of Fisher's geometric model, where mutations (in addition to selection) are assumed to be uncorrelated across the scaled traits. Universal pleiotropy is also assumed, so that each mutation affects all scaled phenotypes. In particular we use the "classic" form of Fisher's geometric model (Harmand *et al.* 2017), where the probability density function of a mutant phenotype is multivariate normal, centred on the current phenotype, with variance λ in each dimension and no covariance. This implies a continuum-of-alleles (Kimura 1965), i.e., phenotype is continuous and each mutation is unique. Mutations are assumed to be additive in phenotype, which induces epistasis in fitness as fitness is a non-linear function of phenotype. We assume asexual reproduction, i.e., no recombination., which is appropriate for many cases of antimicrobial drug resistance.

An obvious and important extension would be to relax the simplifying assumptions of isotropy and universal pleiotropy, which we leave for future work. Note that mild anisotropy yields the same bulk distribution of fitness effects as an isotropic model with fewer dimensions (Martin and Lenormand 2006), but this does not extend to the tails of the distribution. Therefore, whether anisotropy can be reduced to isotropy with fewer dimensions in the case of evolutionary rescue, where the tails are essential, is unknown.

Given this phenotype to fitness mapping and phenotypic distribution of new mutations, the distribution of fitness effects (and therefore growth rates) of new mutations can be derived exactly. Let m be the growth rate of some particular focal genotype and m' the growth rate of a mutant immediately derived from it. Then let $s_0 = m_{max} - m$ be the selective effect of a mutant with the optimum genotype and s = m' - m the selective effect of the mutant with growth rate m'. The probability density function of the selective effects of new mutations, s, is then given by equation 3 in Martin and Lenormand (2015). Converting fitness effects to growth rate (m' = s + m), the probability density function for mutant growth rate m' from an ancestor with growth rate m is (cf. equation 2 in Anciaux et al. 2018)

$$f(m'|m) = \frac{2}{\lambda} f_{\chi_n^2} \left(\frac{2(m_{max} - m')}{\lambda}, \frac{2(m_{max} - m)}{\lambda} \right), \quad (1)$$

281

where $f_{\chi^2_n}(x,c)$ is the probability density function over positive real numbers x of $\chi^2_n(c)$, a non-central chi-square deviate with n degrees of freedom and noncentrality c>0 (equation 26.4.25 in Abramowitz and Stegun 1972).

Lifecycle

We are envisioning a scenario where N_0 wildtype individuals, each of which have phenotype \vec{z}_0 , experience an environmental change, causing population decline, $m_0 \equiv m(\vec{z}_0) < 0$. Each generation, an individual with phenotype \vec{z} produces a Poisson number of offspring, with mean $\ln[m(\vec{z})]$, and dies. Each offspring mutates with probability U, with mutations distributed as described above (see Fisher's geometric model).

Probability of rescue

Let p_0 be the probability that a given wildtype individual is "successful", i.e., has descendants that rescue the population. The probability of rescue is then one minus the probability that none of the initial wildtype individuals are successful,

$$P = 1 - (1 - p_0)^{N_0} \approx 1 - \exp(-N_0 p_0),$$
 (2)

where the approximation assumes small p_0 and large N_0 . What remains is to find p_0 .

Summary of Results

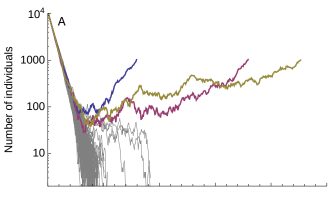
We start with a heuristic explanation of our main results before turning to more detailed derivations in the next section.

Rescue by multiple mutations

A characteristic pattern of evolutionary rescue is a "U"-shaped population size trajectory (e.g., Orr and Unckless 2014). This is the result of an exponentially-declining wildtype genotype being replaced by an exponentially-increasing mutant genotype. On a log scale this population size trajectory becomes "V"-shaped (we denote it a 'V-shaped log-trajectory'). On this scale, the population declines at a constant rate (producing a line with slope $m_0 < 0$) until the growing mutant subpopulation becomes relatively common, at which point the population begins growing at a constant rate (a line with slope $m_1 > 0$). This characteristic V-shaped log-trajectory is observed in many of our simulations where evolutionary rescue occurs (Figure 1A). In contrast, when the wildtype declines faster and the mutation rate is larger we sometimes see 'U-shaped log-trajectories' (e.g., the red and blue replicates in Figure 2A). Here there are three phases instead of two; the initial rate of decline (a line with slope $m_0 < 0$) is first reduced (transitioning to a line with slope $m_1 < 0$) before the population begins growing (a line with slope $m_2 > 0$).

As expected, V-shaped log-trajectories are the result of a single mutation creating a genotype with a positive growth rate that escapes loss when rare and rescues the population (Figure 1B), i.e., 1-step rescue. U-shaped log-trajectories, on the other hand, occur when a single mutation creates a genotype with a negative (or potentially very small positive) growth rate, itself doomed to extinction, which out-persists the wildtype and gives rise to a double mutant genotype that rescues the population (Figure 2B), i.e., 2-step rescue. These two types of rescue comprise the overwhelming majority of rescue events observed in our simulations, across a wide range of wildtype decline rates (e.g., Figure 3).

Of course, with sufficiently high mutation rates rescue by 3 or more mutations comes to dominate (Figure S1). It has recently been suggested that when the mutation rate, U, is substantially less than a critical value, $U_C = \lambda n^2/4$, we are in a "strong selection, weak mutation" regime where essentially all mutations arise on a wildtype background (Martin



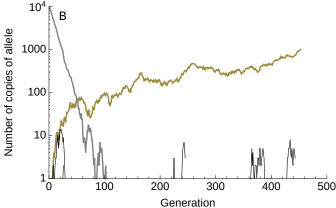
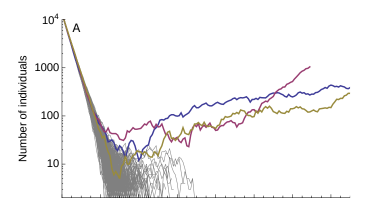


Figure 1 Typical dynamics with a relatively slow wildtype decline and a small mutation rate ($m_0 = -0.1$, $U = 10^{-4}$). **(A)** Population size trajectories on a log scale. Each line is a unique replicate simulation (100 replicates). Replicates that went extinct are grey, replicates that were rescued are in colour (and are roughly V-shaped). **(B)** The number of individuals with a given derived allele, again on a log scale, for the yellow replicate in **A**. The number of individuals without any derived alleles (wildtypes) is shown in grey, the rescue mutation is shown in yellow, and all other mutations are shown in black. Other parameters: n = 4, $\lambda = 0.005$, $m_{max} = 0.5$.

and Roques 2016), consistent with the House of Cards approximation (Turelli 1984, 1985), and thus rescue will occur by a single mutation of large effect (Anciaux *et al.* 2018). In the other extreme, when $U >> U_C$, we are in a weak selection, strong mutation regime where many mutations arise on the same background, creating a multivariate normal phenotypic distribution (Martin and Roques 2016), consistent with the Gaussian approximation (Kimura 1965; Lande 1980), and thus rescue will occur by many mutations of small effect (Anciaux *et al.* 2019). As shown in Figure 3 (where $U = U_C/10$) and Figure S1 (which spans $U_C = 0.02$), rescue by a small number of mutations (but more than one) can become commonplace in the transition zone, $U \sim U_C$, where there are often a considerable number of cosegregating mutations (e.g., Figure 2B, where $U = U_C/2$).

The probability of k-step rescue

Approximations for the probability of 1-step rescue under the strong selection, weak mutation regime were derived by Anci- 318 aux *et al.* (2018). Here we extend this study by exploring the contribution of k-step rescue, deriving approximations for the 320



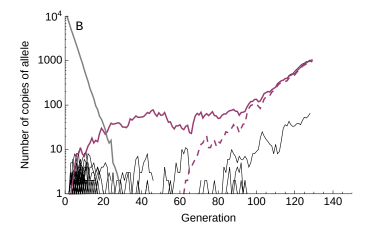


Figure 2 Typical dynamics with a relatively fast wildtype decline and a large mutation rate ($m_0 = -0.3$, $U = 10^{-2}$). (A) Population size trajectories on a log scale. Each line is a unique replicate simulation (500 replicates). Replicates that went extinct are grey, replicates that were rescued are in colour. Note that the blue and red replicates are cases of 2-step rescue (and roughly U-shaped), while the yellow replicate is 1-step rescue (and therefore V-shaped). (B) The number of individuals with a given derived allele, again on a log scale, for the red replicate in **A**. The number of individuals without any derived alleles (wildtypes) is shown in grey, the rescue mutations are shown in red, and all other mutations in black. Here a single mutant with growth rate less than zero arises early and outlives the wildtype (solid red). A second mutation then arises on that background (dashed red), making a double mutant with a growth rate greater than zero that rescues the population. Other parameters: n = 4, $\lambda = 0.005$, $m_{max} = 0.5$.

probability of such events, as well as dissecting the genetic basis of both 1- and 2-step rescue in terms of the distribution of fitness effects of rescue genotypes and their component mutations.

Although requiring a sufficiently beneficial mutations to arise on a rare mutant genotypes doomed to extinction, multi-step evolutionary rescue can be the dominant form of rescue when the wildtype is sufficiently maladapted (Figures 3 and S1). Indeed, on this fitness landscape, the probability of producing a rescue genotype in one mutational step mutant drops very sharply with maladaptation (Anciaux et al. 2018); mutants with intermediate growth rates can thus be a "springboard" – albeit not always a very bouncy one – from which rescue mutants are produced. These intermediates contribute more as mutation

313

315

289

290

29

293

294

295

297

300

301

302

303

304

305

306

307

rates and the decline rate of the wildtype increase (Figures 3 and S1); the former because double mutants become more likely and the latter because the springboard becomes more necessary. With a large enough number of wildtype individuals or a high enough mutation rate (Figure S1), multi-step rescue can not only be more likely than 1-step, but also very likely in an absolute sense.

Alternative regimes of 2-step rescue

321

322

323

324

325

328

333

334

335

336

337

339

340

341

342

348

349

350

35

353

354

355

369

370

371

372

378

370

2-step rescue can occur through first-step mutants with a wide range of growth rates. As shown below, these first-step mutants can be divided into three regimes: "sufficiently subcritical", "sufficiently critical", and "sufficiently supercritical" (we will often drop "sufficiently" for brevity; Figure 4). Critical first-step mutants are defined by having growth rates close enough to zero that occasionally such a mutation will, by chance, persist for long enough that it will almost certainly produce successful double mutants. Subcritical first-step mutants are then defined by having growth rates that are negative enough to prevent such long persistence times. Their dynamics are roughly deterministic; mutations conferring growth rates closer to zero persist longer but are less likely to arise from the wildtype. Similarly, supercritical first-step mutants are defined by having positive growth rates that are large enough to prevent long persistence times once conditioned on extinction; conditioned on extinction these genotypes behave like subcritical mutations with a growth rate of the same absolute value (Maruyama and Kimura 1974). As with subcritical first-step mutants, the dynamics of supercritical mutants are relatively deterministic, but in this case persistence times and mutational input from the wildtype both favour growth rates closer to zero.

The relative contribution of each regime changes with both the initial degree of maladaptation and the mutation rate (Figures 5 and S2). When the wildtype is very maladapted (relative to mutational variance), most 2-step rescue events occur through subcritical first-step mutants (Figure 5A), which are more abundant than critical or supercritical mutants and yet persist longer than the wildtype. If the initial maladaptation is milder, however, the slow decline of the wildtype buys time for critical and supercritical mutations to arise and contribute to 2-step rescue, and also increases the rate at which they are produced. The rate of mutation also plays an interesting role in determining the relative contributions of each regime (Figures 5B and S2). When mutations are rare, only first-step mutations that are very nearly neutral ($m \sim 0$) will persist long enough to give rise to a 2-step rescue mutation. As the mutation rate increases, however, the range of first-step mutant growth rates that can persist long enough to lead to 2-step rescue widens (because fewer individuals carrying the first-step mutation are needed to produce a 423 successful double mutant).

The distribution of fitness effects among rescue mutations

Mutants causing 1-step rescue have growth rates that cluster around small positive values ($m \gtrsim 0$; Figure 6). Consequently, the distribution of fitness effects (DFE) is shifted to the right relative to the constant population size case, beginning at $s=m-m_0 \ge -m_0 > 0$. As a result of this increased threshold, the 1-step rescue DFE has a smaller variance than both the DFE of random mutations and the DFE of mutations that establish in a constant population (Kimura 1983). Further, unlike these other distributions, the variance in the DFE under 1-step rescue decreases as the rate of wildtype decline increases, due to rescue

being restricted to a smaller proportion of the available mutants.

The DFE of genotypes that cause 2-step rescue (the combined effect of two mutations) is also clustered at small positive values, but it has a variance that is less affected by the rate of wildtype decline (Figure 6). This is because double mutant rescue genotypes are created via first-step mutant genotypes that have larger growth rates than the wildtype (i.e., are closer to the optimum), allowing them to create double mutants with a larger range of positive growth rates.

Finally, we can also look at the distribution of growth rates among firsts-step mutations that lead to 2-step rescue (Figures 7 and S2), i.e., 'springboard mutants'. Here there are two main factors to consider: 1) the probability that a mutation with a given growth rate arises on the wildtype background but does not by itself rescue the population and 2) the probability that such a mutation persists long enough for a sufficiently beneficial second mutation to arise on that same background and together rescue the population. As explained above, increasing the rate of wildtype decline (or decreasing the rate of mutation) shifts the contribution from critical first-step mutants to subcritical first-step mutants, lowering the mode and increasing the variance.

Finally, note that, given 2-step rescue, the growth rate of both the first and second mutation may be negative when alone in the wildtype background. This potentially obscures empirical detection of the individual mutations involved in evolutionary rescue.

Mathematical Analysis

The probability of k-step rescue

Generic expressions for the probability of 1- and 2-step rescue were given by Martin *et al.* (2013), using a diffusion approximation of the underlying demographics. The key result that we will use is the probability that a single copy of a genotype with growth rate m, itself fated for extinction but which produces rescue mutants at rate $\Lambda(m)$, rescues the population (equation S1.5 in Martin *et al.* 2013). With our lifecycle this is (c.f., equation A.3 in Iwasa *et al.* 2004a)

$$p(m, \Lambda(m)) = 1 - \exp\left[|m|\left(1 - \sqrt{1 + \frac{2\Lambda(m)}{m^2}}\right)\right]. \tag{3}$$

We can therefore use $p_0 = p(m_0, \Lambda(m_0))$ as the probability that a wildtype individual has descendants that rescue the population and what remains in calculating the total probability of rescue (Equation 2) is $\Lambda(m_0)$. We break this down by letting $\Lambda_i(m)$ be the rate at which rescue genotypes with i mutations are created; the total probability of rescue is then given by Equation 2 with $p_0 = p(m_0, \sum_{i=1}^\infty \Lambda_i(m_0))$.

In 1-step rescue, $\Lambda_1(m_0)$ is just the rate of production of rescue mutants directly from a wildtype genotype. This is the probability that a wildtype gives rise to a mutant with growth rate m (given by $Uf(m|m_0)$) times the probability that a genotype with growth rate m establishes. Again approximating our discrete time process with a diffusion process, the probability that a lineage with growth rate m << 1 establishes, ignoring further mutation, is (Martin $et\ al.\ 2013$)

$$p_{est}(m) \approx \begin{cases} 0 & m \le 0\\ 1 - \exp(-2m) & m > 0 \end{cases}$$
 (4)

This reduces to the $2(s + m_0)$ result in Otto and Whitlock (1997) when $m = s + m_0$ is small, which further reduces to 2s in a

Symbol	Meaning
n	number of (scaled) phenotypic dimensions
λ	variance in mutant phenotypes along each dimension
m_{max}	maximum growth rate
f(m' m)	distribution of growth rates among mutants from a genotype with growth rate m (eq. 1)
U	per genome mutation probability
N_0	initial number of wildtype individuals
m_0	wildtype growth rate
p_0	probability that an initial wildtype individual has descendants that rescue the population
P	probability of rescue (eq. 2)
$p(m,\Lambda(m))$	probability a genotype with growth rate <i>m</i> , itself fated for extinction, has descendants that rescue the population (eq. 3)
$p_{est}(m)$	probability a genotype with growth rate m establishes, i.e., rescues the population (eq. 4)
$\Lambda(m)$	probability that an individual with growth rate m produces a mutant that has descendants that rescue the population
$\Lambda_i(m)$	probability that an individual with growth rate m produces a mutant that has descendants with $i-1$ additional mutations that rescue the population
$\Lambda_{2}^{(-)}(m), \ \Lambda_{2}^{(0)}(m), \ \Lambda_{2}^{(+)}(m)$	probability that an individual with growth rate m produces sufficiently subcritical, critical, or supercritical first-step mutants that eventually lead to 2-step rescue (eq. 8)
ψ	$2(1-\sqrt{1-m/m_{max}})$
ψ_0	$2(1-\sqrt{1-m_0/m_{max}})$
$ ho_{max}$	m_{max}/λ
α	$ ho_{max}\psi_0^2/4$

Table 1 Frequently used notation.

population of constant size, $m_0 = 0$ (Haldane 1927). Using this, the rate of 1-step rescue is

$$\Lambda_1(m_0) = U \int_0^{m_{max}} f(m|m_0) p_{est}(m) dm.$$
 (5)

Taking the first order approximation of $p(m_0, \Lambda_1(m_0))$ with 461 $\Lambda_1(m_0)/m_0^2$ small gives the probability of 1-step rescue given in 462 equation 5 of Anciaux et al. (2018), which effectively assumes 463 deterministic wildtype decline. For completeness we rederive 464 their closed-form approximation in File S1 (and give the results 465 in the Appendix, see Approximating the probability of 1-step

The probability of 2-step rescue is only slightly more complicated. Here $\Lambda_2(m_0)$ is the probability that a mutation arising on $_{469}$ the wildtype background creates a genotype that is also fated 470 for extinction but persists long enough for a second mutation 471 to arise on this mutant background, creating a double mutant genotype that rescues the population. We therefore have

$$\Lambda_2(m_0) = U \int_{-\infty}^{m_{max}} f(m|m_0) \left[1 - p_{est}(m) \right] p(m, \Lambda_1(m)) dm.$$
 (6)

Following this logic, we can retrieve the probability of *k*-step rescue, for arbitrary $k \ge 2$, using the recursion

$$\Lambda_{k}(m_{0}) = U \int_{-\infty}^{m_{max}} f(m|m_{0}) \left[1 - p_{est}(m)\right] p(m, \Lambda_{k-1}(m)) dm,$$
(7)

with the initial condition given by Equation 5.

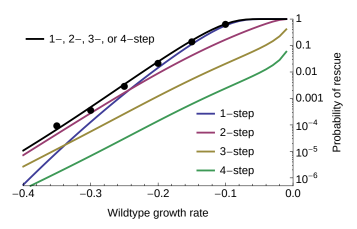


Figure 3 The probability of evolutionary rescue as a function of initial maladaptation. Shown are the probabilities of 1-, 2-, 3-, and 4-step rescue (Equations 2-7), as well as the probability of rescue by up to 4 mutational steps (using $\Lambda(m_0) = \sum_{i=1}^4 \Lambda_i(m_0)$). Dots are individual-based simulation results (ranging from 10³ to 10⁵ replicates per dot), allowing any number of mutations. Simulated populations were considered rescued when there were 1000 individuals with positive growth rates. Parameters: $N_0 = 10^4$, $U = 2 \times 10^{-3}$, n = 4, $\lambda = 0.005, m_{max} = 0.5.$

Approximating the probability of 2-step rescue

452

453

454

455

456

458

459

466

The probability of 2-step rescue is given by Equation 2 with $p_0 = p(m_0, \Lambda_2(m_0))$ (Equations 3-6). We next develop some intuition by approximating this for different classes of single

First, note that when the growth rate of a first-step mutation is close enough to zero such that $m^2 \ll \Lambda_1(m)$, we can approximate the probability such a genotype leads to rescue before itself going extinct, $p(m, \Lambda_1(m))$, with $\sqrt{2\Lambda_1(m)}$ (c.f. equation A.4b in Iwasa et al. 2004a). As shown in the Appendix (see Mutant lineage dynamics), while t < 1/|m| a mutant lineage with growth rate m that is destined for extinction persists for t generations with probability $\sim 2/t$ (Equation 21) and in generation t since it has arisen has $\sim t/2$ individuals (Equation 22). Thus, while T < 1/|m| a mutant lineage that persists for T generations will have produced a cumulative number $\sim T^2/4$ individuals. Such lineages will then lead to 2-step rescue with probability $\sim \Lambda_1(m)T^2/4$ until this approaches 1, near $T=2/\sqrt{\Lambda_1(m)}$. Since the probability of rescue increases like T^2 while the probability of persisting to time T declines only like 1/T, most rescue

435

437

438

439

440

441

443

444

445

446

events will be the result of rare long-lived single mutant genotypes. Considering only the most long-lived genotypes, the probability a first-step mutation leads to rescue is then the probability it survives long enough to almost surely rescue, $\sim 2/\sqrt{\Lambda_1(m)}$ generations. Thus, for first-step mutants with growth rates satisfying $2/\sqrt{\Lambda_1(m)} < 1/|m|$, implying $m^2 << \Lambda_1(m)$, such that they can by chance persist for unusually long times, the probability they lead to rescue is $\sim \sqrt{\Lambda_1(m)}$, consistent with our Taylor series approximation of $p(m,\Lambda_1(m))$. This same reasoning has been used to explain why the probability a neutral mutation segregates long enough to produce a second mutation is $\sim \sqrt{U}$ in a population of constant size (Weissman *et al.* 2009).

472

473

474

475

477

478

480

481

482

485

486

487

488

489

492

493

494

495

497

499

500

50

502

504

505

507

511

512

513

515

At the other extreme, when the growth rate of a first-step mutation is far enough from zero such that $m^2 >> \Lambda_1(m)$, we can approximate $p(m, \Lambda_1(m))$ with $\Lambda_1(m)/|m|$ (c.f. equation A.4c in Iwasa et al. 2004a). As just discussed, conditioned on extinction such genotypes cannot persist long enough to almost surely lead to 2-step rescue. Instead, we expect such mutations to persist for at most $\sim 1/|m|$ generations (Equation 21) with a lineage size of ~ 1 individual per generation (Equation 22), and thus produce a cumulative total of $\sim 1/|m|$ individuals. The probability of 2-step rescue from such a first-step mutation is therefore $\Lambda_1(m)/|m|$, again consistent with our Taylor series approach. This is the same reasoning that explains why a rare mutant genotype with selection coefficient |s| >> 0 in a constant population size model is expected to have a cumulative number $\sim 1/|s|$ individuals, given it eventually goes extinct (Weissman et al. 2009).

The transitions between these two regimes occur when $\Lambda_1(m)/|m|=\sqrt{2\Lambda_1(m)}$, i.e., when $|m|=\sqrt{\Lambda_1(m)/2}$. We call single mutants with growth rates $m<-\sqrt{\Lambda_1(m)/2}$ "sufficiently subcritical", those with $|m|<\sqrt{\Lambda_1(m)/2}$ "sufficiently critical", and those with $m>\sqrt{\Lambda_1(m)/2}$ "sufficiently supercritical". Given that U and thus $\Lambda_1(m)$ will generally be small, m will also be small at these transition points, meaning we can approximate them with $m^*=\sqrt{\Lambda_1(0)/2}$ and $-m^*$. We then have an approximation for the rate of 2-step rescue,

$$\begin{split} &\Lambda_{2}(m_{0}) = \Lambda_{2}^{(-)}(m_{0}) + \Lambda_{2}^{(0)}(m_{0}) + \Lambda_{2}^{(+)}(m_{0}) \\ &\Lambda_{2}^{(-)}(m_{0}) = U \int_{-\infty}^{-m^{*}} f(m|m_{0})\Lambda_{1}(m)/|m|dm \\ &\Lambda_{2}^{(0)}(m_{0}) = U \int_{-m^{*}}^{m^{*}} f(m|m_{0}) \left[1 - p_{est}(m)\right] \sqrt{2\Lambda_{1}(m)}dm \\ &\Lambda_{2}^{(+)}(m_{0}) = U \int_{m^{*}}^{m_{max}} f(m|m_{0}) \left[1 - p_{est}(m)\right] \Lambda_{1}(m)/|m|dm \end{split} \tag{8}$$

523

524

525

526

where $\Lambda_2^{(-)}(m_0)$ is the rate of 2-step rescue through sufficiently subcritical first-step mutants, $\Lambda_2^{(0)}(m_0)$ is the rate of 2-step rescue through sufficiently critical first-step mutants, and $\Lambda_2^{(+)}(m_0)$ is the rate of 2-step rescue through sufficiently supercritical first-step mutants. A schematic depicting the 1- and 2-step genetic paths to rescue is given in Figure 4.

Closed-form approximation for sufficiently critical rescue When U is small m^* is also small, allowing us to use m=0 within the integrand of $\Lambda_2^{(0)}(m_0)$, giving

$$\Lambda_2^{(0)}(m_0) \approx Uf(0|m_0)\sqrt{2\Lambda_1(0)}2m^*
= 2Uf(0|m_0)\Lambda_1(0).$$
(9)

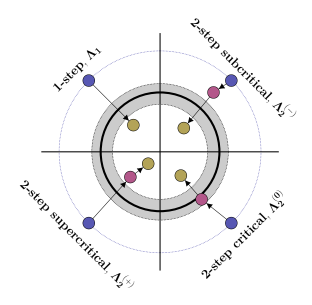


Figure 4 1- and 2-step genetic paths to evolutionary rescue. Here we show an n=2 dimensional phenotypic landscape. Continuous-time (Malthusian) growth rate (m) declines quadratically from the centre, becoming negative outside the thick black line. The grey zone indicates where growth rates are "sufficiently critical" (see text for details). Blue circles show wildtype phenotypes, red circles show intermediate first-step mutations, and yellow circles show the phenotypes of rescue genotypes.

We can then approximate $\Lambda_1(m)$ with $\tilde{\Lambda}_1(m)$ (Equation 19) and take $m \to 0$ (Equation 20), giving a closed form approximation for the rate of 2-step rescue through critical single mutants in Fisher's geometric model

$$\Lambda_2^{(0)}(m_0) \approx 4U^2 f(0|m_0) \sqrt{m_{max} \lambda / \pi}.$$
 (10)

This well approximates numerical integration of $\Lambda_2^{(0)}(m_0)$ (Equation 8; see Figure 5 and File S1) . In general, it will perform better as the range of critical growth rates, and thus $U\sqrt{m_{max}\lambda}$, becomes smaller.

To get a better understanding of how the rate of 2-step critical rescue depends on the underlying parameters of Fisher's geometric model, we approximate $f(m|m_0)$, assuming $\rho_{max} = m_{max}/\lambda$ is large, and convert this to a distribution over $\psi = 2(1-\sqrt{1-m/m_{max}})$, a convenient rescaling (for details seeFile S1 and Anciaux *et al.* 2018). Evaluating this at m=0 gives

$$\Lambda_2^{(0)}(m_0) \approx U^2 (1 - \psi_0/2)^{(1-n)/2} e^{-\alpha} \frac{2}{\pi},\tag{11}$$

where $\psi_0 = 2(1 - \sqrt{1 - m_0/m_{max}})$ and $\alpha = \rho_{max}\psi_0^2/4$.

Closed-form approximations for sufficiently non-critical rescue We can also approximate $\Lambda_1(m)$ in $\Lambda_2^{(-)}(m_0)$ and $\Lambda_2^{(+)}(m_0)$ with $\tilde{\Lambda}_1(m)$ (Equation 19), leaving us with just one integral over the growth rates of the first-step mutations. We then replace $f(m|m_0)$ with its approximate distribution over ψ .

In the case of subcritical rescue we can then make two contrasting approximations. First, when the ψ (and thus m) that contribute most are close enough to zero (meaning maladaptation is not too large relative to mutational variance) and we ignore mutations that are less fit than the wildtype, we have

$$\Lambda_2^{(-)}(m_0) \approx U^2 \frac{(1 - \psi_0/2)^{1-n}}{1 - \psi_0/4} e^{-\alpha} \frac{\log(\psi_0/\psi_-^*)}{\pi},$$
(12)

where $\psi_-^* = 2(1-\sqrt{1+\tilde{m}^*/m_{max}})$ and $\tilde{m}^* = \sqrt{\tilde{\Lambda}_1(0)/2}$. Second, when the mutational variance, λ , is very small relative to maladaptation, implying that mutants far from m=0 substantially contribute, we find

544

545

546

548

549

550

551

553

554

555

556

557

560

561

565

566

567

568

570

571

572

573

574

575

578

579

580

581

584

585

586

587

588

589

$$\Lambda_2^{(-)}(m_0) \approx -U^2 \frac{(1 - \psi_0/2)^{1-n}}{1 - \psi_0/4} \left(e^{-\alpha} \frac{1}{(\alpha/2)^3 \pi} \right)^{1/2}.$$
 (13)

These two approximations do well compared with $\Lambda_2^{(-)}(m_0)$ (Equation 8; see Figure 5 and File S1). As expected, we find that Equation 13 does better under fast wildtype decline while Equation 12 does better when the wildtype is declining more slowly.

For supercritical 2-step rescue, only first-step mutants with growth rates near m^* will contribute (larger m will rescue themselves and are also less likely to arise by mutation), and so we can capture the entire distribution with a small m approximation (following the same approach that led to Equation 12). As shown in File S1, this approximation works well for $\psi < \sqrt{2/\rho_{max}}$, beyond which the rate of 2-step rescue through such first-step mutants falls off very quickly due to a lack of mutational input. Thus, considering only supercritical single mutants with scaled growth rate less than $\sqrt{2/\rho_{max}}$, our approximation is

$$\Lambda_2^{(+)}(m_0) \approx U^2 \frac{(1 - \psi_0/2)^{1-n}}{1 - \psi_0/4} e^{-\alpha} \frac{\log(\psi_{max}/\psi_+^*)}{\pi},$$
(14)

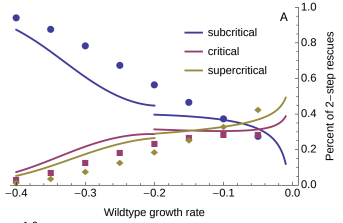
with $\psi_+^*=2(1-\sqrt{1-\tilde{m}^*/m_{max}})$ and $\psi_{max}=\sqrt{2/\rho_{max}}$. This approximation tends to provide a slight overestimate of $\Lambda_2^{(+)}(m_0)$ (Equation 8; see Figure 5 and File S1).

Comparing 2-step regimes These rough but simple closed-form approximations (Equations 11–14) show that, while the contribution of critical mutants to 2-step rescue scales with U^2 , the contribution of non-critical single mutants scales at a rate less than U^2 due to an increase in the range of critical single mutants (both ψ_{-}^{*} and ψ_{+}^{*} increase with U). This difference in scaling with *U* is stronger when the wildtype is not very maladapted relative to the mutational variance, i.e., when Equation 12 is the better approximation for subcritical rescue. The approximations 592 also show that, when initial maladaptation is small the ratio of 593 supercritical to subcritical contributions (Equation 12 divided by 14) primarily depends only on the range of growth rates in- 595 cluded in each regime, while with larger initial maladaptation this ratio (Equation 13 divided by 14) begins to depend more strongly on initial maladaptation and mutational variance (α). The effect of maladaptation and mutation rate on the relative contributions of each regime is shown in Figure 5.

The distribution of growth rates among rescue genotypes

We next explore the distribution of growth rates among rescue genotypes, i.e., the distribution of growth rates that we expect to observe among the survivors across many replicates.

We begin with 1-step rescue. The rate of 1-step rescue by genotypes with growth rate m is simply $Uf(m|m_0)p_{est}(m)$. Disolating this by the rate of 1-step rescue through any m (Equation 604 5) gives the distribution of growth rates among the survivors 605



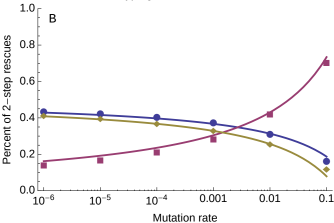


Figure 5 The relative contribution of sufficiently subcritical, sufficiently critical, and sufficiently supercritical single mutants to 2-step rescue. The curves are drawn using Equations 10–14 (Equation 12 is used for $m_0 < 0.2$ while Equation 13 is used for $m_0 > 0.2$). The dots are numerical evaluations of Equation 8. Parameters: n = 4, $\lambda = 0.005$, $m_{max} = 0.5$, (A) $U = 10^{-3}$, (B) $m_0 = -0.1$.

$$g_1(m) = \frac{Uf(m|m_0)p_{est}(m)}{\Lambda_1(m_0)},$$
(15)

where U cancels out. This distribution is shown in blue in Figure 6. The distribution has a mode at small but positive m as a result of two conflicting processes: smaller growth rates are more likely to arise from a declining wildtype but larger growth rates are more likely to establish given they arise. As the rate of wildtype decline increases, the former process exerts more influence, causing the mode to move towards zero and reducing the variance.

We can also give a closed form approximation here using the same approach taken to reach Equation 19. On the ψ scale we have

$$\tilde{g}_{1}(\psi) = \frac{\exp(\alpha)\sqrt{\alpha\rho_{max}}}{[\exp(\alpha)\sqrt{\pi\alpha}\mathrm{Erfc}(\sqrt{\alpha}) - 1]\psi_{0}} e^{-\rho_{max}(\psi - \psi_{0})^{2}/4}\psi, \quad (16)$$

implying the ψ are distributed like a normal truncated below $\psi = 0$ and weighted by ψ . This provides a very good approximation (see dashed blue curves in Figure 6).

In 2-step rescue, the rate of rescue by double mutants with growth rate m_2 is given by Equation 6 with $\Lambda_1(m)$ replaced

by $Uf(m_2|m)p_{est}(m_2)$. Normalizing gives the distribution of growth rates among the double mutant genotypes that rescue the population

$$g_{2}(m_{2}) \approx \frac{A(m_{2})}{\int_{0}^{m_{max}} A(m_{2}) dm_{2}}$$

$$A(m_{2}) = \int_{-\infty}^{m_{max}} f(m|m_{0}) [1 - p_{est}(m)]$$

$$p(m, Uf(m_{2}|m) p_{est}(m_{2})) dm.$$
(17)

This distribution, $g_2(m)$, is shown in red in Figure 6. Because the first-step mutants contributing to 2-step rescue tend to be nearer the optimum than the wildtype, this allows them to produce double mutant rescue genotypes with higher growth rates than in 1-step rescue (as seen by comparing the mode between blue and red curves in Figure 6). The fact that these first-step mutants are closer to the optimum also allows for a greater variance in the growth rates of rescue genotypes than in 1-step rescue. This also allows the 2-step distribution to maintain a more similar mode and variance across wildtype decline rate than the 1-step distribution. Note that because the *U* in $Uf(m_2|m)p_{est}(m_2)$ does not cancel, $g_2(m_2)$ depends on U and the buffering effect of first-step mutants depends on the mutation rate. In principle, decreasing the mutation rate disproportionately increases the contribution of first-step mutants with growth rates closer to zero, as these genotypes are more likely to persist long enough to produce rescue mutants (see The distribution of growth rates among rescue intermediates below for more discussion). This implies that the distribution of rescue genotype growth rates is more strongly buffered from the rate of wildtype decline at lower mutation rates.

610

614

615

616

617

620

621

622

623

624

625

629

63

635

636

637

638

639

642

643

644

645

650

651

653

The distribution of growth rates among rescue intermediates

Finally, our analyses above readily allow us to explore the distribution of first-step mutant growth rates that contribute to 2-step rescue. Analogously to Equation 15, we drop the integral in $\Lambda_2(m_0)$ (Equation 6) and normalize, giving

$$h(m) = \frac{Uf(m|m_0) [1 - p_{est}(m)] p(m, \Lambda_1(m))}{\Lambda_2(m_0)}, \quad (18)$$

where the first *U* cancels but the *U* within $\Lambda_1(m)$ does not. This distribution is shown in black in Figure 7. At slow wildtype decline rates the overwhelming majority of 2-step rescue events arise from first-step mutants with growth rates near 0. As indicated by Equation 8, the contribution of first-step mutants with growth rate *m* declines as $\sim 1/|m|$ as *m* departs from zero, due to shorter persistence times given eventual extinction. As wildtype growth rate declines the relative importance of mutational input, $f(m|m_0)$, grows, causing the distribution to flatten and first-step mutants with substantially negative growth rates begin to contribute (see also Figure 5A). Decreasing the mutation rate disproportionately increases the contribution of firststep mutants with growth rates near zero (while simultaneously shrinking the range of growth rates that are sufficiently critical; Figure 5B) making the distribution of first-step mutant growth rates contributing to 2-step rescue more sharply peaked around m = 0 (Figure S2). Correspondingly, with a higher mutation rate a greater proportion of the contributing single mutants have substantially negative growth rates.

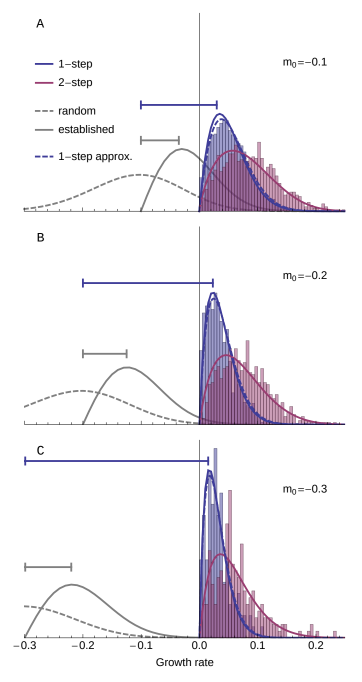


Figure 6 The distribution of growth rates among rescue genotypes under 1-step (blue; Equation 15 solid and 16 dashed) and 2-step (red; Equation 17) rescue for three different levels of initial maladaptation. For comparison, the distribution of random mutations (dashed; Equation 1) and the distribution of beneficial mutations that establish in a population of constant size (solid grey; Equation 1 times Equation 4 and normalized) are shown. Horizontal lines indicate the most common fitness effect ($s = m_0 - m$) in a population of constant size (grey) and in 1-step rescue (blue). The histograms show the distribution of growth rates among rescue genotypes observed across (A) 10^4 , (B) 10^5 , and (C) 10^6 simulated replicates. Populations were considered rescued once there were \geq 100 individuals with positive growth rate. The most common genotype at this point was considered the rescue genotype, and the number of mutational steps to rescue was set as the number of mutations in that genotype. Other parameters: $N_0 = 10^4$, $U = 2 \times 10^{-3}$, n = 4, $\lambda = 0.005$, $m_{max} = 0.5$.

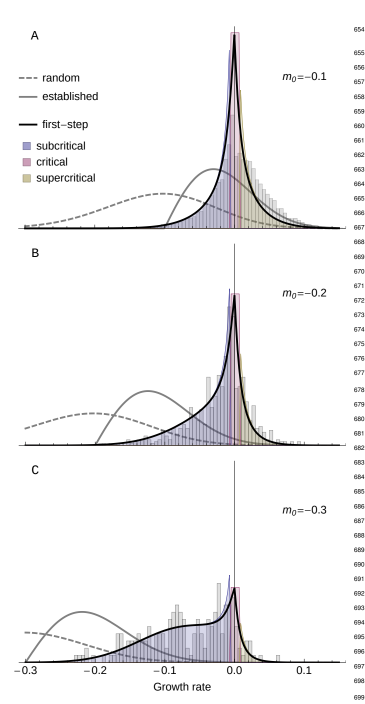


Figure 7 The distribution of growth rates among first-step mutations that lead to 2-step rescue (black; Equation 18) for three different levels of initial maladaptation. Shading represents our sufficiently subcritical approximation (blue; replacing $p(m, \Lambda_1(m))$ with $\Lambda_1(m)/|m|$ in the numerator of Equation 18), our sufficiently critical approximation (red; using $Uf(0|m_0)\sqrt{2\Lambda_1(0)}$ as the numerator in Equation 18), and our sufficiently supercritical approximation (yellow; replacing $p(m, \Lambda_1(m))$ with $\Lambda_1(m)/|m|$ in the numerator of Equation 18). The histograms show the distribution of growth rates among first-step mutations in rescue genotypes with 2 mutations observed across (\mathbf{A}, \mathbf{B}) 10^5 or (\mathbf{C}) 10^6 simulated replicates. Note that the discrepancy in panel \mathbf{A} is likely due to us sampling only the most common rescue genotype in each replicate. See Figure 6 for additional details.

Discussion

700

701

702

703

704

705

706

707

712

713

714

Here we've explored the possibility of evolutionary rescue by multiple mutations on a simple fitness landscape. We find that rescue by multiple mutations can be the most likely path to persistence under high mutation rates or when the population is initially very maladapted. Under these scenarios, intermediate genotypes that are declining less quickly provide a 'springboard' from which rescue genotypes emerge. In 2-step rescue these springboard single mutants come from one of three regimes: those that have growth rates near enough to zero (i.e., replacement) to, by chance, persist for unusually long periods of time and therefore reach unusually large subpopulation sizes and those with growth rates either too negative or too positive to do so. The relative contribution of each regime shifts with initial maladaptation and mutation rate; rare mutations that can occasionally reach unusually large subpopulation sizes play a larger role as maladaptation declines and mutation increases. In contrast, when initial maladaptation is very high most first-step mutations are themselves also very maladapted and thus restricted in the subpopulation sizes they reach. All three regimes help maintain the variance in the distribution of fitness effects among rescue genotypes as initial maladaptation increases; meanwhile, in 1-step rescue the variance declines due to ever more extreme sampling of the tail of the mutational distribution.

How often rescue arises as a result of multiple mutations is an open question. It is clear that more than one mutation can contribute to adaptation to near-lethal stress, but experiments are often designed to avoid extinction (reviewed in Cowen et al. 2002), and therefore greatly expand the scope for multiple mutations to arise on a single genotype. A few exceptions provide some insight. For example, populations of Saccharomyces cervisae that survived high concentrations of copper acquired multiple mutations (Gerstein et al. 2015) - in fact the authors argue for the 'springboard effect' discussed above, where first step mutations prolong persistence and thereby allow further mutations to arise. In Pseudomonas flourescens, fluctuation tests with nalidixic acid showed that nearly a third of the most resistant surviving strains were double mutants (Bataillon et al. 2011), which were able to tolerate 10x higher drug concentrations than single mutants, suggesting 2-step rescue might dominate at high drug concentrations. It is unclear if our prediction – that rescue takes more mutational steps with greater initial maladaptation – holds true. Verification will require more experiments that allow extinction and uncover the genetic basis of adaptation at different severities of environmental change (e.g., drug concentration).

In describing the genetic basis of adaptation in populations of constant size, Orr (1998) showed that the mean phenotypic displacement towards the optimum scales roughly linearly with initial displacement. Converting phenotype to fitness, this implies the mean fitness effect of fixed mutations, $s = m - m_0$, increases as $\sim \exp(-m_0)$ as initial Malthusian fitness, m_0 , declines, which is roughly linear when m_0 is small. Here we see that, under 1-step rescue, the mean fitness effect also increases roughly linearly as the initial growth rate declines (see horizontal blue lines in Figure 6). However, the rate of this linear increase in fitness effect is much larger under rescue than in a population of constant size (compare blue and grey horizontal lines in Figure 6), where declines in wildtype fitness not only allow larger mutations to be beneficial but also require larger mutations for persistence. Thus the race between extinction and adaptation during evolutionary rescue is expected to produce a genetic basis of adaptation with fewer mutations of larger effect.

While under 1-step rescue the fitness effect of the first mutation increases roughly linearly as wildtype fitness declines, 779 below some wildtype fitness most rescue events will be 2-step 780 (e.g., at $m_0 \approx -0.25$ in Figure 3). At this junction the effect size 781 of the first mutation will no longer increase as quickly (and po-782 tentially even decrease), as its expected fitness will itself begin 783 to decline substantially with the fitness of the wildtype (Figures 5 and 7). Thus as rescue switches from dominantly *k*-step to dominantly (k + 1)-step the genetic basis of adaptation becomes more diffuse, with each mutation having a smaller individual fitness effect as the contributing fitness effects spread over more loci. In the limit of large *k* (due to large initial maladaptation or high mutation rates), the genetic basis of adaptation should at some point converge to many loci with small effect, as would also be expected in a population of constant size. Indeed, at very high mutation rates the rate of adaptation (the change in mean fitness) is the same under rescue as it is in populations of constant size (Anciaux et al. 2019), implying that the genetic 795 basis of adaptation no longer depends on demography. It is therefore at intermediate levels of initial maladaptation and low 797 mutation rates, where rescue primarily occurs from a few large effect mutations, that the race between adaptation and persistence is predicted to have the largest effect on the genetic basis of adaptation.

716

717

718

719

720

72

722

72

728

720

730

73

734

735

736

737

742

743

744

745

746

747

750

75

752

758

759

760

761

765

766

767

772

773

774

775

776

Fluctuation tests (?) provide a means to isolate rescue genotypes, whose growth rates can then be measured under the selective conditions. Consistent with our theory (Figure 6), the resulting growth rate distributions in both bacteria and yeast often find modes that are substantially greater than zero (as opposed to, say, an exponential distribution; Kassen and Bataillon 2006; MacLean and Buckling 2009; Gerstein et al. 2012; Lindsey et al. 2013; Gerstein et al. 2015). A number of these conform even more closely to our expected shape (Kassen and Bataillon 2006; 810 Gerstein et al. 2015) while the others appear to be substantially 811 more clumped around the mode, perhaps due to a very restricted number of possible rescue mutations in any one circumstance, 813 the size of the experiment, or the way in which growth rates are 814 measured. Further, these experiments are designed to begin with 815 substantial standing genetic variation (at least across replicates), which should increase the contributions of mutations with small growth rates (?), although these could be outcompeted by mutations with higher growth rates and/or be undersampled. The role of standing genetic variance becomes particularly important when we consider rescue by multiple steps, as the ability of the 821 intermediate genotypes to persist long enough to accumulate 822 further mutations will differ strongly between environments. 823 Finally, Gerstein et al. (2015) not only provides the distribution 824 of growth rates among rescue genotypes, but also the growth 825 rates of individual mutations that compose multi-step rescue 826 genotypes. In two of three cases they see that most component 827 mutations have growth rates near that of the multi-mutation 828 genotype while in the other case there appears to be multiple 829 mutations that provide intermediate benefits, consistent with their hypothesis of multi-step rescue. These growth rates were measured in a less stressful environment than the original selective environment, which unfortunately makes it impossible to tell if these individual mutations had positive or negative growth rates (i.e., were subcritical or supercritical) during rescue.

Pinpointing the mutations responsible for adaptation is hampered by genetic hitchhiking, as beneficial alleles elevate the frequency of linked neutral and mildly deleterious alleles (Barton 2000). The problem is particularly severe under strong selection 839

and low recombination, and therefore reaches an extreme in the case of evolutionary rescue in asexuals, especially if many neutral and deleterious mutations are segregating at the time of environmental change. To circumvent this, mutations that have risen to high frequency in multiple replicates are often introduced in a wildtype background, in isolation and sometimes also in combination with a small number of other common high-frequency mutations, and grown under the selective conditions (e.g., Jochumsen et al. 2016; Ono et al. 2017). As we have demonstrated above, however, under multi-step rescue there may be no one mutation that individually confers growth in the selective conditions. Thus, a mutation driving rescue may go undetected or be mistaken as a hitchhiker if the appropriate multiple-mutation genotypes are not tested. Unfortunately reverse engineering all combinations of mutations quickly becomes unwieldy as the number of mutations grows, and thus this approach will not be practical under severe initial maladaptation and high mutation rates, where we predict rescue to occur by many mutations. Interestingly, our simulations show that the population dynamics themselves may help differentiate how many mutations contribute to rescue (e.g., V- vs. U-shaped logtrajectories; Figures 1 and 2), and fitting models of k-step rescue could produce estimates for the growth rates of the *k* genotypes.

Environmental change often selects for mutator alleles, which elevate the rate at which beneficial alleles arise and subsequently increase in frequency with them (Tenaillon et al. 2001). When beneficial alleles are required for persistence, as in evolutionary rescue, mutator alleles can reach very high frequencies or rapidly fix (e.g., Mao et al. 1997). Consistent with this, mutator alleles are often associated with antibiotic resistance in clinical isolates (see examples in Bell 2017). Further, the more beneficial mutations available the larger the advantage of a mutator allele; for a mutator that increases the mutation rate *m*-fold, its relative contribution to the production of *n* beneficial mutations scales as m^n (Tenaillon et al. 1999). Thus multi-step rescue can impose much stronger selection for mutator alleles than 1-step rescue. There are a number of examples where lineages with higher mutation rates acquired multiple mutations and persisted at higher doses of antibiotics (Couce et al. 2015; San Millan et al. 2017). The number of mutations required for persistence is, however, often unknown, making it difficult to compare situations where rescue requires different numbers of mutations. Experiments with a combination of drugs may provide a glimpse; for instance, Escherichia coli populations only evolved resistance to a combination of two drugs (presumably through the well-known mutations specific to each drug) when mutators were present, despite the fact that mutators were not required for resistance to either drug in isolation (Gifford et al. 2019). In cases where we have less information on the genetic basis of resistance, our model suggests that mutators will be more advantageous when initial maladaptation is severe (e.g., higher drug concentrations or a larger number of drugs), as rescue will then be dominated by genetic paths with more mutational steps.

Here we have investigated the genetic basis of evolutionary rescue in an asexual population that is initially genetically uniform. Extending this work to allow for recombination and standing genetic variation at the time of environmental change will likely be interesting. The effect of standing genetic variation on the probability of 1-step rescue is relatively straight-forward to incorporate, depending only on the expected number of rescue mutations initially present and their mean establishment probability (Martin *et al.* 2013). In the case of mutation-selection

balance, the probability of 1-step rescue from standing genetic 900 variance in Fisher's geometric model was given by Anciaux et al. 901 (2018), whose equations 3 and 5 immediately give the distribu- 902 tion of fitness effects among those that rescue. Allowing these 903 standing genetic variants to be springboards to multi-step res- 904 cue will help clarify the role of standing genetic variation on the 905 genetic basis of rescue more generally. Recombination can help 906 combine such springboard mutations into rescue genotypes but will also break these combinations apart, as demonstrated in a 2-locus 2-allele model of rescue (Uecker and Hermisson 2016). How recombination affects the genetic basis of evolutionary rescue when more loci can potentially contribute remains to be seen.

Acknowledgements

842

843

845

851

852

853

854

855

858

859

860

867

868

869

872

873

874

875

876

879

881

882

883

884

887

888

889

895

896

897

898

We would like to thank the Otto and Doebeli labs for helpful feedback at various stages, Ophélie Ronce and Thomas Lenormand for their hospitality and valuable input at the beginning of this project, and Mike Whitlock, Amy Angert, and Luis-Miguel Chevin for constructive criticism on previous versions of the manuscript. Funding provided by the National Science and Engineering Research Council (CGS-D 6564 to M.M.O., RGPIN-2016-03711 to S.P.O.), the University of British Columbia, Banting, and the University of California - Davis (fellowships to M.M.O.), the National Institute of General Medical Sciences of the National Institutes of Health (NIH R01 GM108779 to Graham Coop), the Agence Nationale de la Recherche (ANR-18-CE45-0019 "RESISTE" to G.M.), and the Centre Méditerranéen Environment et Biodiversité ("BACTPHI" to G.M.).

Literature Cited

- Abramowitz, M. and I. A. Stegun, editors, 1972 Handbook of mathematical functions with formulas, graphs, and mathematical tables. United States Department of Commerce, Washington,
- Alexander, H. K. and T. Day, 2010 Risk factors for the evolutionary emergence of pathogens. Journal of the Royal Society Interface 7: 1455-1474.
- Alexander, H. K., G. Martin, O. Y. Martin, and S. Bonhoeffer, 939 2014 Evolutionary rescue: linking theory for conservation and medicine. Evolutionary Applications 7: 1161–1179.
- Allen, L. J., 2010 An introduction to stochastic processes with applications to biology. CRC Press.
- Anciaux, Y., L.-M. Chevin, O. Ronce, and G. Martin, 2018 Evolutionary rescue over a fitness landscape. Genetics 209: 265–279.
- Anciaux, Y., A. Lambert, O. Ronce, L. Roques, and G. Martin, 2019 Population persistence under high mutation rate: from evolutionary rescue to lethal mutagenesis. BioRxiv p. 521203.
- Antia, R., R. R. Regoes, J. C. Koella, and C. T. Bergstrom, 2003 The role of evolution in the emergence of infectious diseases. Nature 426: 658.
- Barton, N. H., 2000 Genetic hitchhiking. Philosophical Transactions of the Royal Society of London. Series B: Biological Sciences 355: 1553-1562.
- Bataillon, T. and S. F. Bailey, 2014 Effects of new mutations on fitness: Insights from models and data. Annals of the New York Academy of Sciences 1320: 76–92
- Bataillon, T., T. Zhang, and R. Kassen, 2011 Cost of adaptation 958 and fitness effects of beneficial mutations in pseudomonas 959 fluorescens. Genetics 189: 939-949.
- Bell, G., 2009 The oligogenic view of adaptation. Cold Spring Harbor Symposia on Quantitative Biology 74: 139–144.

- Bell, G., 2017 Evolutionary rescue. Annual Review of Ecology, Evolution, and Systematics 48: 605–627.
- Couce, A., A. Rodríguez-Rojas, and J. Blázquez, 2015 Bypass of genetic constraints during mutator evolution to antibiotic resistance. Proceedings of the Royal Society B: Biological Sciences 282: 20142698.
- Cowen, L. E., J. B. Anderson, and L. M. Kohn, 2002 Evolution of drug resistance in candida albicans. Annual Reviews in Microbiology **56**: 139–165.
- Dettman, J. R., N. Rodrigue, A. H. Melnyk, A. Wong, S. F. Bailey, et al., 2012 Evolutionary insight from whole-genome sequencing of experimentally evolved microbes. Molecular ecology 21: 2058-2077.

912

913

914

929

930

931

935

936

937

942

943

950

953

957

960

- Fisher, R. A., 1918 The correlation between relatives on the supposition of mendelian inheritance. Transactions of the Royal Society of Edinburgh 52: 399–433.
- Fisher, R. A., 1930 The genetical theory of natural selection. Clarendon Press, London.
- Foll, M., Y.-P. Poh, N. Renzette, A. Ferrer-Admetlla, C. Bank, et al., 2014 Influenza virus drug resistance: a time-sampled population genetics perspective. PLoS Genetics 10: e1004185.
- Gerstein, A. C., D. S. Lo, and S. P. Otto, 2012 Parallel genetic changes and nonparallel gene-environment interactions characterize the evolution of drug resistance in yeast. Genetics **192**: 241-252.
- Gerstein, A. C., J. Ono, D. S. Lo, M. L. Campbell, A. Kuzmin, et al., 2015 Too much of a good thing: the unique and repeated paths toward copper adaptation. Genetics **199**: 555–571.
- Gifford, D. R., E. Berríos-Caro, C. Joerres, T. Galla, and C. G. Knight, 2019 Mutators drive evolution of multi-resistance to antibiotics. bioRxiv p. 643585.
- Gomulkiewicz, R. and R. D. Holt, 1995 When does evolution by natural selection prevent extinction? Evolution 49: 201–207.
- Haldane, J. B. S., 1927 A Mathematical Theory of Natural and Artificial Selection, Part V: Selection and Mutation. Mathematical Proceedings of the Cambridge Philosophical Society 23: 838.
- Harmand, N., R. Gallet, R. Jabbour-Zahab, G. Martin, and T. Lenormand, 2017 Fisher's geometrical model and the mutational patterns of antibiotic resistance across dose gradients. Evolution 71: 23–37.
- Iwasa, Y., F. Michor, and M. A. Nowak, 2004a Evolutionary dynamics of invasion and escape. Journal of Theoretical Biology **226**: 205-214.
- Iwasa, Y., F. Michor, and M. A. Nowak, 2004b Stochastic tunnels in evolutionary dynamics. Genetics 166: 1571–1579.
- Jochumsen, N., R. L. Marvig, S. Damkiær, R. L. Jensen, W. Paulander, et al., 2016 The evolution of antimicrobial peptide resistance in pseudomonas aeruginosa is shaped by strong epistatic interactions. Nature communications 7: 13002.
- Kassen, R. and T. Bataillon, 2006 Distribution of fitness effects among beneficial mutations before selection in experimental populations of bacteria. Nature genetics 38: 484.
- Kimura, M., 1965 A stochastic model concerning the maintenance of genetic variability in quantitative characters. Proceedings of the National Academy of Sciences 54: 731–736.
- Kimura, M., 1983 The neutral theory of molecular evolution. Cambridge University Press, Cambridge, UK.
- Lande, R., 1980 The genetic covariance between characters maintained by pleiotropic mutations. Genetics **94**: 203–215.
- Lindsey, H. A., J. Gallie, S. Taylor, and B. Kerr, 2013 Evolutionary rescue from extinction is contingent on a lower rate of

environmental change. Nature 494: 463–467.

962

963

964

965

966

967

972

973

974

975

976

977

979

980

981

982

990

99

992

995

996

997

1002

1003

1004

1005

1006

100

1008

1009

1010

1011

1012

1013

1014

1017

1018

1019

1020

1021

1022

- MacLean, R. C. and A. Buckling, 2009 The distribution of fitness 1025 effects of beneficial mutations in *Pseudomonas aeruginosa*. PLoS 1026 Genetics 5: e1000406.
- MacLean, R. C., A. R. Hall, G. G. Perron, and A. Buckling, 2010 1028 The population genetics of antibiotic resistance: integrating 1029 molecular mechanisms and treatment contexts. Nature Re- 1030 views Genetics 11: 405.
- Mao, E. F., L. Lane, J. Lee, and J. H. Miller, 1997 Proliferation 1032 of mutators in a cell population. Journal of Bacteriology **179**: 1033 417–422.
- Martin, G., R. Aguilee, J. Ramsayer, O. Kaltz, and O. Ronce, 2013 1035 The probability of evolutionary rescue: towards a quantita- 1036 tive comparison between theory and evolution experiments. 1037 Philosophical Transactions of the Royal Society of London B: 1038 Biological Sciences 368: 20120088.
- Martin, G. and T. Lenormand, 2006 The fitness effect of muta- 1040 tions across environments: a survey in light of fitness land- 1041 scape models. Evolution **60**: 2413–2427.
- Martin, G. and T. Lenormand, 2015 The fitness effect of muta-1043 tions across environments: Fisher's geometrical model with 1044 multiple optima. Evolution **69**: 1433–1447.
- Martin, G. and L. Roques, 2016 The nonstationary dynamics of 1046 fitness distributions: asexual model with epistasis and stand-1047 ing variation. Genetics **204**: 1541–1558.
- Maruyama, T. and M. Kimura, 1974 A note on the speed of gene 1049 frequency changes in reverse directions in a finite population. 1050 Evolution pp. 161–163.
- Ono, J., A. C. Gerstein, and S. P. Otto, 2017 Widespread genetic 1052 incompatibilities between first-step mutations during parallel 1053 adaptation of saccharomyces cerevisiae to a common environ- 1054 ment. PLoS biology **15**: e1002591.
- Orr, H. A., 1998 The Population Genetics of Adaptation: The Distribution of Factors Fixed during Adaptive Evolution. Evolution **52**: 935.
- Orr, H. A., 2005 The genetic theory of adaptation: a brief history. Nature Reviews Genetics **6**: 119–27.
- Orr, H. A. and R. L. Unckless, 2014 The population genetics of evolutionary rescue. PLoS Genetics **10**: e1004551.
- Otto, S. P. and M. C. Whitlock, 1997 The probability of fixation in populations of changing size. Genetics **146**: 723–733.
- Pennings, P. S., S. Kryazhimskiy, and J. Wakeley, 2014 Loss and recovery of genetic diversity in adapting populations of hiv. PLoS Genetics **10**: e1004000.
- Robbins, N., T. Caplan, and L. E. Cowen, 2017 Molecular evolution of antifungal drug resistance. Annual Review of Microbiology 71: 753–775.
- San Millan, A., J. A. Escudero, D. R. Gifford, D. Mazel, and R. C. 1064 MacLean, 2017 Multicopy plasmids potentiate the evolution 1065 of antibiotic resistance in bacteria. Nature ecology & evolution 1066 1: 0010.
- Schlötterer, C., R. Kofler, E. Versace, R. Tobler, and S. Franssen, 2015 Combining experimental evolution with next-generation sequencing: a powerful tool to study adaptation from standing genetic variation. Heredity 114: 431–440.
- Stapley, J., J. Reger, P. G. Feulner, C. Smadja, J. Galindo, *et al.*, 1068 2010 Adaptation genomics: the next generation. Trends in 1069 ecology & evolution **25**: 705–712.
- Tenaillon, O., 2014 The utility of Fisher's geometric model in 1071 evolutionary genetics. Annual Review of Ecology, Evolution, 1072 and Systematics 45: 179–201.
- Tenaillon, O., F. Taddei, M. Radman, and I. Matic, 2001 Second-1074

- order selection in bacterial evolution: selection acting on mutation and recombination rates in the course of adaptation. Research in microbiology **152**: 11–16.
- Tenaillon, O., B. Toupance, H. Le Nagard, F. Taddei, and B. Godelle, 1999 Mutators, population size, adaptive land-scape and the adaptation of asexual populations of bacteria. Genetics **152**: 485–493.
- Turelli, M., 1984 Heritable genetic variation via mutationselection balance: Lerch's zeta meets the abdominal bristle. Theoretical population biology **25**: 138–193.
- Turelli, M., 1985 Effects of pleiotropy on predictions concerning mutation-selection balance for polygenic traits. Genetics 111: 165–195.
- Uecker, H. and J. Hermisson, 2016 The role of recombination in evolutionary rescue. Genetics **202**: 721–732.
- Uecker, H., S. P. Otto, and J. Hermisson, 2014 Evolutionary rescue in structured populations. The American Naturalist 183: E17– E35.
- Weinreich, D. M., N. F. Delaney, M. A. DePristo, and D. L. Hartl, 2006 Darwinian evolution can follow only very few mutational paths to fitter proteins. science **312**: 111–114.
- Weissman, D. B., M. M. Desai, D. S. Fisher, and M. Feldman, 2009 The rate at which asexual populations cross fitness valleys. Theoretical Population Biology 75: 286–300.
- Weissman, D. B., M. W. Feldman, and D. S. Fisher, 2010 The rate of fitness-valley crossing in sexual populations. Genetics **186**: 1389–1410.
- Wolfram Research Inc., 2012 Mathematica, Version 9.0. Champaign, IL.
- Yilmaz, N. K., R. Swanstrom, and C. A. Schiffer, 2016 Improving viral protease inhibitors to counter drug resistance. Trends in Microbiology **24**: 547–557.

Appendix

1024

Approximating the probability of 1-step rescue

The probability of 1-step rescue in this model has been derived by Anciaux *et al.* (2018). As replicated in File S1 and given by their equation 7, when $\rho_{max} = m_{max}/\lambda$ is large a closed-form approximation is

$$\Lambda_1(m_0) \approx \tilde{\Lambda}_1(m_0) \equiv -m_0 U \frac{(1 - \psi_0/2)^{(1-n)/2}}{1 - \psi_0/4} g(\alpha), \quad (19)$$

where $\psi_0=2(1-\sqrt{1-m_0/m_{max}}), g(\alpha)=\exp(-\alpha)/\sqrt{\pi\alpha}-\exp(-\sqrt{\alpha}),$ and $\alpha=\rho_{max}\psi_0^2/4$, with erfc(.) the complimentary error function. When the wildtype declines slowly m_0 and thus ψ_0 is small and $\Lambda_1(m_0)\approx Ug(\alpha)$. In the limit $m_0\to 0$, Equation 19 becomes

$$\tilde{\Lambda}_1(0) \equiv \lim_{m_0 \to 0} \tilde{\Lambda}_1(m_0) = 2U\sqrt{m_{max}\lambda/\pi}.$$
 (20)

Mutant lineage dynamics

Here we follow the lead of Weissman *et al.* (2010) and Uecker and Hermisson (2016) in approximating our discrete-time process with a continuous-time branching process (see chapter 6 in Allen 2010). Consider a birth-death process, where individuals give birth at rate b and die at rate d. One can then obtain the probability generating function for the number of individuals at a given time, n(t), given the initial number, n(0). We are primarily interested in new mutant lineages, n(0) = 1. The generating

function then allows us to calculate the probability that a lineage persists at least until time t and the distribution of n(t) given it does so (see below).

To convert between birth and death rates and our compound Malthusian parameter we follow Uecker and Hermisson (2016) in equally distributing the growth rate m between birth and death, b = (1+m)/2 and d = (1-m)/2, such that m = b - d and the continuous-time process exhibits the same amount of drift as the discrete time process (and matches discrete-time simulations well; Uecker *et al.* 2014). We can now report the necessary results in terms of m (assuming |m| < 1).

Denoting the extinction time as T, the probability a mutant with growth rate m persists until time t is approximately (see File S1 for derivation)

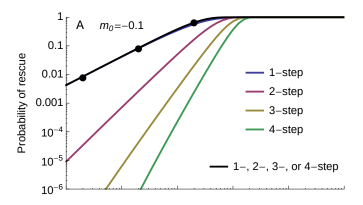
$$P(T > t) \approx \begin{cases} 2/t & t << |1/m| \\ -2m \exp(mt) & t >> -1/m > 0 \end{cases}$$
 (21)

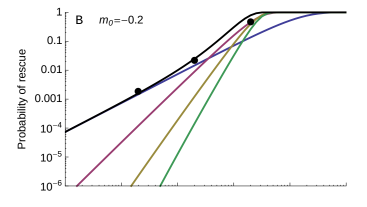
As pointed out in Weissman *et al.* (2010) (whose equation A2 differs from Equation 21 by a factor of 2 because they have b+d=2), the distribution of persistence times has a long tail (like 1/t) until being cut off (declining exponentially) at t=-1/m.

Given a lineage persists until t, the distribution of n(t) is roughly (see File S1 for derivation)

$$P(n(t) = n|n(t) > 0) \approx \begin{cases} 2(1/t)(1 + 2/t)^{-n} & t << |1/m| \\ -2m(1+m)^{n-1} & t >> -1/m > 0 \end{cases}$$
(22)

As pointed out in Weissman *et al.* (2010) (whose equation A3 only differs from Equation 22 by constants), the distribution of n(t) is approximately geometric for small or large t, implying n(t) is very unlikely to be greater than the minimum of t and -1/m.





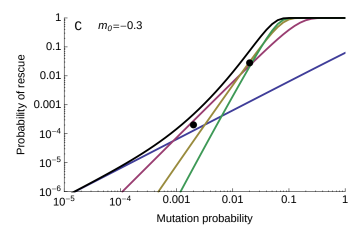


Figure S1 The probability of rescue as a function of mutation rate for three different levels of initial maladaptation. See Figure 3 for details. Other parameters: n=4, $\lambda=0.005$, $m_{max}=0.5$, $N_0=10^4$.

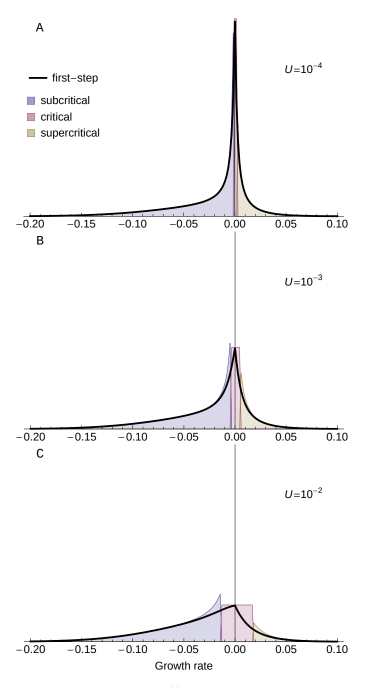


Figure S2 The distribution of first-step mutant growth rates given 2-step rescue under three mutation rates. See Figure 7 for details. Parameters: n=4, $\lambda=0.005$, $m_{max}=0.5$, $m_0=-0.2$.