

Why there are so many definitions of fitness in models

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Data availability

No new data were used in this study. Code is available at:
https://github.com/DanielSmithEcology/Fitness_Definitions_Code

Conflicts of interest

We declare no conflicts of interest.

Abstract

“Fitness” quantifies the ability to survive and reproduce, but is operationalized in many different ways. Generally, short-term fitness (e.g., expected number of surviving offspring) is *assigned* to genotypes or phenotypes, and used to non-trivially *derive* longer-term operationalizations of fitness (e.g. fixation probability or sojourn time), providing insight as to which organismal strategies tend to evolve due to natural selection. Assigned fitness operationalizations vary, but all summarize currently expected organismal vital rates (i.e. births, deaths, organismal growth). Derived operationalizations depend also on assumptions regarding demographic stochasticity, environmental stochasticity, feedbacks whereby births, deaths, and organismal growth cause environmental change, and the impact of migration and niche construction on which environment is experienced. After reviewing existing derived fitness operationalizations, we propose a new one tailored to balancing selection. Population genetic models generally sidestep ultra-high-dimensional phenotype space and genotype spaces by instead deriving the long-term evolutionary fate of a lower-dimensional set of genetically encoded “strategies”. Strategies (e.g. costly developmental commitment to producing armaments) are causally upstream from realized phenotypes (e.g. armament size). While selection is best understood in terms of differences in *organismal* vital rates, its derived outcomes are most easily understood as properties of *genetic lineages*.

Keywords:

Invasion fitness, Malthusian parameter, individuality, theoretical population genetics, bet-hedging, life history strategy, density-dependent selection

“Fitness: Something everyone understands but that no one can define precisely” (Stearns 1976)

51 Introduction

52 Darwin's theory of evolution by natural selection did not launch a professional discipline of
53 evolutionary biology until the Modern Synthesis of the early twentieth century, in which the
54 role of mathematical population genetics was key (Provine 1978). Central to this
55 mathematization was "fitness," which turned intuitions about "the ability to survive and
56 reproduce" or "what tends to be favored in the struggle for existence" into more formal
57 quantitative operationalizations. Resulting models are used to derive non-obvious insights
58 (Servedio et al. 2014). One important historical role for fitness models was to deduce that
59 the timescale of evolution by natural selection is fast enough to make selection a
60 predominant factor in evolution, even with reasonably small selection coefficients (Provine
61 1978; Charlesworth 2020). Models of fitness can also be fit to sequence data, e.g. to detect
62 loci under recent selection (Enard 2021); this falls outside the scope of the current
63 manuscript.

64 From the outset of its mathematization, fitness has been operationalized in different
65 ways (Ariew & Lewontin 2004; Orr 2009). Haldane (1927) used the expected *absolute*
66 number of surviving offspring, while the influential Wright-Fisher model used the expected
67 *relative* contribution to the gene pool in the next generation (Fisher 1930; Wright 1931).
68 Theoretical population genetic models *assign* some version of expected short-term fitness to
69 genotype-environment combinations, from which they mathematically *derive* longer-term
70 outcomes. For example, Haldane (1927) assigned births per generation to a mutant
71 genotype, and then derived its probability of fixation. Each such model thus involves at least
72 two operationalizations of fitness: the assigned short-term fitness, and the derived long-term
73 outcome or probability distribution of outcomes.

74 We review a variety of models and corresponding fitness operationalizations, their
75 motivation, and simplifying assumptions, using annual plants and the Hawk-Dove game
76 (Maynard Smith & Price 1973) as illustrative examples. We first focus on operationalizations
77 that are at least sometimes assigned, then on operationalizations in which fitness is always

78 derived. We then propose a conceptual scheme describing how models give insights into the
79 fate, under natural selection, of the organismal strategies of interest to biologists. We argue
80 that short-term fitness is best operationalized via *organismal* vital rates (births, deaths,
81 organismal growth) plus organismal effects on the experienced environment through
82 migration and/or niche construction, while long-term fitness is best operationalized for
83 *genetic lineages*.

84 Assigned Fitness

85 Absolute fitness

86 Absolute fitness W describes the expected number of surviving offspring that a
87 (hermaphroditic or asexual) individual produces after reaching reproductive maturity.
88 Equivalently, it describes a juvenile's expected number of offspring (reversing the order of
89 survival and reproduction). Either way, it is the expectation over one complete life cycle or
90 'generation' of both survival and reproduction.

91 The seminal use of assigned absolute, per-generation fitness was to derive the
92 fixation probability of a new beneficial mutation. Haldane (1927) considered a resident (R)
93 population of constant size, such that $W_R = 1$. He then considered the fate of a new lineage
94 produced by a beneficial mutation. Individuals carrying the mutation have $W_I = 1 + s$, where
95 the selective advantage $s > 0$. With some simplifying assumptions, including a Poisson
96 distribution of offspring and $s \ll 1/2$, Haldane (1927) derived the probability that the
97 beneficial mutation escapes extinction to "invade" as $2s$ (Fig. 1). Beyond the Poisson
98 distribution, invasion probability is $2s/\sigma^2$ where σ^2 is the variance in offspring number
99 (Barton et al. 2007, p. 25).

100 This example illustrates how the long-term fate of a mutant (probability of extinction)
101 is derived from the short-term probability distribution of offspring number. Evolutionary
102 success under natural selection cannot be reduced, even in a very simple model, to a single
103 number such as W . Larger variance in reproduction σ^2 increases the extinction probability,
104 which can loosely be understood in terms of a lower signal (s) to noise (σ^2) ratio.

105 Haldane's assignment of absolute fitness W to genotypes is rarely used outside of
106 this example of a rare beneficial mutant. All biological populations are density regulated,
107 meaning that high W causes an increase in population density, which in turn reduces W
108 (Haldane 1956; Nicholson 1957). For assigned constant W , the invading mutant lineage
109 instead experiences unbounded exponential growth.

110 Relative fitness

111 Assigning *relative fitness* instead of *absolute fitness* sidesteps the issue of unbounded
112 exponential growth. Relative fitness models treat the proportions of variants, rather than their
113 absolute abundances. To motivate this, Crow and Kimura (1970, pp. 25-26) derived relative
114 fitnesses w_k from assignments of absolute fitnesses W_k in the context of exponential
115 population growth or decline. On this basis, they argued for simplified models in which w_k
116 values are directly assigned, sidestepping assignments of W_k . Measurement theory has also
117 been invoked as supporting the use of relative fitness over alternatives (Wagner 2010).

118 In the simplified models that have become standard within population genetics,
119 relative fitness is defined as proportional to the expected fraction of the next generation that
120 is descended from the focal genotype or individual. In the simple case of asexual
121 reproduction, if $p_i(t)$ is the proportion of the population with genotype i at time t , and w_i is
122 the relative fitness of type i , then its expected proportion in the next generation is

123
124
$$p_i(t + 1) = p_i(t) \frac{w_i}{\bar{w}}.$$

125
126 Normalization by the population mean of relative fitness $\bar{w} = \sum_{\text{all } k} p_k(t) w_k$ can be used
127 either to keep the population size constant, or to impose a different demographic model such
128 as exponential growth. Importantly, this normalization derives absolute fitness W in a
129 frequency-dependent way from relative fitness w - the opposite direction of Crow and
130 Kimura's justification for the assignment of relative fitness.

131 A classic use of relative fitness assignments is in Wright-Fisher models that select
132 among parent genotypes to produce expected fecundity w_i/\bar{w} . All adults then die - a
133 potentially appropriate model for an annual plant. The finite size N of Wright-Fisher
134 populations enables the derivation of fixation probabilities also for deleterious mutations,
135 which never avoid extinction under the branching process treatment of Haldane (1927).
136 Finite population size also enables derivation of the expected “sojourn” time prior to
137 extinction or fixation (Charlesworth 2020). Conditional on fixation, $\bar{\tau} \sim 2 (\ln(sN) + \gamma)/s$
138 generations in a haploid Wright-Fisher model (Fig. 1) where $\gamma = 0.5772$ is Euler’s constant
139 (Hermisson & Pennings 2005). Sojourn times were historically key to proving that natural
140 selection works sufficiently rapidly to be a major cause of evolution (Provine 1978). In the
141 modern era, sojourn times (or, if backward time is considered, coalescence times) are used
142 when inferring a population’s history of selection and demography from sequence data
143 (Gutenkunst et al. 2009; Keightley & Halligan 2011; Ronen et al. 2013; Charlesworth 2020;
144 Liu & Fu 2020; Excoffier et al. 2021).

145 A key limitation of models in which relative fitness is assigned to genotypes is that
146 they do not allow the consequences of selection to feed back onto population density. In
147 other words, the population size N is externally set, independently of mean population
148 fitness. Problematically, no matter how low population fitness w drops, the externally set
149 population size N will not decline, contradicting the desired behavior that low fitness should
150 indicate an increased tendency to go extinct. A second, related limitation is that relative
151 fitness cannot be compared across populations.

152 Vital rates are the “ultimate” assigned values

153
154 Vital rates describe rates of organismal growth, deaths, and reproduction. To complete a
155 generation, seeds must germinate and survive to become seedlings, then survive from
156 seedlings until they reach reproductive maturity, and then produce and disperse seeds. This
157 description of three “fitness components” encompasses three vital rates for three life history

158 transitions: the first two include both survival and growth, while the third includes only
159 reproduction. Per-generation absolute fitness is the product of fitness components, each
160 describing survival and/or reproduction during a different life history transition, within a fixed
161 sequence. However, when the sequence varies, different values of fitness components are
162 derived from the same vital rates, e.g. for a seed that survives within a seed bank for a
163 variable number of years, each time without growth.

164 Like Metcalf (2007), Doebeli (2017), and Matheson (2023), we propose making
165 survival and reproduction core to our scheme, and assigning corresponding values of death
166 rate d and birth rate b , rather than of “fitness”, to phenotypes in an environment. On the
167 surface, many models assign relative or absolute fitness values. In fact, a classic model
168 such as Wright-Fisher is better seen as assigning a variable birth rate combined with a
169 constant adult death rate, from which per-generation fitness is implicitly and trivially derived.
170 In more complex models, e.g. of populations perturbed away from demographic equilibrium,
171 selection on fecundity/juveniles does not produce the same allele frequency trajectory as
172 selection on adult death rates (Benton & Grant 2000; Bertram & Masel 2019).

173 Derived fitness operationalizations

174 The fitness operationalizations presented so far are sometimes assigned to genotypes (as a
175 function of their current environment). Next, we consider properties that are rarely if ever
176 assigned to genotypes, but derived from assigned fitness operationalizations. To illustrate
177 them, we add a seed bank to our annual plant example.

178 The Malthusian parameter

179
180 The Malthusian parameter (Malthus 1798; Fisher 1930) or intrinsic growth rate r (Lotka
181 1907) quantifies how quickly a genetic lineage tends to grow or shrink, in absolute time units
182 (e.g. days), rather than in the per-generation time units of the relative and absolute fitness
183 operationalizations above. While usually specified as a form of absolute fitness, a relative
184 fitness version can be obtained as $r_i' = r_i - \bar{r}$ where r is the mean Malthusian parameter,
185 with r_i' analogous to w_i/\bar{w} discussed above. Occasionally, r is an assigned parameter as a

186 technical matter to allow the use of differential equations (Desai & Fisher 2007). In most
187 studies, however, r is a derived fitness operationalization. Doebeli et al. (2017) argue that r
188 should always be derived rather than assigned.

189 For the non-overlapping generations treated by the Wright-Fisher model, r and W
190 contain the same information, albeit in different units. However, consider a simple scenario
191 of overlapping generations, where individuals produce offspring at rate b and die at rate d .
192 The Malthusian parameter is $r = b - d$ with time units, whereas per-generation absolute
193 fitness is $W = b/d$ (births occurring during expected lifespan $1/d$). For example, when $b =$
194 0.2 and $d = 0.1$, then $W = b/d = 2$ (average of 2 offspring per generation), while $r = b -$
195 $d = 0.1$ (lineage is growing with exponential growth rate 0.1 per external time unit such that
196 $y(t) = y(0)e^{rt}$).

197 When generations overlap, neither W nor r can be derived given information only
198 about the other, and they provide information about different things (De Jong 1994). The
199 Malthusian parameter tells us what allele frequencies to expect at a specified time in the
200 future (r is a *rate*). For example, sojourn time (Fig. 1, $\bar{\tau}$ in mutation 4) depends on
201 differences in r , whereas differences in W , combined with σ^2 , tell us the *probability* that a
202 rare beneficial mutation will escape extinction (Fig. 1, mutations 1-4).

203 Selection can act on differences in one quantity (r or W) even given equality for the
204 other. For example, consider a trade-off between b and d such that $W = b - d$ remains
205 constant. Importantly, r need not be constant under this constraint. In the wake of a
206 disturbance that kills many individuals from a population previously at equilibrium, selection
207 will favor larger b and d , because this increases $r = b - d$, enabling the type with the faster
208 life history strategy to more quickly rise back up to carrying capacity (Stearns 1992).

209 The Malthusian parameter generally depends on all three kinds of vital rate: deaths,
210 births, and growth. For example, consider adult plants (A) that die at rate d and give birth at
211 rate b to seeds (S) that grow into reproductively mature adults at rate g . For simplicity, we
212 neglect seed death. This yields the following differential equations:

213
$$\begin{matrix} dS(t)/dt \\ dA(t)/dt \end{matrix} = \begin{pmatrix} -g & b \\ g & -m \end{pmatrix} \begin{pmatrix} S(t) \\ A(t) \end{pmatrix}$$

214 The Malthusian parameter is the dominant eigenvalue of the 2×2 matrix above: $r =$
 215 $\frac{1}{2}(\sqrt{4bg + d^2 + g^2 - 2dg} - d - g)$. This summary of the short-term fitness consequences of
 216 vital rates illustrates the need to include g ; note that $r \rightarrow b - d$ as $g \rightarrow \infty$. In contrast, $W =$
 217 b/d , with no dependence on g .

218 A common use of the Malthusian parameter is to describe “invasion fitness”, meaning
 219 whether and at what speed a new mutant genotype I deterministically invades a population
 220 of “resident” genotype R at equilibrium abundance $\widehat{N}_R(t + 1) = \widehat{N}_R(t)$ (Metz et al. 1992). To
 221 illustrate this, consider an annual plant population in which a seed germinates with
 222 probability g per year to produce an expected f seeds, or else survives with probability $1/d$
 223 in the seed bank. Now our vital rates are f , g , and d . We capture the dependence of
 224 fecundity on seedling density using parameter α . In external timesteps $t = 1$ (rather than
 225 per-generation terms), types $k = R, I$ (resident and invader) obey:

226
$$E[N_k(t + 1)] = \underbrace{\frac{(1/d)(1 - g_k)N_k(t)}{\text{Number of non-germinating seeds that survive}}}_{\text{Number of non-germinating seeds that survive}} + \underbrace{\frac{N_k f g_k}{1 + \alpha \sum_{\text{all } j} g_j N_j(t)}}_{\substack{\text{New seeds produced by germinating} \\ \text{individuals that survive density effects}}} \quad (1)$$

227 For a rare invader ($N_I(0) \ll \widehat{N}_R(t)$), invasion fitness is equal to the absolute Malthusian
 228 parameter:

229
$$r_I = E \left(\ln \frac{N_I(1)}{N_I(0)} \right) \quad (2)$$

230 which depends on resident density \widehat{N}_R via the denominator in the rightmost term of Equation
 231 1. Invader I tends to invade if and only if $r_I > 0$.

232 So far, the optimal strategy is always to germinate, i.e. $r_I > 0$ if and only if $g_I > g_R$.
 233 This is because there is so far no advantage to being dormant, to offset the risk of dying
 234 while in the seed bank. This changes when we consider fluctuating environments below, in
 235 which germination is sometimes futile.

236 Fitness across a variable environment

237 Most organisms experience environmental heterogeneity that affects their vital rates. E.g.,
238 plant seed production f depends on abiotic (e.g. rainfall), and biotic density-dependent
239 (MacArthur 1962; Tilman 1982; Travis et al. 2023) and frequency-dependent (Tilman et al.
240 2020) environmental factors. Environmental variation can be spatial and/or temporal.

241 Given spatial environmental variation, migration enables organisms to affect which
242 environment(s) they encounter. Some forms of migration, e.g. seed dispersal, are closely
243 coupled to a life history transition, but can be conceptually separated into a migration
244 phenotype in the old location, followed by vital rates of birth, death, and growth in the new
245 location. Similarly, organisms can indirectly modify their vital rates via phenotypes that
246 physically alter their local environment (niche construction; Odling-Smee et al. 1996).
247 Selection on migration and niche construction phenotypes is included within the Malthusian
248 parameter calculated across spatial environmental variation. I.e., the Malthusian parameter
249 is derived not just from assigned vital rates, but also from assigned migration and niche
250 construction rates.

251 We consider temporal variation in the environment $e(t)$ via an extension of Equation
252 (1) in which germinating seeds produce zero offspring during drought years, such that
253 fecundity

$$254 \quad f(e(t)) = \begin{cases} f & \text{in good years with probability } p \\ 0 & \text{in bad years with probability } 1 - p \end{cases}$$

255 Instead of the instantaneous Malthusian parameter in a single environment, we take, as
256 invasion fitness, its expected value across the distribution of environments $e(t)$:

$$257 \quad r_I = E_{e(t)} \left(\ln \frac{N_I(t+1)}{N_I(t)} \right). \quad (3)$$

258 This is known as the *geometric mean fitness* because it corresponds to the geometric mean
259 of absolute per-generation or per-time-step W . It is equivalent to the arithmetic mean of the
260 Malthusian parameter over environments (Takacs & Bourrat 2022, 2024). In more complex

261 scenarios when multiple life stages are affected by the environment, a generalization of the
262 Malthusian parameter known as the Lyapunov exponent can be used
263 (Cohen 1979; Metz et al. 1992; Kussell & Leibler 2005).

264 While germination probability $g = 1$ maximizes r_I in a constant environment, it results
265 in complete extinction in a bad year, and so a more conservative $g_I < 1$ maximizes r_I in a
266 temporally varying environment. This is an example of evolutionary *bet hedging* (Cohen
267 1966; Seger & Brockmann 1987; Frank 2011a).

268 In adaptive dynamics (Metz et al. 1995), the standard practice is to assume that
269 evolution moves in the direction that maximizes invasion fitness, given infinitesimal
270 perturbations to parameters controlling strategies (e.g., g_I infinitesimally differs from g_R). In
271 the seed bank model, evolved g_R then achieves $r_I < 0$ for all $g_I \neq g_R$ (an “evolutionary stable
272 strategy”; Geritz et al. 1998).

273 274 Fixation Probability Ratio

275 Derived fitness operationalizations attempt to capture which strategies will become
276 prevalent, if present, under evolution by natural selection. Although individuals die within a
277 short timescale, they embody a strategy/type (e.g., germination probability) that lasts over a
278 longer timescale, due to being genetically encoded. Consider a genetic lineage (Akçay &
279 Van Cleve 2016; Graves & Weinreich 2017) consisting of all gene copies descended from a
280 new mutation encoding a change in germination probability. In the long term, this lineage
281 either fails (goes extinct), or succeeds (fixes in the population). The probabilities of lineage
282 fate can be used to construct a derived operationalization of fitness.

284 By equating $r_I > 0$ with success, invasion fitness (equations 2-3) neglects chance
285 extinction. Recalling that the probability of invasion $2s/\sigma^2$, invasion fitness does nothing to
286 capture genetic variation affecting demographic stochasticity σ^2 . Stochasticity in the series
287 of environments also contributes to extinction (King & Masel 2007).

288 Consider an extension of the annual plant example in which genotype abundance is
289 a discrete random variable, X

290
$$N_k(t + 1) = X(\mu, \sigma^2). \quad (4)$$

291

292 Equation 1 is the special case with $\sigma^2 = 0$. As in Haldane (1927), $N_k(t + 1)$ can be 0 even
293 if $\mu > N_k(t)$. The maximal probability that an invader lineage fixes requires at a lower value
294 of g_I than the maximal geometric mean growth rate (Fig. 2). Larger g causes greater
295 fluctuations in N – the strategy with largest r therefore increases extinction risk (Adler &
296 Drake 2008; Constable et al. 2016; Pande et al. 2020; Pande et al. 2022). Invasion fitness r
297 thus does not fully capture the long-term fates of lineages.

298 To capture demographic stochasticity, we can compare fixation probabilities to those
299 of neutral alleles (Nowak et al. 2004). To also capture environmental stochasticity, we can
300 use the ratio of the probability with which allele 1 invades a population in which allele 2 is
301 resident : the probability with which allele 2 invades a population in which allele 1 is resident
302 (Masel 2005). When mutation between the two alleles is symmetric and rare, the fixation :
303 counterfixation ratio describes the odds with which a population will be found fixed for allele
304 1 vs. allele 2. This makes it directly applicable to empirical situations such as quantifying
305 preferences among codons (Bulmer 1991), in which there is sufficient data across an
306 ensemble of comparable instances.

307 Note that when mutation is not symmetric, the direction and degree of mutational
308 asymmetry also affect the odds with which a population will be found fixed for allele 1 vs.
309 allele 2, which are given by $\mu_{j \rightarrow i} p_{\text{fix}}(j \rightarrow i) : \mu_{i \rightarrow j} p_{\text{fix}}(i \rightarrow j)$. This ratio includes both our
310 fitness operationalization $p_{\text{fix}}(j \rightarrow i) : p_{\text{fix}}(i \rightarrow j)$, and mutation bias $\mu_{j \rightarrow i} : \mu_{i \rightarrow j}$. The relative
311 mutation rates matter because a variant must first appear in the population before it can be
312 subject to natural selection. Fitness cannot be equated with quantifying “what evolution
313 makes prevalent”, because natural selection is not the only cause of evolution (Stoltzfus &
314 Yampolsky 2009).

315 The evolved mutation rate is a good example of an outcome determined in part by
316 mutation bias. There are more mutations that increase the mutation rate (mutators) than
317 decrease it (antimutators). However, indirect selection against deleterious mutation load

318 favors a lower mutation rate (Johnson 1999a, b). This results in a mutation-selection-drift
319 balance at some fairly low mutation rate (Lynch 2008). Operationalizing fitness as the ratio of
320 fixation : counterfixation probabilities readily handles the complexities of indirect selection
321 that arise during the evolution of mutation rate. A lineage approach is also useful for
322 understanding the evolution of cooperation (Akçay & Van Cleve 2016).

323 324 How do we operationalize fitness under balancing selection? 325

326 Balancing selection is a challenge to all three derived operationalizations presented above.
327 Sometimes two alleles can each invade an equilibrium population of the other, such that
328 both variants are maintained by balancing selection (Fig. 3A). Characterizing cases of
329 *mutual invasibility* is common in evolutionary game theory (Maynard Smith & Price 1973),
330 adaptive dynamics (Metz et al. 1995), and theoretical community ecology (Turelli 1978;
331 Chesson 2000). While coexisting at equilibrium, both types have a geometric mean fitness of
332 1. Both fixation probabilities are much lower than the neutral $1/N$ or $1/2N$, and taking the
333 ratio of fixation probabilities contains little information about the outcomes natural selection
334 tends to produce.

335 The qualitative intuition that “both types are fit” can be operationalized in stochastic
336 terms by noting that both types invade with a high probability of “establishment” (rather than
337 fixation) when rare. Establishment means reaching high enough abundance such that
338 deterministic dynamics dominate, with subsequent stochastic extinction being rare (Desai &
339 Fisher 2007). A “high” establishment probability can be operationalized by comparing an
340 invader’s probability of reaching a given frequency to that of a neutral reference invader (i.e.
341 one indistinguishable from the resident).

342 To quantitatively operationalize fitness under balancing selection, we propose taking
343 the time-integral of mutant lineage abundance from introduction into a resident population of
344 the other type, until stochastic extinction. We then take the ratio of these integrals, switching
345 which is the resident and which is the invader. This is illustrated in Fig. 3 for the Hawk-Dove
346 game. The time-integral is slightly larger than the product of three informative components:

347 establishment probability, sojourn time from introduction until extinction conditional on
348 establishment, and mean abundance during its sojourn (Fig. 3B-G). Minor deviation of
349 overall fitness from the product of these three components comes from neglecting
350 abundance conditional on non-establishment. As a technical matter to prevent the sojourn
351 time from being inflated by fixation events, a model should disallow transitions to the
352 absorbing boundary of invader fixation. Our metric captures the potential vulnerability of an
353 abundant type to extinction e.g. from disturbance (Tilman et al. 1994), which would be
354 missed if we used abundance or biomass (Van Valen 1975) from the corresponding mean
355 field model.

356

357 The role of fitness within evolution by natural 358 selection

359

360 Fig. 4 illustrates how models describe causality during evolution by natural selection. We
361 distinguish between three aspects of the environment. The *selective environment* interacts
362 with phenotypes to give rise to a particular organism's vital rates. (Note that our use of
363 "selective environment" better corresponds to the "ecological environment" of Brandon
364 (1990).) Here, we restrict the term *phenotype* to realized organismal properties (e.g., body
365 size) or behaviors (e.g., migration, aggression). Extended phenotypes (Dawkins 1982) are
366 captured by feedback from phenotype to the environment (Fig. 4). We refer to aspects of the
367 environment that directly change phenotypes as the *developmental environment*.

368 Organismal *strategies* describe allocation of scarce resources in pursuit of phenotypes.

369 Strategies are shaped by genotype and/or by a plastic response to the *informational*

370 *environment* – the cues that organisms respond to, prior to the direct effects of the

371 environment on development. Responses to the informational environment (e.g., using

372 locally low resource levels as a cue to migrate), if any, reflect the history of adaptation. In

373 contrast, we consider intrinsic effects of the environment on phenotypes (e.g., reactions

374 proceed faster at higher temperatures; Brown et al. 2004) to be part of the developmental

375 environment. Note that the same environmental factor (e.g., temperature) can be part of all
376 three aspects of the environment, by giving information, altering development, and imposing
377 selection.

378 Different models simplify the Fig. 4 scheme in different ways. Commonly assigned
379 fitness operationalizations, e.g. per-generation absolute fitness $W = b/d$, summarize the
380 differential *vital rates* that embody natural selection in the short-term (Fig. 4, small shadow).
381 In the Wright-Fisher model, genotypes vary in b , whereas in Haldane's model and the Moran
382 model (Moran 1958), they could also vary in d . Haldane holds the environment constant,
383 whereas the Wright-Fisher model lets the selective environment (represented by allele
384 frequencies) affect the absolute vital rate b produced by a given genotype.

385 Derived fitness operationalizations are more complex summaries of the longer-term
386 fate of genetic lineages, including the influences of demographic stochasticity, migration,
387 niche construction, and spatial and temporal environmental variation (Fig. 4, large shadow).
388 Natural selection produces differential vital rates, while the long-term outcomes of natural
389 selection are embodied in long-term lineage fate. Simple population genetic models provide
390 insights into the efficacy and timescale over which natural selection may operate (e.g.,
391 invasion probability $\sim 2s/\sigma^2$, and sojourn time $\sim 2(\ln(sN) + \gamma)/s$. However, phenotype-
392 agnostic assigned fitness operationalizations do not provide insights into the underlying
393 biological mechanisms through which natural selection favors particular traits.

394 Directly assigning vital rates enables us to ask, for example, how natural selection
395 acts during the evolution of dormancy, operationalized as a genetically encoded 1-locus
396 strategy to germinate with probability g per year. More sophisticated strategies might involve
397 active sensing to exploit the informational environment (Kussell & Leibler 2005). For
398 example, selection might favor a reaction norm of higher g given higher soil moisture. A
399 sufficiently reliable environmental cue begets a shift from bet hedging to plasticity (Botero et
400 al. 2015). Selection acts on phenotypes (germinating vs. not) as a function of both biotic
401 environment (population density) and abiotic environment (drought vs. non-drought year), to

402 produce vital rates whose impact on genetic lineages, over time, can be summarized by
403 derived fitness operationalizations. This type of model provides insights into the biological
404 mechanism through which a lineage with a mutation (g_I) “wins”.

405

406 Strategies

407 Strategies are intermediate between genotype and phenotype. In a broader sense,
408 strategies are a form of phenotype, describing what an organism prioritizes given
409 constraints, often entailing commitment to developmental pathways and/or behaviors.
410 Strategies can be seen as setting organismal goals (at least in organisms capable of
411 cognition). The decision to commit is informed by genotype and by the informational
412 environment, with its success in achieving the anticipated phenotype affected by the
413 developmental environment.

414 As a simple example, consider a “Hawk” strategy from the Hawk-Dove game in
415 evolutionary game theory (Maynard Smith & Price 1973). Briefly, Hawks fight for resources
416 while Doves avoid conflict. In classic models, the developmental environment is neglected,
417 and having a Hawk strategy fully specifies behavioral phenotypes. One’s opponent (Hawk or
418 Dove) constitutes one’s selective environment, and knowledge of their past behavior (if
419 included in the model variant) constitutes the informational environment. In contrast, we
420 conceptualize a Hawk *strategy* not just as behaviors within the narrow confines of game
421 theory, but as a developmental commitment toward *developing a set of phenotypes* (both
422 armaments and behaviors) that are relevant for implementing aggression. This allows for the
423 possibility that developmental conditions (e.g., insufficient resources) may prevent a Hawk
424 from e.g., achieving large enough body size or armaments to be successful. The individual
425 may then switch strategies, treating developmental inputs as part of the informational
426 environment.

427 Applying our distinction between strategy and phenotype to the seed bank model
428 (equation 1) is more subtle. A seed’s realized phenotype is defined by germination (or lack
429 thereof) while its strategy is embodied in the stochastic gene circuitry that is an adaptation

430 for achieving a probability of germination g within the historical range of environments. An
431 organism's realized phenotype arises from the latter via a noise within the developmental
432 environment (Frank 2011b). An unanticipated developmental environment (e.g. a prolonged
433 hard freeze) could cause the outcome (germinating with probability g) to deviate from the
434 strategy.

435 Strategies include investing in rapid growth given low population density, or in
436 competitiveness or persistence given high population density (Grime 1988; Bertram & Masel
437 2019). This was originally formalized as r - vs. K -selected "strategies" (MacArthur 1962;
438 Roughgarden 1971), where r is the Malthusian parameter at low density (and a prefactor of
439 it also at higher densities), and K describes susceptibility to density-dependence (similar to
440 $1/\alpha$ in equation (1)). A trade-off between investment in r vs. K was assumed, with the
441 resulting "strategy" reflecting an organism's position along that trade-off. However, r and K
442 are often positively correlated with slope near 1 in empirical studies (Luckinbill 1978, 1979;
443 Valle et al. 1989; Kuno 1991; Hendriks et al. 2005; Fitzsimmons et al. 2010), in agreement
444 with some process-based theoretical models (Travis et al. 2023). While there does seem to
445 be a fast-slow continuum, contemporary life history theory categorizes strategies in other
446 ways (Salguero-Gómez et al. 2016; Healy et al. 2019; Malik et al. 2020; Bruggeman et al.
447 2023).

448 Organisms are capable of an extraordinary variety of phenotypes. The "functional
449 trait" literature in community ecology attempts to reduce this dimensionality, by focusing on
450 phenotypes (e.g., wood density, seed size, metabolic rate) that are most closely tied to
451 strategies and vital rates (McGill et al. 2006; Yang et al. 2018). In contrast, vital rates come
452 in only three key varieties, applied to different life history stages. Organismal strategies
453 might have far lower dimensionality than downstream functional traits or other organismal
454 phenotypes, in a manner that helps provide generalizable insights. Strategy space might be
455 both small enough and concrete enough to give coherence to the organism's developmental
456 commitments, as well as to scientists studying them. Assigned and derived fitness

457 operationalizations are key components of the models that serve to clarify how natural
458 selection acts on strategies.

459 We find the 3-dimensional scheme of Grime to be a promising starting point for
460 characterizing strategies. Grime (1977, 1988, 2001) hypothesized that trade-offs shape
461 species into three types of specialization – “ruderals” tolerate harsh abiotic environments,
462 “competitors” excel at high population density, and “colonizers” rapidly disperse to
463 ephemeral resources. Each strategy is closely tied to vital rates. High-dimensional
464 phenotype space among e.g., coral species can be simplified via a space of just these three
465 strategies (Darling et al. 2012). Our simple example of a seed bank illustrates how
466 organismal strategies can be described with reference to vital rates (and potentially also
467 migration and niche construction phenotypes) in order to gain insight into how populations
468 evolve within strategy space.

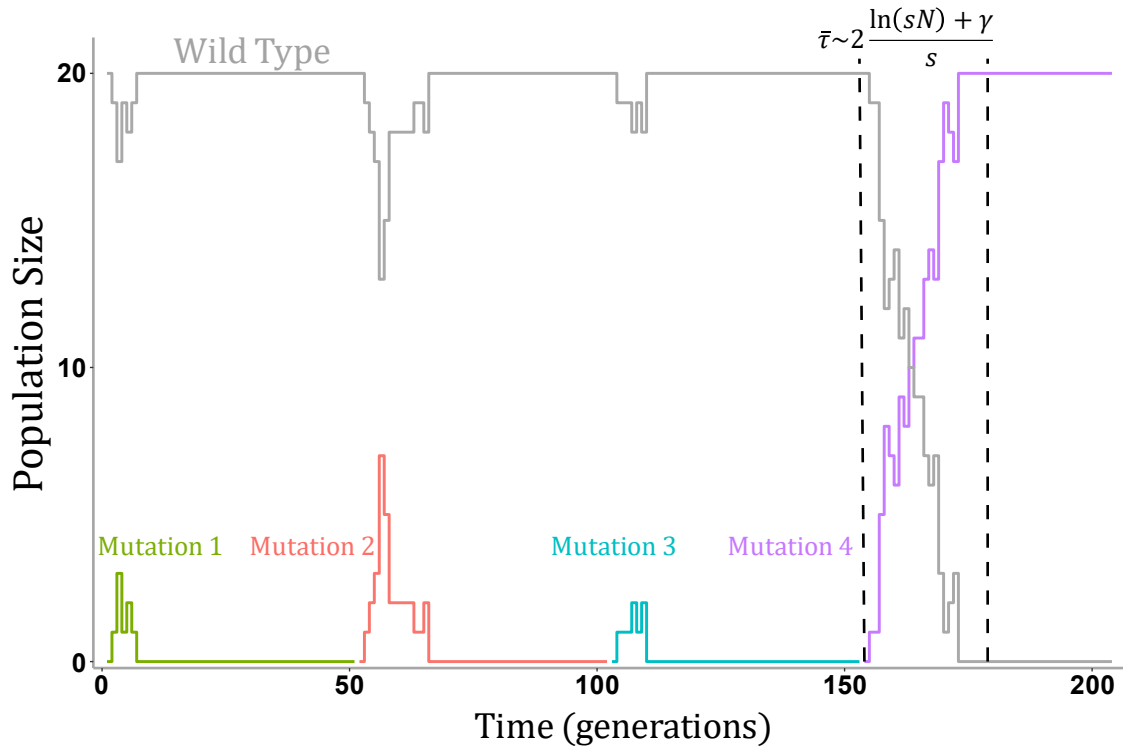
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470 Conclusion

471 Both genotype space and phenotype space are huge and must be simplified to produce
472 generalizable biological insight. Organismal “strategies”, intermediate between genotypes
473 and phenotypes, capture biological questions of interest, and give rise to vital rates,
474 migration rates, and niche construction phenotypes, which influence the quantifiable fate of
475 genetic lineages. Traditional relative fitness and absolute fitness implicitly assign vital rates
476 to organisms. From assigned vital rates, other fitness operationalizations (i.e. ways of
477 quantifying what natural selection favors) are derived to describe evolutionary outcomes.
478 Variations on the Malthusian parameter capture adaptation speed, while the probability of
479 invasion is captured by the fixation : counterfixation probability ratio. We build on the latter to
480 propose a new, lineage-based fitness operationalization suitable for describing fitness under
481 balancing selection.

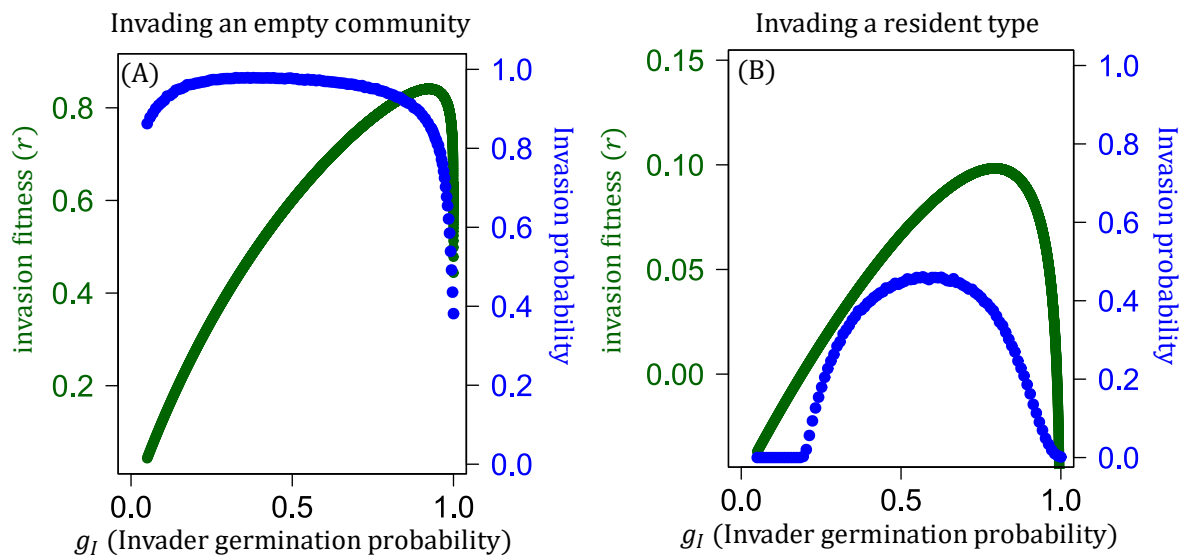
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Figures



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Fig 1: Fixation probability and sojourn time capture different long-term consequences of natural selection. Representative Wright-Fisher simulation of a population of size $N = 20$ in which an allele with selection coefficient $s = 0.125$ appears repeatedly by mutation. The mutant fixes with probability $\approx 2s/\sigma^2 = 0.25$. The sojourn time τ describes the number of generations before a mutation fixes (given it does not go extinct) with mean $\bar{\tau} = 2(\ln(sN) + \gamma)/s$. Each color indicates a different mutation. The interval between the dashed lines depicts $\bar{\tau}$; slightly shorter than the realized value of τ in this simulation.

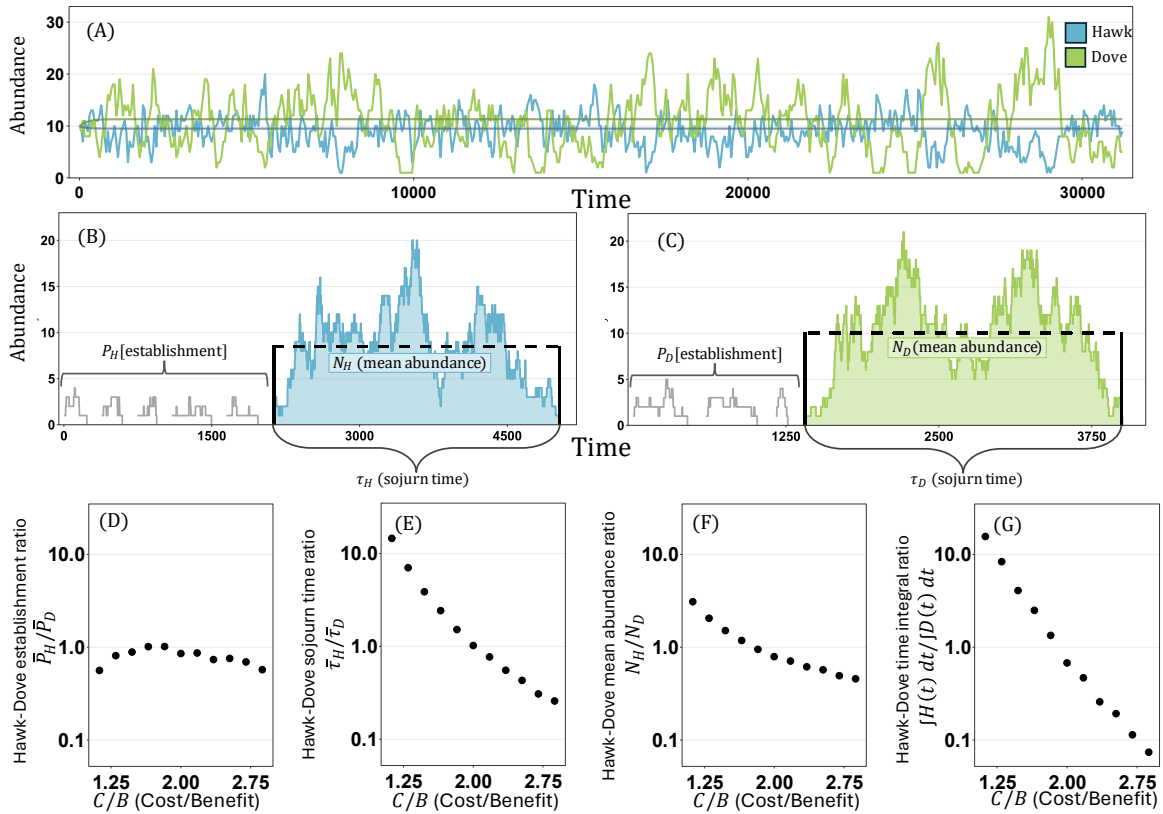


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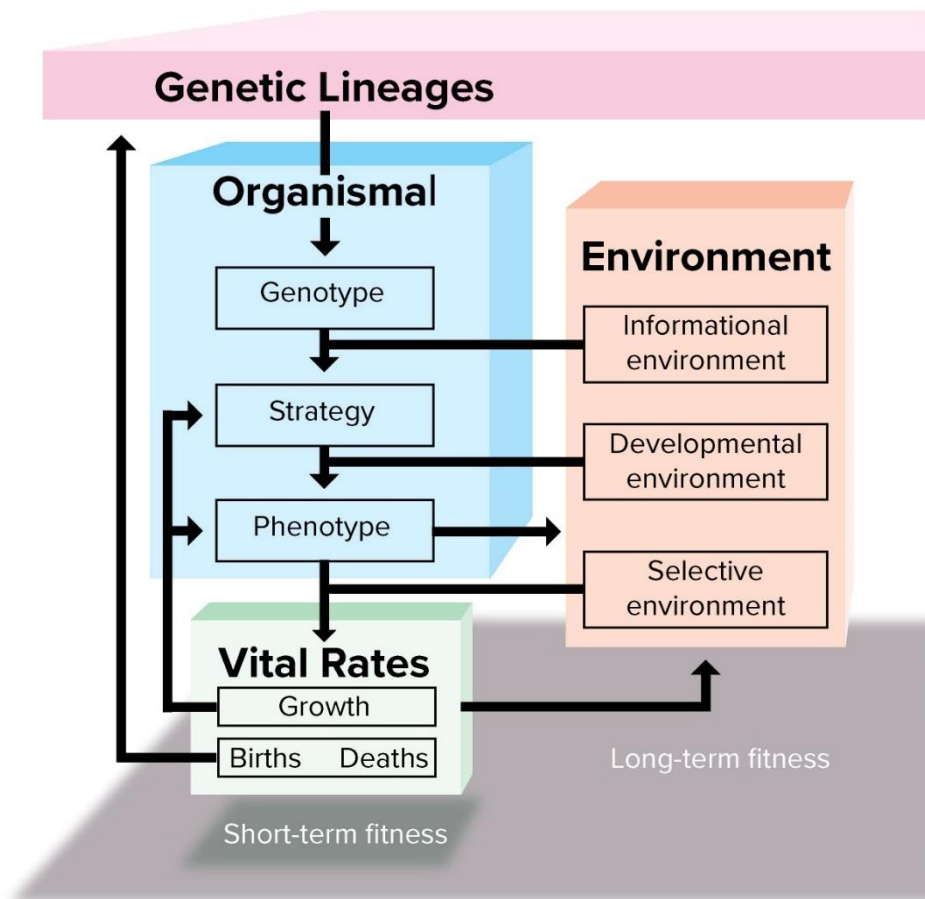
Fig 2: The invader's initial geometric mean growth rate is maximal for a higher germination probability than that which maximizes invasion probability. Seed banks are simulated given demographic stochasticity, and probability $p = 0.95$ that a given year allows reproductive success. An invader with germination probability g_I is introduced into an empty community (A) or a resident population of $g_R = 0.2$ (B); note the different y-axis scales for r in green. Invasion probability (blue) is defined in A as the probability that invader persists for at least 20 generations, and in B as the resident going extinct before the invader does. Invasion fitness (green) peaks at $g \sim 0.8$, but invasion probability (blue) is highest for moderately low values of g_I . Invasion fitness relative to a resident (B) is only positive for $g_I > g_R$. Invasion probability peaks at an intermediate value for which $g_I > g_R$ and is ≈ 0 when $g_I < g_R$. Adaptive dynamics models consider only infinitesimal changes in g_I relative to g_R , and only consider the invasion fitness. Parameters: $d = 1.053, f = 3, \alpha = 0.025$.

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Fig 3: Our proposed operationalization of long-term fitness for a balanced polymorphism. We simulated a discrete-time stochastic Hawk-Dove game, code available on GitHub. A Hawk competing against a Dove always obtains the contested resource and receives a benefit B ; a Hawk competing with a Hawk either gains the benefit B , or experiences a cost of fighting C , with equal probability; competing Doves split the benefit B evenly. Each timestep, individuals (1) die with probability d and then, if alive (2) produce offspring according to a Poisson distribution. The mean of the Poisson distribution for a type (Hawk or Dove) is determined by a baseline birth rate, payoffs that depend on the frequencies of Hawks and Doves in the population as well as B and C , and a density-dependent parameter such that births decrease with increasing density. A stable polymorphism requires $B - C < 0$. All points shown are in the parameter regime for which coexistence occurs under a mean field approximation. (A) Throughout most of the time series, Hawks and Doves coexist with abundances near the corresponding mean (horizontal lines). However, occasionally, one type falls to low abundance, and would go extinct in the absence of the reflecting boundary used in the simulation. The three components of our novel fitness operationalization are illustrated for the Hawk (B) and Dove (C). When a previously absent Hawk or Dove is introduced by mutation or migration, it must establish (increase from rarity when the other type is at equilibrium). We operationalized establishment as reaching the equilibrium frequency in the corresponding mean field model. Establishment probability (P_H and P_D) depends on various parameters of the model; gray time series data depict failures to establish. After establishment (colored blue and green time series data), the Hawks and Doves persist for a sojourn time (τ_H and τ_D) until eventual extinction. During the sojourn, the abundance of Hawks and Doves fluctuate around the mean (N_H and N_D). (D) – (E) show the ratio of the fitness components as a function of C/B . Each point shows the ratio of mean values from 7500 simulations of the Hawk invading the Dove and *vice versa*. (D) represents the establishment : counter-establishment probability ratio, which captures the relative tendencies to invade. (E) is the ratio of expected sojourn times conditional on establishment, which captures the relative tendency of each type to evade extinction over time. (F) is the ratio of average abundances throughout the sojourn. Our proposed fitness operationalization (G), the ratio of time-integrals from introduction to extinction, is negligibly different than the product of its components (D) – (E).



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542 **Fig. 4:** Causal diagram of the key components/factors underlying operationalizations of fitness. All
 543 arrows imply causality. Evolution by natural selection involves feedback between genes, environment,
 544 organismal phenotypes, and vital rates. Short-term fitness operationalizations (e.g. traditional
 545 assigned relative or absolute fitness, or derived Malthusian parameter) are summaries of current vital
 546 rates, while long-term, derived fitness operationalizations reflect lineage fate within more complete
 547 feedback systems. Both are illustrated here as shadows, indicating projections in a mathematical
 548 sense. Short-term fitness reflects instantaneous vital rates, while long-term fitness reflects longer-term
 549 projections of the fate of genetic lineages. The environment experienced by an organism broadly
 550 includes all abiotic factors (mean physical conditions, including the effects of biotic resource depletion
 551 and ecosystem engineering) and biotic factors (direct effects of conspecific and heterospecific
 552 abundances). Births, deaths, and organismal growth all feed back to the environment, because
 553 population density and its consequences are important aspects of the environment. Note that all three
 554 vital rates feed back into all three aspects of the environment, as do phenotypes. Genotypes and the
 555 informational environment (i.e. interpretable cues that organisms plastically respond to, via phenotypic
 556 plasticity and epigenetics) give rise to the strategies used by organisms. Strategies consist of
 557 investment allocations subject to life history trade-offs such as Grime's CSR triangle (Grime 1977),
 558 the competition-colonization trade-off (Tilman 1994), and bet-hedging. Phenotypes emerge from
 559 strategies deployed within a developmental environment. Niche construction and migration
 560 phenotypes affect the environment, or which environment is experienced, respectively. Selection on
 561 phenotypes gives rise to differences in vital rates. While the authors differ in their metaphysical
 562 interpretations of this figure (i.e., whether the objects shown in 3D are in fact appropriately depicted
 563 as "real" objects with fitness as a mere shadow, or whether the objects shown in 3D are rather
 564 themselves shadow-like, imperfect measures of fitness as a "real" property), what the figure shows
 565 regarding various considerations for operationalizing fitness and the relationships among alternative
 566 operationalizations is compatible with either metaphysical picture (Pence & Ramsey 2013; Walsh et
 567 al. 2017).

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