Evolutionary branching under slow directional evolution

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HIGHLIGHTS

- We derive conditions for evolutionary branching in directionally evolving populations.
- The derived conditions extend those for univariate trait spaces to bivariate trait spaces.
- Numerical analyses demonstrate their robustness.
- Our conditions are further extended to multivariate trait spaces.

ARTICLE INFO

Article history:
Received 9 December 2011
Received in revised form 20 August 2013
Accepted 24 August 2013
Available online 5 September 2013

Keywords:
Frequency-dependent selection
Speciation
Adaptive dynamics
Two-dimensional traits
Multi-dimensional traits

ABSTRACT

Evolutionary branching is the process by which ecological interactions induce evolutionary diversification. In asexual populations with sufficiently rare mutations, evolutionary branching occurs through trait-substitution sequences caused by the sequential invasion of successful mutants. A necessary and sufficient condition for evolutionary branching of univariate traits is the existence of a convergence stable trait value at which selection is locally disruptive. Real populations, however, undergo simultaneous evolution in multiple traits. Here we extend conditions for evolutionary branching to bivariate trait spaces in which the response to disruptive selection on one trait can be suppressed by directional selection on another trait. To obtain analytical results, we study trait-substitution sequences formed by invasions that possess maximum likelihood. By deriving a sufficient condition for evolutionary branching of bivariate traits along such maximum-likelihood-invasion paths (MLIPs), we demonstrate the existence of a threshold ratio specifying how much disruptive selection in one trait direction is needed to overcome the obstruction of evolutionary branching caused by directional selection in the other trait direction. Generalizing this finding, we show that evolutionary branching of bivariate traits can occur along evolutionary-branching lines on which residual directional selection is sufficiently weak. We then present numerical analyses showing that our generalized condition for evolutionary branching is a good indicator of branching likelihood even when trait-substitution sequences do not follow MLIPs and when mutations are not rare. Finally, we extend the derived conditions for evolutionary branching to multivariate trait spaces.

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1. Introduction

Real populations have undergone evolution in many quantitative traits. Even when such populations experience contemporary selection pressures, their selection response will usually be highly multivariate. However, not all responding adaptive traits evolve at the same speed: in nature, such evolutionary speeds exhibit a large variation (Hendry and Kinnison, 1999; Kinnison and Hendry, 2001). Past speciation processes may have been driven mainly by traits undergoing fast evolution (Schluter, 1996), while gradual evolutionary differentiation among species, genera, and families may derive from traits undergoing slow evolution. These differences in evolutionary speed can have two fundamentally different causes. First, they may be due to less genetic variation being available for evolution to act on: in asexual populations this occurs when mutation rates and/or magnitudes are smaller in some traits than in others, while in sexual populations this occurs when standing genetic variation is smaller in some traits than in others. Second, differences in evolutionary speed are also expected when fitness is much less sensitive to changes in some traits than to changes in others.

For brevity we refer to the slowly evolving and the rapidly evolving traits as slow traits and fast traits, respectively. If the slow traits are sufficiently slow, it is tempting to neglect their effects on the evolution of the fast traits. As far as evolutionary responses to directional selection are concerned, this simplification is usually

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0022-5193/$ - see front matter © 2013 Elsevier Ltd. All rights reserved.
http://dx.doi.org/10.1016/j.jtbi.2013.08.028
unproblematic: the directional trend of evolution (Rice et al., 2011) resulting from such selection, which we can refer to as directional evolution, is described effectively by ordinary differential equations or difference equations focusing only on those fast traits (Price, 1970; Lande, 1979; Dieckmann and Law, 1996; Rice et al., 2011).

On the other hand, such a simplification may not be safe where more complex evolutionary dynamics are involved. A typical example is adaptive speciation, i.e., evolutionary diversification driven by ecological interactions (Dieckmann et al., 2004; Rundle and Nosil, 2004). Ito and Dieckmann (2007) have shown numerically that when populations undergo disruptive selection in a fast trait, their evolutionary diversification can be suppressed by directional evolution of another trait, even if the latter is slow. Conversely, if the slow directional evolution is sufficiently slow, disruptive selection in the fast trait can drive evolutionary diversification, both in asexual populations and in sexual populations (Ito and Dieckmann, 2007). The suppression of evolutionary diversification can occur even when the slow and fast traits are mutationally and ecologically independent of each other. Thus, in a multivariate trait space, evolutionary diversification in one trait can be suppressed by slow directional evolution in just one of the many other traits. Moreover, such slow directional evolution may never cease, as the environments of populations are always changing, at least slowly, due to changes in abiotic components (e.g., climatic change) or biotic components (i.e., evolution in other species of the considered biological community). It is therefore important to improve the theoretical understanding of this phenomenon by deriving conditions for evolutionary diversification under slow directional evolution.

As a starting point to this end, we consider the special situation in which there is only a single fast trait, while all other traits of the considered population are evolving extremely slowly, such that they are completely negligible. In this case, the question whether the selection on the fast trait favors its evolutionary diversification can be examined through conditions that have been derived for the evolutionary branching of univariate traits (Metz et al., 1992; Geritz et al., 1997, 1998). In general, evolutionary branching is the process through which a unimodal phenotype distribution of a population becomes bimodal in response to frequency-dependent disruptive selection (Metz et al., 1992; Geritz et al., 1997, 1998; Dieckmann et al., 2004), which can occur through all fundamental types of ecological interaction, including competition, exploitation, mutualism, and cooperation (Doebeli and Dieckmann, 2000; Doebeli et al., 2004). This kind of diversifying evolution provides ecological underpinning for the sympatric or parapatric speciation of sexual populations (e.g., Doebeli, 1996; Dieckmann and Doebeli, 1999; Kisdi and Geritz, 1999; Doebeli and Dieckmann, 2003; Dieckmann et al., 2004; Claessen et al., 2008; Durinx and Van Dooren, 2009; Heinz et al., 2009; Payne et al., 2011). Moreover, evolutionary branching may lead to selection pressures that favor further evolutionary branching, inducing recurrent adaptive radiations and extinctions (e.g., Ito and Dieckmann, 2007), and thus community assembly (e.g., Jensen and Mulder, 1999; Bonsall et al., 2004; Johansson and Dieckmann, 2009; Brännström et al., 2012) and food-web formation (e.g., Loeuille and Loreau, 2005; Ito et al., 2009; Brännström et al., 2010; Takahashi et al., 2014). Therefore, evolutionary branching may be one of the important mechanisms underlying the evolutionary diversification of biological communities.

Conditions for the evolutionary branching of univariate traits can be extended to bivariate trait spaces, if all traits considered evolve at comparable speeds (Bolnick and Doebeli, 2003; Vukics et al., 2003; Ackermann and Doebeli, 2004; Van Dooren et al., 2004; Egas et al., 2005; Leimar, 2005; Van Dooren, 2006; Ito and Shimada, 2007; Ravnigé et al., 2009). However, if their evolutionary speeds are significantly different, the resultant conditions for bivariate traits can fail to predict evolutionary branching observed in numerical analyses (Ito and Dieckmann, 2007, 2012; Ito et al., 2009). In the present study, we therefore derive conditions for a population’s evolutionary branching in a fast trait when, at the same time, such a population is directionally evolving in one or more slow traits. The resultant conditions reveal when slow directional evolution either prevents or permits evolutionary branching.

This paper is structured as follows. Section 2 explains heuristically how the likelihood of evolutionary branching in asexual populations depends on selection pressures and mutational step sizes. Section 3 derives a normal form for strong disruptive selection and weak directional selection in a bivariate trait space and explains when arbitrary bivariate fitness functions can be mapped onto this normal form. Section 4 introduces the concept of maximum-likelihood-invasion paths, formed by mutants with maximum likelihood of invasion. On that basis, Section 5 derives sufficient conditions for evolutionary branching. Section 6 numerically examines the robustness of these conditions when the simplifying assumptions underlying our derivation are relaxed. Section 7 summarizes all conditions needed for identifying evolutionary-branching lines and extends these conditions to multivariate trait spaces. Section 8 discusses how our results generalize previously derived conditions for evolutionary branching that ignored slow directional evolution, and how our maximum-likelihood-invasion paths are related to existing methods for determining evolutionary dynamics or reconstructing evolutionary histories.

2. Heuristics

We start by describing, in a heuristic way, how disruptive selection in one direction, directional selection in another direction, and mutational step sizes may affect the likelihood of evolutionary branching. We then explain the analyses required for deriving the conditions for evolutionary branching, which are conducted in the subsequent sections.

When a population undergoes disruptive selection in trait $x$, as well as directional selection in trait $y$, its fitness landscape resembles that illustrated in Fig. 1a. The strength of disruptive selection in $x$ is given by the fitness landscape’s curvature (i.e., second derivative) along $x$, denoted by $D_{xx}$, while the strength of directional selection in $y$ is given by the fitness landscape’s slope (i.e., first derivative) along $y$, denoted by $G_y$. For simplicity, we assume that the population is monomorphic with a resident phenotype $(x,y)$, indicated by a small black circle in Fig. 1a, and that mutational step sizes are identical in all directions. In this case, possible mutants are located on a circle around the resident phenotype, as shown in Fig. 1b–g. Then, small $G_y$ means slow evolution in $y$. Roughly speaking, the direction of evolution favored by selection is indicated by the mutants possessing maximum fitness (small white circles in Fig. 1b–g). These mutants are located where the circle of considered mutants is tangential to the fitness contours.

From this simple setting, we can already draw the following geometrically evident conclusions. If $G_y$ is large compared to $D_{xx}$, which results in low curvatures for the fitness contours (Fig. 1b), the mutant having the maximum $y$ has maximum fitness, in which case directional evolution along $y$ is expected. On the other hand, if $G_y$ is sufficiently small compared to $D_{xx}$ (Fig. 1c and d), the high curvatures mean that two different mutants are sharing the same maximum fitness. In this case, evolutionary diversification in $x$ may be expected. In addition, we can easily see (Fig. 1e–g) that the smaller the mutational step size, the smaller the $G_y$ and/or larger the $D_{xx}$ required for two different mutants jointly having maximum fitness (Fig. 1e–g).
It turns out that these qualitative and heuristic insights can be corroborated by formal analysis (Sections 3–6). For this, two things have to be done properly. First, we have to clarify the conditions under which a population undergoes disruptive selection in one direction and sufficiently weak directional selection in the other direction. To compare the strengths of selection among different directions, trait spaces have to be normalized so that mutation becomes isotropic in all directions, as in Fig. 1b–g. Second, because the existence of disruptive selection is a necessary but not a sufficient condition for evolutionary branching (Metz et al., 1996; Geritz et al., 1997, 1998), the emergence of an initial dimorphism and the subsequent process of divergent evolution have to be analyzed. Conducting these analyses in the subsequent sections, we end up being able quantitatively to predict the likelihood of evolutionary branching in terms of $G_Y$, $D_{xx}$, and mutational step sizes.

3. Normal form for bivariate invasion-fitness functions causing slow directional evolution

In this section, we first derive a normal form that applies when evolution is slow in one direction. As mentioned before, this may occur when mutational steps or fitness sensitivities are strongly asymmetrical. Second, we explain the evolutionary dynamics that are expected under this normal form. Third, we outline the fundamental ideas underlying our subsequent analyses.

3.1. Bivariate invasion-fitness functions causing slow directional evolution

We start by considering arbitrary bivariate trait spaces; accordingly, each phenotype $S = (X, Y)^T$ comprises two scalar traits $X$ and $Y$. We assume an asexual population with a large population size and sufficiently small mutation rates. The latter assumption has two consequences. First, the population dynamics have sufficient time to relax toward their equilibrium after a new mutant emerges. Second, as long as the population experiences directional selection, only the phenotype with the highest fitness among the existing phenotypes survives as a result of selection. Thus, the population is essentially always close to equilibrium and monomorphic. This allows its directional evolution to be translated into a trait-substitution sequence based on the invasion fitness of a mutant phenotype $s'$ arising from a resident phenotype $S$ (Metz et al., 1992, 1996; Dieckmann and Law, 1996).

The invasion fitness of $s'$ under $S$, denoted by $F(s'; S)$, is defined as the initial per capita growth rate of $s'$ in the monomorphic population of $S$ at its equilibrium population size. The function $F(s'; S)$ can be treated as a fitness landscape in $s'$, whose shape depends on $S$. When a mutant emerges, which occurs with probability $\mu$ per birth, we assume that its phenotype follows a fitness function which can be expanded in two dimensions, $V_{XX}$ and $V_{YY}$ (Fig. 2b). Then, stretching the trait space in the Y-direction by $\sigma_Y$ gives an isotropic mutation distribution with standard deviation $\sigma$, where $\sigma_X = \sqrt{V_{XX}}$ and $\sigma_Y = \sqrt{V_{YY}}$ (Fig. 2b). Then, $\Lambda = M(\delta S)S\delta S^T d\delta X d\delta Y$, where $\delta S = s' - S$. The distribution is assumed to be symmetric, unimodal, and smooth. As long as the mutational step sizes are sufficiently small, such that $M(\delta S)$ has sufficiently narrow width, the distribution is well characterized by its variance-covariance matrix $\Lambda$.

\[
\Lambda = \int M(\delta S)S\delta S^T d\delta X d\delta Y = \begin{pmatrix} V_{XX} & V_{XY} \\ V_{XY} & V_{YY} \end{pmatrix},
\]

where $V_{XX} = \int \delta X^2 M(\delta S)d\delta X d\delta Y$, $V_{YY} = \int \delta Y^2 M(\delta S)d\delta X d\delta Y$, and $V_{XY} = \int \delta X \delta Y M(\delta S)d\delta X d\delta Y$. The standard deviation of mutational step sizes along each direction is thus described by an ellipse, $\delta S^T \Lambda^{-1} \delta S = 1$, as shown in Fig. 2a. Since $\Lambda$ is symmetric, its two eigenvectors are orthogonal, and these vectors determine the directions of the long and short principal axes of this ellipse. Through a coordinate rotation, we align the axes of the coordinate system with the eigenvectors of $\Lambda$. In the rotated coordinate system, $\Lambda$ is diagonal, $V_{XY} = 0$, and we can choose the axes such that $\sigma_X > \sigma_Y$, where $\sigma_X = \sqrt{V_{XX}}$ and $\sigma_Y = \sqrt{V_{YY}}$ (Fig. 2b). Then, the trait space in the Y-direction by $\sigma_Y$ gives an isotropic mutation distribution with standard deviation $\sigma = \sigma_X$ in all directions (Fig. 2c). This is achieved by introducing a new coordinate system $s = (x, y)^T$, where $x = X$ and $y = (\sigma_X/\sigma_Y)Y$. We assume that $\sigma$ is small, such that $\sigma < 1$. In this normalized trait space, the invasion-fitness function can be expanded in $s$ and $s$ around a base point $s_0 = (x_0, y_0)^T$ as

\[
f(s'; s) = G(s' - s_0)^T c + \frac{1}{2} s'(D_0 s) + O(\sigma^2),
\]
where \( \delta s = s' - s = O(\sigma) \) and \( s - s_0 = O(\sigma) \) are assumed. The \( G, C, \) and \( D \) are given by

\[
G = (G_x, G_y) = f_m, \quad C = \begin{pmatrix} C_{xx} & C_{xy} \\ C_{yx} & C_{yy} \end{pmatrix} = f_{mm} + f_{rm},
\]

\[
D = \begin{pmatrix} D_{xx} & D_{xy} \\ D_{yx} & D_{yy} \end{pmatrix} = f_{mm},
\]

\[
f_m = (f_x f_y) \text{ for } s = s_0, \quad f_{mm} = (f_{xx} f_{xy}) \text{ for } s = s_0,
\]

\[
f_{rr} = (f_{xx} f_{yy}) \text{ for } s = s_0.
\]

where the subscripts ‘m’ and ‘r’ refer to mutants and residents, respectively, and where \( f_m \) for \( \alpha = x, y \) and \( f_{mr} \) for \( \alpha, \beta = x, y \) denote the first and second derivatives of \( f(s'; s) \), respectively. See Appendix A for the derivation of Eq. (2). Note that \( G, C, \) and \( D \) are functions of the base point \( s_0 \). The vector \( G \) describes the fitness gradient at \( s \) when \( s = s_0 \). The matrix \( C \) describes how the fitness gradient at \( s \) changes as \( s \) deviates from \( s_0 \). The matrix \( D \) describes the curvature (i.e., second derivative) of the fitness landscape, which is approximately constant as long as the third-order terms can be neglected.

If \( \sigma_y \) is much smaller than \( \sigma_x \), such that \( \sigma_y = O(\sigma_x^2) \), the stretching of the trait space in the \( Y \)-direction for the normalization makes derivatives with respect to \( y \) very small, in the sense \( G_y = O(\sigma), \ C_{xy} = O(\sigma), \ C_{yy} = O(\sigma), \ D_{xy} = O(\sigma), \ C_{yy} = O(\sigma^2) \), and \( D_{yy} = O(\sigma^2) \). This simplifies Eq. (2a),

\[
f'(s'; s) = G_x \delta x + C_{xy}(x-x_0) \delta x + \frac{1}{2} D_{xx} \delta x^2 + G_y \delta y + O(\sigma^3),
\]

where terms with \( C_{yy}, C_{xy}, D_{xx}, D_{xy}, \) and \( D_{yy} \) are subsumed in the higher-order terms \( O(\sigma^3) \). Notice that on the right-hand side only the first term is of order \( O(\sigma) \), while the other terms, including \( G_y \delta y \), are of order \( O(\sigma^2) \).

This means that this normal form describes fitness functions with significantly weak directional selection along \( y \) compared to that along \( x \) as long as \( G_x \) and \( G_y \) in the original trait space have similar magnitudes. We call Eq. (3) the normal form for invasion-fitness functions with significant sensitivity difference.

A comparison of Eqs. (2a) and (3) shows that the latter can be obtained even for \( \sigma_y = \sigma_x \), provided the sensitivity of the fitness function to variation in trait \( Y \) is significantly weaker than that in \( X \), so that \( G_y, C_{xy}, C_{yy}, D_{xy}, \) and \( D_{yy} \) are all relatively small, satisfying

\[
|G_y| + |C_{xy}| + |C_{yy}| + |D_{xy}| + |D_{yy}| = O(\sigma).
\]

For a derivation of Eq. (4), see Appendix B.

Notice that the assumption needed for the derivation of Eq. (3), i.e., \( \sigma_y = O(\sigma_x^2) \), also satisfies Eq. (4). Thus, the condition for a significant difference between mutational step sizes in the original trait space \( S \) can naturally be integrated with the condition for a significant sensitivity difference of the invasion-fitness function in the normalized trait space \( s \). Based on Eq. (4), we therefore define the condition for significant sensitivity difference as follows.

**Significant sensitivity difference.** After normalization to make mutation isotropic, the invasion-fitness function can be made to satisfy Eq. (4) by rotating the \( x- \) and \( y- \) axes.

### 3.2. Evolutionary dynamics expected under normal form

We now consider the expected evolutionary dynamics induced by the normal form in Eq. (3). For this purpose we first recap expectations for the simpler case in which \( G_y \) is so small that \( G_y = O(\sigma^2) \). In that case, \( G_y \delta y = O(\sigma^3) \) is negligible, so that \( y \) vanishes from Eq. (3), and the evolutionary dynamics therefore become univariate in \( x \), so that phenotypes are characterized by that trait value alone. In this simpler case, conditions for evolutionary branching are easier to understand (Metz et al., 1996; Geritz et al., 1997, 1998), as follows.

Suppose that the base point \( x_0 \) of the expansion of \( f(s'; s) \) can be chosen such that \( G_{xx} = 0 \). Such a point, which we denote by \( x_0 \), is called an evolutionarily singular point (or simply a singular point or an evolutionary singularity) because a resident located at \( x_0 \) experiences no directional selection. In contrast, a resident located close to \( x_0 \) experiences directional selection along \( x \).

\[
\frac{df(s'; s)}{dx} \bigg|_{x=x_0} = C_{xx}(x-x_0).
\]

If \( C_{xx} < 0 \), the fitness gradient is positive for \( x < x_0 \), and negative for \( x > x_0 \), which means that it attracts a monomorphic population through directional evolution. In other words, the singular point is then convergence stable (Christiansen, 1991).

When a population comes close to \( x_0 \), it may become possible for a mutant \( s' \) to coexist with a resident \( s \). Mutual invasibility between \( s' \) and \( s \), which gives rise to protected dimorphism (Prout, 1968), is defined by \( f(s'; s) > 0 \) and \( f(s; s') > 0 \), which requires \( D_{xx} = C_{xx} > 0 \). Following the emergence of a protected dimorphism of trait values denoted by \( s_1 \) and \( s_2 \), the resultant fitness landscape \( f(s'; s_1, s_2) \) maintains approximately the same curvature (i.e., second derivative) \( D_{xx} \) along \( x \). If this curvature is positive, i.e., this point is not a local evolutionarily stable strategy, or ESS (Maynard Smith and Price, 1973), the two subpopulations evolve in opposite directions, keeping their coexistence, in a process called dimorphic divergence. When dimorphic divergence occurs in univariate trait spaces, it can never collapse (if it is assumed that mutual invasibility among phenotypes ensures their coexistence). This is because only mutants outside of the interval between the two residents can invade, and such an invading mutant then always excludes the closer resident and is mutually invasible with the other more distant resident, resulting in a new protected dimorphism with a larger phenotypic distance.

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**Fig. 2.** Coordinate transformations for normalizing the mutation probability distribution. The first transformation, (a) to (b), is a rotation, while the others cause scaling.
(Geritz et al., 1998). As we will see below, such collapses, however, become crucial when analyzing dimorphic divergence in bivariate trait spaces.

The evolutionary process described above is called evolutionary branching. It requires monomorphic convergence ($C_{xx} < 0$), mutual invasibility ($D_{xx} - C_{xx} > 0$), and dimorphic divergence ($D_{xx} > 0$). We therefore see that the necessary and sufficient conditions for univariate evolutionary branching are given by the existence of a point $x = x_0$ satisfying

$$G_x = 0, \qquad (6a)$$

$$C_{xx} < 0, \qquad (6b)$$

$$D_{xx} > 0, \qquad (6c)$$

with $x_0 - x_0$. When trait $y$ is also taken into account, the point $x = x_0$ forms a line in the bivariate trait space. Thus, the aforementioned conditions for univariate evolutionary branching can be translated into the following statement.

**Bivariate translation of conditions for univariate evolutionary branching.** When directional selection in $y$ is very weak, such that $G_y = O(\sigma^2)$, monomorphic populations around a line $x = x_0$ converge to that line and bring about evolutionary branching, if and only if Eqs. (6) are all satisfied.

Using these simple results as a baseline for comparison, we now consider the case that $G_y$ is of order $\sigma^1$, which, according to Eq. (3), implies that the evolution in $y$ can affect the evolution in $x$. When the population is not close to the singular line $x = x_0$, directional evolution in $x$ dominates the effects of $y$. Thus, the singular line still attracts monomorphic populations if and only if $C_{xx} < 0$. We thus call such a singular line a convergence-stable line. When a population is in the neighborhood of a convergence-stable line and evolves toward it, directional selection in $x$ inevitably becomes small, such that $G_y$ may affect the evolutionary dynamics.

When $x$ is close to $x_0$, a dimorphism in $x$ may emerge, but invasion by a mutant in $y$ with higher fitness may exclude both of the coexisting resident phenotypes and thereby abort the incipient evolutionary branching. Such an abortion is especially likely when $G_y$ is large. Thus, the larger $G_y$ becomes, the more difficult evolutionary branching is expected to be. Below, we examine how the resultant likelihood of evolutionary branching can be estimated.

### 3.3. Motivation for further analyses

In principle, bivariate evolutionary branching is possible even for very large $G_y$, as long as trait-substitution sequences comprise invasions only in $x$ for an adequately large number of substitutions after the inception of dimorphism. However, for large $G_y$, sequential invasions of this kind are unlikely, because the fitness advantage of mutants in $y$ then is large, which favors their invasion, which in turn easily destroys any initial dimorphisms. Thus, the average number of invasions required for evolutionary branching is expected to be quite large in this case. We can thus measure the likelihood of evolutionary branching as the probability of its successful completion within a given number of invasions.

It is difficult to calculate this probability directly, and thus to determine its dependence on the parameters $G_y$, $C_{xx}$, and $D_{xx}$ of the normal form. To avoid this difficulty, we focus on invasions that individually have maximum likelihood for a given composition of residents. We can loosely interpret the successions of residents formed by such invasions as describing typical evolutionary paths. Because of their special construction, it is possible analytically to derive sufficient conditions for evolutionary branching along these paths. It is expected that the conditions thus obtained can serve as useful indicators for the probability of evolutionary branching along the more general evolutionary paths formed by arbitrary stochastic invasions.

Notice that when we refer to stochastic invasions, we refer to the stochasticity of mutations and to the stochasticity of the initial survival of rare mutants, but not to the effects resulting from small resident population size, which occasionally allow mutants to invade even when they have negative fitness. In formal terms, these clarifications are implied by our assumption of sufficiently large resident population size.

In our analyses below, we assume that the conditions for evolutionary branching in univariate trait spaces, Eqs. (6), are satisfied. Our goal is to determine how conditions for bivariate evolutionary branching in $x$ under weak directional selection in $y$ differ from Eqs. (6).

### 4. Maximum-likelihood-invasion paths

In this section, we define evolutionary paths formed by sequential invasions each of which has maximum likelihood. Among all possible evolutionary paths formed by arbitrary stochastic invasions, these paths have high likelihood and may therefore be regarded as typical. Our reason for introducing these maximum-likelihood-invasion paths (MLIPs) is that we can derive, in Section 5, conditions for evolutionary branching along those typical paths.

#### 4.1. Definition of oligomorphic stochastic invasion paths

We start by explaining how probabilities of invasion events are formally defined. We consider a monomorphic population with phenotype $s$, as a trait vector with an arbitrary dimension, at equilibrium population size $n$ that is uniquely determined by $s$. The birth and death rates (i.e., the number of birth and death events per individual per unit time, respectively) of a rare mutant phenotype $s'$ are denoted by $b(s' ; s)$ and $d(s' ; s)$, where $b(s, s)$ and $d(s, s)$ denote the birth and death rates of the resident $s$, which must satisfy $f(s, s) = b(s, s) - d(s, s) = 0$ because the resident is at population dynamical equilibrium. The invasion fitness of the mutant $s'$ in the environment determined by the resident $s$ is given by

$$f(s, s') = b(s, s') - d(s', s).$$

(7)

Once a mutant $s'$ has arisen, the probability of its successful invasion in a population of resident $s$ is approximately given by $f(s', s)/b(s, s)$ (Dieckmann and Law, 1996). Here, the subscript "+" denotes conversion of negative values to zero. The probability density per unit time for the emergence of a successfully invading mutant $s'$ in a population of residents $s$ is given by multiplying the density $\mu b(s, s) M(s' - s)$ of mutants emerging per unit time with their probability of successful invasion,

$$E(s', s) = \mu b(s, s) M(s' - s) f(s', s)/b(s, s),$$

(8a)

where $\mu$ is the mutation probability per birth event, and $M(s' - s)$ is the mutation probability distribution. The above approximation applies in the leading order of $s' - s$, when $\sigma$ is sufficiently small such that $b(s'; s) - b(s, s)$ is much smaller than $b(s, s)$ (i.e., $[b(s'; s) - b(s, s)]/b(s, s) = O(\sigma^2)$) (see Appendix C for the derivation). The expected waiting time for the next invasion event is given by $T = 1 / \int E(s', s) ds'$. When an invasion event occurs, the successfully invading mutants $s'$ follow the invasion-event probability density

$$P(s', s) = TE(s', s).$$

(8b)

For a polymorphism with resident phenotypes $s_0 = (s_1, \ldots, s_V)$, the probability density (per unit time) of successful invasion by a mutant phenotype $s'$ originating from the resident $s_0$ is
given by an approximation analogous to the monomorphic case, Eq. (8a),
\[ E_i(s'; s_1, \ldots, s_k) = \hat{n}_i M(s' - s_i)f(s'; s_1, \ldots, s_N), \]
where \( \hat{n}_i \) is the equilibrium population size of the resident \( s_i \). As in Eq. (8b), we can also define an invasion-event probability density
\[ P_i(s'; s_1, \ldots, s_N) = T E_i(s'; s_1, \ldots, s_N), \]
where \( T = 1/\sum_{i=1}^{N} E_i(s'; s_1, \ldots, s_N)ds' \) is the expected waiting time for the next invasion event. Notice that the invasion event is identified by the combination (\( s' \), i) of the mutant phenotype \( s' \) and its parental resident \( s_i \). Consequently, the invasion event probability densities are normalized according to \( \int \sum_{i=1}^{N} P_i(s'; s_1, \ldots, s_N)ds' = 1 \). When only a single resident exists, Eqs. (9a) and (9b) are identical to Eqs. (8a) and (8b), respectively.

The invasion by a mutant leads the community to a new population dynamical equilibrium. In most cases, the mutant replaces only its parental resident, while under certain conditions the coexistence of both, extinction of both, or extinction of other residents may occur. A sequence of such invasions specifies a succession dynamics of resident phenotypes, which is called a trait-substitution sequence (Metz et al., 1996).

If the invasion event is calculated stochastically according to Eqs. (8b) and (9b), the resultant trait substitution is called an oligomorphic stochastic process (Ito and Dieckmann, 2007). When considering an initial monomorphic resident \( s_i \) and a mutant-invasion sequence
\[ I = (s'(1), \ldots, s'(k), \ldots, s'(K)), \]
where \( s'(k) \) is the mutant that invades in the kth invasion event, the trait-substitution sequence is denoted by
\[ \text{RI}(I, s_i) = \left( s_i(1), \ldots, s_i(k), \ldots, s_i(K) \right), \]
where \( s_i(k) = (s_i(k), \ldots, s_{N_i}(k)) \) is an \( (N-k) \)-dimensional vector composed of the \( N-k \) resident phenotypes that coexist after the invasion of \( s'(k) \). This kind of trait-substitution sequence constitutes an evolutionary path, which we call an oligomorphic stochastic invasion path (OSIP).

If the kth invasion event leads to the extinction of the entire community, no further invasions occur. In this case, the lengths of \( I \) and \( \text{RI}(I, s_i) \) are limited by \( k \). In this study, we condition all analyses on the absence of complete community extinction.

4.2. Definition of maximum-likelihood-invasion paths

We now introduce the concept of maximum-likelihood invasion. Specifically, we define a maximum-likelihood-invasion event as the combination of the mutant \( s_{\text{MLI}} \) and its parental resident \( s_i \) with \( i = i_{\text{MLI}} \) that maximizes the invasion-event probability density, Eq. (9b),
\[ \arg \max_{s', s_i} P_i(s'; s_1, \ldots, s_N), \]
where \( s_{\text{MII}} \) and \( s_{\text{MLI}} \) are the MLI mutant and MLI resident, respectively, and denote \( s_{\text{MLI}} \) by \( s_{\text{MII}} \) for convenience. A maximum-likelihood-invasion path (MLIP) is a trait-substitution sequence formed by MLI events, denoted by \( \text{MLI} = \left( s_{\text{MLI}}(0), \ldots, s_{\text{MLI}}(k), \ldots, s_{\text{MLI}}(K) \right) \). The MLIP, which is expressed as \( \text{RI}(I_{\text{MLI}}, s_i) \) for an initial monomorphic resident \( s_i \), is included in the set of all corresponding possible OSIPs \( \text{RI}(I, s_i) \).

Note that the MLI mutational steps \( s_{\text{MII}} - s_{\text{MLI}} \) are bounded by \( \sqrt{2} \sigma \), if invasion-fitness functions are approximated by quadratic forms of \( s' \) (e.g., Eqs. (2)) and if mutation probability distributions are approximated by multivariate Gaussian functions (Appendix F).

Also note that the MLIP does not give the maximum-likelihood OSIP, which would require maximization of the likelihood at the level of the mutant-invasion sequence rather than at the level of individual mutant-invasion events. Although such sequence-level maximization would be more appropriate for our purpose, it seems analytically intractable. On the other hand, the event-level maximization defined by MLIPs is analytically tractable, and the MLIP is still expected to have a relatively large likelihood among corresponding OSIPs. Likewise, as illustrated by our numerical results in Section 6, when an MLIP \( \text{RI}(I_{\text{MLI}}, s_i) \) exhibits evolutionary branching, then a large fraction of the corresponding OSIPs \( \text{RI}(I, s_i) \) also exhibit evolutionary branching.

5. Conditions for evolutionary branching along MLIPs

In this section, we derive sufficient conditions for evolutionary branching along MLIPs, in terms of the properties of the normal form for invasion-fitness functions with significant sensitivity difference, Eq. (3).

5.1. Further rescaling

Here we assume that the base point of expansion \( s_0 = (x_0, y_0)^T \) is on a convergence-stable line \( x = x_b \) that satisfies univariate conditions for evolutionary branching, Eqs. (6). To simplify the analysis, we adjust the trait space as follows, without loss of generality. First, we shift the origin of the trait space to the base point \( s_0 \) so that \( s_0 = (0, 0)^T \) and \( y_0 = 0 \). Second, we rescale the trait space so that \( \sigma = 1 \). (In this case, magnitude differences among \( |d_s| = O(\sigma), \) \( |d_s|^2 = O(\sigma^2) \) with \( \sigma < 1 \) are transformed into those among the corresponding derivative coefficients \( G, C, \) and \( D \), while the magnitudes of \( |d_s|, |d_s|^2 \) themselves become similar to each other.) Third, we rescale time and potentially flip the direction of the y-axis so that \( C_0 = 1 \). For simplicity, we consider the first- and second-order terms only. Consequently, \( f(s; s') \) is given by
\[ f(s; s') = \delta y + D \delta x^2 + C \delta x, \]
with \( \sigma \) before the rescaling (i.e., \( \sigma < 1 \)). In the simplified normal form in Eq. (12a), only two dimensionless parameters \( D \) and \( C \) determine the geometry of the fitness landscape. This geometry not only determines the fitness landscape’s shape (\( D \)), but also how the landscape changes when the resident phenotype is varied (\( C \)). Eq. (12a) then shows that any possible fitness landscape \( f(s; s') \) can be obtained from \( f(s'; (0, 0)^T) = \delta y + D \delta x^2 \) by a parallel shift, i.e.,
\[ f(s; s') = f(s' - s_0; (0, 0)^T) \]
with \( s_0 = (x_b, y_0)^T \). This means that the contour curve \( f(s; s') = 0 \) given by \( y = -Dx^2 - x - y_0 \) always has a constant parabolic shape specified by \( D \), so that the position of this curve determines the fitness landscape (Fig. 3a).

In the next two subsections, we derive conditions on \( D \) and \( C \) for evolutionary branching along MLIPs. We first obtain conditions on MLI mutants, \( s_{\text{MLI}} \), for evolutionary branching. Then we analyze these conditions considering the dependence of \( s_{\text{MLI}} \) on \( D \) and \( C \), which provides conditions on these two parameters for evolutionary branching.

5.2. Conditions on MLI mutants for evolutionary branching

Here we obtain conditions on \( s_{\text{MLI}} \) for evolutionary branching. The process of evolutionary branching can be decomposed into two
Conditions for dimorphic emergence

The dark gray and white regions touch each other only when the MLIP condition (e) and (f) are satisfied, respectively. The thin parabolic curves giving the boundaries share the same shape. In panels (c) and (d), the white regions indicate mutants that satisfy the conditions for dimorphic emergence and divergence, respectively. The thick gray curves in (a) and (b) indicate the zero-contour of the invasion. Clearly, if conditions for dimorphic emergence and for dimorphic divergence both hold, evolutionary branching along MLIPs inevitably occurs for an arbitrary initial resident.

Lemma 1. Suppose the conditions for dimorphic divergence below hold. Then for any initial protected dimorphism of $s_1$ and $s_2$ emerging under the conditions for dimorphic emergence, subsequent invasions by $s_{MLI}$ continue directional evolution of the two morphs in opposite directions in $x$ without collapse.

Conditions for dimorphic divergence. Any $s_{MLI}$ satisfies

\[ x_{MLI} < x_1 \quad \text{and} \quad |y_{MLI} - y_1| < D(x_{MLI} - x_1)^2 \]  

or

\[ x_{MLI} > x_2 \quad \text{and} \quad |y_{MLI} - y_2| < D(x_{MLI} - x_2)^2, \]

where $x_1 < x_2$ is assumed without loss of generality.

The set of mutants satisfying inequalities (14) is indicated by the white regions in Fig. 3d. In each invasion step, $s_{MLI}$ replaces only its parental resident, so the divergence of the new dimorphism in $x$ is larger than that of $s_1$ and $s_2$.

Clearly, if conditions for dimorphic emergence and for dimorphic divergence both hold, evolutionary branching along MLIPs inevitably occurs for an arbitrary initial resident $s_i$.

5.3. MLIP condition

As $s_{MLI}$ is a function of $D$ and $C$, substituting this function into the conditions for dimorphic emergence and divergence above and solving those for $D$ and $C$ gives conditions on these parameters for evolutionary branching.

To derive $s_{MLI}$ as a function of $D$ and $C$, we explicitly define the mutation distribution. For analytical tractability, we assume that the mutation distribution is approximated by a two-dimensional Gaussian distribution, which is expressed in the normalized and rescaled trait space $s = (x, y)^T$ as

\[ M(s) = \frac{1}{2\pi} \exp\left(-\frac{1}{2} |s|^2\right). \]

where the standard deviation of mutational step sizes is scaled to $1$.

Under monomorphism with phenotype $s$, the MLI mutant $s_{MLI}$, which maximizes the invasion-event probability density, is given by the $s'$ that maximizes Eq. (8b),

\[ P(s' ; s) = T \mu \delta x \exp\left(-\frac{1}{2} [\delta x]^2 + [\delta y]^2 |\delta y + D \delta x^2 + C \delta x| \right). \]

We first focus on the special case that $s$ is located exactly on the convergence-stable line, i.e., $s = (0, y)^T$ with arbitrary $y$. In this

\[ x_{MLI} > x \quad \text{for} \quad x < 0 \]

\[ x_{MLI} < x \quad \text{for} \quad x > 0, \] (13a)
case, $s_{nu}$ is given by
\[
\begin{align*}
    s_{nu} &= \begin{cases} 
        (0, 1)^T + \bar{s} & \text{for } D < \frac{1}{2} \\
        \left( \pm \frac{\sqrt{D^2-1}}{2D}, \frac{1}{2} \right)^T + \bar{s} & \text{for } D \geq \frac{1}{2}
    \end{cases}
\end{align*}
\]  

Substitution of Eq. (17) and $s = (0, y)^T$ into the inequality condition (13b) for dimorphic emergence yields
\[
D > \frac{1}{\sqrt{2}}
\]  

We do not need to examine inequalities (13a), which are of interest only for $s = (x, y)^T$ with $x \neq 0$. Even if the resident is not located on the convergence-stable line, i.e., $s = (x, y)^T$ with $x \neq 0$, inequalities (13) still hold under $D > 1/\sqrt{2}$, as shown in Appendix D. Therefore, conditions for dimorphic emergence are satisfied if $D > 1/\sqrt{2}$ holds. In this case, $s_{nu}$ is always located inside the dark gray rectangle in Fig. 3e.

Moreover, under $D > 1/\sqrt{2}$, the derivation in Appendix D shows that $s_{nu}$ always satisfies the condition for dimorphic divergence. Therefore, inequality (18), $D > 1/\sqrt{2}$, is a sufficient condition for evolutionary branching along MLIPs starting from an arbitrary initial monomorphic resident. More specifically, we have

**Theorem 1.** Suppose a normalized and rescaled invasion-fitness function having significant sensitivity difference, expressed in form of Eq. (3) with $\sigma = 1$, satisfies both $C < 0$ and inequality (18), $D > 1/\sqrt{2}$. Then any MLIP starting from an arbitrary initial monomorphic resident monotonically converges toward the convergence-stable line, $x = 0$, and brings about protected dimorphism, which leads to dimorphic divergence without collapse, i.e., evolutionary branching.

We thus call inequality (18), $D > 1/\sqrt{2}$, the MLIP condition for evolutionary branching, and refer to a convergence-stable line satisfying this condition as an *evolutionary-branching line*.

**5.4. Directional evolution sufficient for evolutionary branching**

Under the MLIP condition, dimorphism with $|x_2 - x_1| \geq \Delta x^*$ for arbitrary $\Delta x^* > 0$ emerges before the population directionally has evolved by
\[
y - y_0 = L_0(y_1, \Delta x^*) = (|x_1| + \Delta x^*)/\sqrt{2},
\]  

where the second equality defines the function $L_0(|x_1|, \Delta x^*)$, and $s_0 = (x_0, y_0)^T$ is the initial monomorphic resident (see Appendix J for the derivation). The $\bar{y}$ is the mean value of $y_j$, given by $\bar{y} = \bar{y}$ for monomorphism or by $\bar{y} = (\bar{n_1}y_1 + \bar{n_2}y_2)/(\bar{n_1} + \bar{n_2})$ for dimorphism, where $\bar{n}_1$ and $\bar{n}_2$ are the equilibrium population sizes of $s_1$ and $s_2$, respectively.

**6. Numerical examination of MLIP condition**

In this section we investigate how the MLIP condition is related to the likelihood of evolutionary branching in numerically calculated MLIPs, OSIPs, and polymorphic stochastic invasion paths (PSIPs) in which mutation rates are not small. See Appendices K, M, and L for details on algorithms and initial conditions.

When deriving the MLIP condition, we assumed the bivariate Gaussian mutation distribution defined in Eq. (15). The resultant MLIP condition may also be applicable to other types of mutation distributions. To examine this kind of robustness, below we investigate additional three different mutation distributions for the calculation of OSIPs and PSIPs. The *bivariate fixed-step* distribution has possible mutations that are bounded on a circle (Fig. 5b). The *univariate Gaussian* distribution applies when mutations in $x$ and $y$ occur separately, each following a one-dimensional Gaussian distribution (Fig. 5c). The *univariate fixed-step* distribution also limits possible mutations to affect either $x$ or $y$, but with fixed step sizes (Fig. 5d). See Appendix L for details on these mutation distributions.

The cumulative likelihood of evolutionary branching is measured as a probability $p(L_y - L_0)$, where $L_y$ is the length of directional evolution in $y$ along MLIPs, OSIPs, or PSIPs, while $L_0$, calculated through Eq. (19), is the length of directional evolution in $y$ along MLIPs sufficient for the occurrence of evolutionary branching (see Appendix K for details on $L_0$). Thus, $p(L_y - L_0)$ measures the cumulative probability of evolutionary branching when the population has directionally evolved in $y$ by $L_y$, beyond what is implied by the MLIP condition ($L_0$). In the case of MLIPs, $p(L_y - L_0) = 1$ clearly holds for $L_y - L_0 = 0$. In the case of OSIPs and PSIPs, when values of $p(L_y - L_0)$ for $L_y - L_0 = 0$ are close to 1, this indicates that the MLIP condition is working well also under such relaxed conditions. However, $p(L_y - L_0)$ never reaches 1 in OSIPs, differently from MLIPs. One reason is that even under very large $D$ there are non-zero probabilities for repeated mutant invasions only in the $y$-direction, causing directional evolution in the $y$-direction. Another reason is that even after the emergence of a protected dimorphism, this dimorphism may collapse by subsequent mutant invasions in the case of OSIPs. When a dimorphism has collapsed, leaving behind a monomorphic resident, by the definition of OSIPs, the information about the collapse itself is lost, and it is only the remaining resident that determines the likelihood of evolutionary branching in the “next trial.” A sufficiently large $D$ is expected to induce evolutionary branching within a few trials, keeping the total directional evolution in the $y$-direction short, which results in a high value of $p(L_y - L_0)$ for $L_y - L_0 = 0$.

**6.1. Sufficient vs. necessary conditions: MLIPs**

Fig. 4a shows the branching likelihood in MLIPs under the bivariate Gaussian mutation distribution for varying $C < 0$ and $D > 0$; the contour curves indicate where a 97% cumulative probability of evolutionary branching is reached for $L_y - L_0 = 0$, 100, and 200 (i.e., $p(100) = 0.97$, $p(1000) = 0.97$, and $p(2000) = 0.97$). For $D > 1/\sqrt{2}$, MLIPs quickly undergo evolutionary branching in the gray area in Fig. 4a, while they do not undergo evolutionary branching in the white area in Fig. 4a. Examples of branching and non-branching MLIPs are shown as gray curves in Fig. 4b–d, respectively. Importantly, the threshold boundary for evolutionary branching is well located in the white area in Fig. 4a. In particular, the MLIP condition $D > 1/\sqrt{2}$ seems to give a necessary and sufficient condition as $C$ converges to 0.

**6.2. Robustness of MLIP condition: OSIPs**

When the MLIP condition $D > 1/\sqrt{2}$ holds, OSIPs tend to undergo immediate evolutionary branching (black curves in Fig. 4b). On the other hand, even for $D < 1/\sqrt{2}$, OSIPs may still undergo evolutionary branching (black curves in Fig. 4c). In this case, however, the required $L_y - L_0$ becomes large as $D$ is decreased. As $D$ is decreased further, evolutionary branching may not be observed even for very large $L_y - L_0$ (black curves in Fig. 4d).

Fig. 5a shows the branching likelihood in OSIPs under the bivariate Gaussian mutation distribution for varying $C < 0$ and $D > 0$; the contour curves indicate where a 97% cumulative probability of evolutionary branching is reached for $L_y - L_0 = 0$, 100, and 200 (i.e., $p(0) = 0.97$, $p(100) = 0.97$, and $p(200) = 0.97$). We see that more than 97% branching likelihood is attained for $L_y - L_0 = 0$, as expected by the MLIP condition. Similarly, more than 90% branching likelihood is attained for $L_y - L_0 = 0$ for each of the three other mutation distributions (Fig. 5b–d), as long as the mutation rate in $y$ is not very small.
-sized equals 1 in all directions.

The stochastic effects become large when mutation distributions. Population dynamics of PSIPs are calculated based on (as for OSIPs), but by gradual changes of polymorphic phenotype.

6.3. Robustness of MLIP condition: PSIPs

For PSIPs, mutation rates need not be low. In this case, evolutionary dynamics are no longer given by trait-substitution sequences (as for OSIPs), but by gradual changes of polymorphic phenotype distributions. Population dynamics of PSIPs are calculated based on the stochastic sequence of individual births and deaths (Dieckmann and Law, 1996). The stochastic effects become large when fitness gradients and curvatures are both weak and/or population sizes are small. In this case, the likelihood of evolutionary branching in PSIPs, in contrast with OSIPs, may be affected not only by \( C \) and \( D \), but also by other parameters, such as the mutational step size \( \mu \), the mutation rate \( \mu \), and the carrying capacity along the evolutionary-branching line, \( K_0 \). We have numerically confirmed that the MLIP condition is still useful for characterizing evolutionary branching in PSIPs across a certain range of parameter values. For example, \( D > 1 / \sqrt{2} \) provides \( p(0) > 0.9 \) under all four mutation distributions for \( 0.001 \leq \sigma \leq 0.01, 300 \leq K_0 \leq 10000 \), and \( 1 \times 10^{-1} \geq \mu \geq 3.3 \times 10^{-5} \), with \( 3 \times 10^{-3} \leq \mu \sigma K_0 \leq 3 \times 10^{-2} \) (results not shown).

Fig. 5e–h show the branching likelihood in PSIPs for varying \( C \) and \( D \), with \( \sigma = 0.01, K_0 = 600 \), and \( \mu = 5.1 \times 10^{-5} \); the contour curves indicate where a 95% cumulative probability of evolutionary branching is reached for \( L_y-L_0 = 0 \), 40, and 80 (i.e., \( p(0) = 0.95 \), \( p(40) = 0.95 \), and \( p(80) = 0.95 \). We see that more than 95% branching likelihood is attained for \( L_y-L_0 = 0 \) under all four mutation distributions, as long as the mutation rate in \( y \) is not very small compared to that in \( x \) (i.e., \( \mu_y / \mu_x \geq 0.05 \) for the univariate Gaussian and univariate fixed-step mutation distributions. Thus, for the examined PSIPs, the MLIP condition turns out to be robust as a good indicator for evolutionary branching, even when mutation rates are not small and/or mutation distributions other than bivariate Gaussian are considered.

7. Conditions for evolutionary-branching lines

In this section, we first summarize the conditions for evolutionary-branching lines in bivariate trait spaces. Second, we
extend these conditions to multivariate trait spaces. Third, we explain how to find evolutionary-branching lines or manifolds in arbitrary trait spaces with arbitrary dimensionality.

7.1. Conditions for evolutionary-branching lines in bivariate trait spaces

By an appropriate affine transformation, arbitrary trait spaces \( S = (X, Y)^T \) can be normalized into a representation \( s = (x, y)^T \) with isotropic mutation with standard deviation \( \sigma \). In this normalized trait space, the invasion-fitness function can be expanded around \( s_0 \) as shown in Eq. (2a),

\[
f(s; s) = Gss + \frac{1}{2} \delta s^T D \delta s + O(\sigma^3),
\]

where \( \delta s = (\delta x, \delta y)^T = s - s_0 \). If the \( x \)-and \( y \)-axes can be adjusted such that Eq. (4) holds,

\[
|G_y| + |C_{x}y| + |C_{yx}| + |D_{xy}| + |D_{yy}| = O(\sigma),
\]

then \( f(s; s) \) is significantly less sensitive to trait \( y \) than to trait \( x \). In this case, Eq. (2a) is transformed into Eq. (3),

\[
f(s; s) = G_x \delta x + C_{xx}(x - x_0) \delta x + \frac{1}{2} D_{xx} \delta x^2 + G_y \delta y + O(\sigma^3).
\]

If Eq. (6a) holds,

\[
G_x = 0,
\]

then there exists a singular line \( x = x_0 \) denoted by \( x_0 \). This line is convergence stable if Eq. (6b) holds,

\[
C_{xx} < 0
\]

(i.e., if \( C < 0 \)). The convergence-stable line causes evolutionary branching along MLIPs (maximum-likelihood-invasion paths), if inequality (18), i.e., the MLIP condition, holds,

\[D = \frac{\sigma D_{xx}}{2|C_{yy}|} > \frac{1}{\sqrt{2}}\].

In this case, any MLIP starting from a monomorphic resident \( s_0 = (x_0, y_0)^T \) with \( |s_0 - s_0| = O(\sigma) \) inevitably converges to the line and brings about evolutionary branching. Here we refer to Eqs. (4), (6a), (6b), and (18) as the conditions for evolutionary-branching lines, which is summarized as follows.

**Theorem 2.** Suppose that a point \( s_0 \) in a normalized trait space \( s \) satisfies the conditions for evolutionary-branching lines below. Then there exists an evolutionary-branching line passing through \( s_0 \) parallel with the \( y \)-axis of the trait space. In this case, any MLIP starting from a monomorphic resident \( s_0 = (x_0, y_0, y_0, y_0)^T \) with \( |s_0 - s_0| = O(\sigma) \) monotonically converges to the line and brings about a protected dimorphism, which leads to dimorphic divergence in \( x \) without collapse, as long as the deviations from \( s_0 \) are \( O(\sigma) \).

**Conditions for evolutionary-branching lines along MLIPs in normalized bivariate trait spaces.** The \( x \)-and \( y \)-axes can be adjusted by rotation such that the first and second derivatives of the invasion-fitness function at \( s_0 \) satisfy Eqs. (4), (6a), (6b), and (18).

Rescaling trait spaces such that \( \sigma = 1 \) and applying Theorem 1 proves this theorem.

If these conditions for evolutionary-branching lines hold, then evolutionary branching occurs with high likelihood in evolutionary paths even under relaxed assumptions (i.e., in OSIPs and PSIPs, as shown in Section 6). As the sensitivity difference goes to infinity, which means that the right-hand side of Eq. (4) converges to zero,
the conditions for evolutionary-branching lines converge to the univariate conditions for evolutionary branching, given by Eqs. (6).

Notice that the MLIP condition requires that $\sigma$ is not infinitesimally small, but finite; otherwise, satisfying this inequality is impossible. Thus, as long as the population is directionally evolving, its evolutionary branching requires finite mutational step sizes. Conversely, $\sigma$ can have large magnitudes, as long as approximating the invasion-fitness functions by the normal form in Eq. (3) is appropriate at the scale of that $\sigma$. In this case, a single large mutational step may generate a mutant such that the mutant and resident together straddle an evolutionary-branching line, resulting in protected dimorphism with a relatively large phenotypic difference. Although this sounds different from the process of evolutionary branching with small phenotypic difference, the two cases are formally equivalent as long as the invasion-fitness function is well approximated by Eq. (3), i.e., terms beyond $\sigma$ insignificantly small, but finite. Thus, as long as the population is directionally evolving, the conditions for evolutionary branching given by Eqs. (6) can be satisfied by the normal form in Eq. (3), as explained in Appendix O.

Theorem 3. Suppose that a point $s_0$ in a normalized $L$-dimensional trait space satisfies the conditions for evolutionary-branching manifolds below. Then there exists an $(L-1)$-dimensional evolutionary-branching manifold passing through $s_0$ and vertical to the sensitive direction in the space, denoted by $x$. In this case, any MLIP starting from a monomorphic resident $s_0$ with $|s_0 - s_0| = O(\sigma)$ monotonically converges to the manifold and brings about a protected dimorphism, which leads to dimorphic divergence in $x$ without collapse, as long as the deviations from $s_0$ are $O(\sigma)$.

Conditions for evolutionary-branching manifolds along MLIPs in normalized multivariate trait spaces. Trait axes can be adjusted by rotation such that the first and second derivatives of the invasion-fitness function at $s_0$ satisfy Eq. (21) with $L=1$, and then the simplified invasion-fitness function, Eq. (22), satisfies Eqs. (6a), (6b), and (18).

Even if the sensitive subspace is more than univariate, conditions for evolutionary branching might be constructed in a similar form, as explained in Appendix O.

7.2. Conditions for evolutionary-branching lines in multivariate trait spaces

The conditions for evolutionary-branching lines explained above can be applied also to multivariate trait spaces: for this we only have to extend the condition for significant sensitivity difference, as explained below.

As before, an arbitrary $L$-variate trait space $S = (U_1, ..., U_L)^T$ can be normalized by an appropriate affine transformation into a representation $s = (u_1, ..., u_L)^T$ with isotropic mutation with standard deviation $\sigma$ (see Appendix P). In this normalized trait space, the invasion-fitness function can be expanded as in the bivariate case,

$$f(s'; s) = G_0 s + (s-s_0)^T C_0 s + \frac{1}{2} s^T D s + O(\sigma^3),$$

(20)

where $\delta s = (\delta u_1, ..., \delta u_L)^T = s - s$, $G$ is a $L$-dimensional row vector, and $C$ and $D$ are $L$-by-$L$ matrices.

In a manner similar to the bivariate case, the trait space can be decomposed into a $L$-variate sensitive subspace $\mathbf{x} = (x_1, ..., x_L)^T = (u_1, ..., u_L)^T$ and an $(L-1)$-variate insensitive subspace $\mathbf{y} = (y_1, ..., y_{L-1})^T = (u_{L-1}, ..., u_L)^T$, if trait axes can be adjusted such that

$$[G_{ij} + |C_{ij}| + |C_{ji}| + |D_{ij}| + |D_{ji}|] \lambda = 0 \quad \text{for all } i = 1, ..., L, j = 1, ..., L, \text{ where } G_j \text{ is the jth component of } G \text{, and } C_{ij} \text{ and } D_{ij} \text{ are the (i,j)th components of } C \text{ and } D, \text{ respectively.}$$

(21)

If $\lambda L = 1$, i.e., the sensitive subspace is univariate, then Eq. (20) simplifies to the normal form for invasion-fitness functions with significant sensitivity difference,

$$f(s'; s) = G_0 e \delta x + C_{xx}(x-x_0) \delta x + \frac{1}{2} D_{xx} \delta x^2 + G_0 \delta y + O(\sigma^3),$$

(22)

where $G_0 = G_{xx}$, $C_{xx} = C_{xx}$, $D_{xx} = D_{xx}$, $G_0 = |G_{ij}| = |G_{ji}| = |G_{ji}| = |G_{ji}|$, and $\delta y = (\delta x_0 - y_0)^T/C_{yy}$. See Appendix O for derivations of Eqs. (21) and (22). Notice that the insensitive subspace $y$ contributes to the invasion fitness only through the element parallel to the fitness gradient $G_0$. Thus, local evolutionary dynamics based on Eq. (22) can be contracted into a bivariate trait space $(x, y)$. As Eq. (22) is identical to the bivariate invasion-fitness functions with significant sensitivity difference, Eq. (3), the conditions for evolutionary-branching lines in bivariate trait spaces, Eqs. (4), (6a), (6b), and (18), can be applied directly. If $s = s_0$ satisfies those conditions, it forms in the trait space $s$ an $(L-1)$-dimensional evolutionary-branching manifold, $\mathbf{x} = x_0$. Thus, Theorem 2 is translated as follows.

Theorem 3. Suppose that a point $s_0$ in a normalized $L$-dimensional trait space $s$ satisfies the conditions for evolutionary-branching manifolds below. Then there exists an $(L-1)$-dimensional evolutionary-branching manifold passing through $s_0$ and vertical to the sensitive direction in the space, denoted by $x$. In this case, any MLIP starting from a monomorphic resident $s_0$ with $|s_0 - s_0| = O(\sigma)$ monotonically converges to the manifold and brings about a protected dimorphism, which leads to dimorphic divergence in $x$ without collapse, as long as the deviations from $s_0$ are $O(\sigma)$.

7.3. Finding evolutionary-branching lines without prior normalization

For checking conditions for evolutionary-branching lines (or manifolds) in an arbitrary trait space $S$ with arbitrary dimension $L$, the vector $G$ and the matrices $C$ and $D$ of the invasion-fitness function $F(S; S)$ are all that is needed. These are given by

$$G = F_m Q_1^T B,$$

$$C = B' Q F_{mm} + F_{rm} Q_1^T B,$$

$$D = B' Q F_{mm} Q_1^T B,$$

(23)

where $F_m$ is the gradient vector of $F(S; S)$ (comprising first derivatives) with respect to $S$ at $S_0$, while $F_{mm}$, $F_{rm}$, and $F_{mm}$ are the Hessian matrices (comprising second derivatives) there, where the subscripts ‘m’ and ‘r’ correspond to the mutant $S$ and the resident $S$, respectively. The matrix $Q$, which describes the normalization of the trait space to attain isotropic mutation with standard deviation $\sigma$, fulfills $A = (\sigma Q)^T (\sigma Q)$, while the matrix $B$, which describes the adjustment of the axes by rotation, is given by

$$B = (v_{d1}, ..., v_{dL}).$$

(24)

where $v_{d1}, ..., v_{dL}$ are the eigenvectors of $Q F_{mm} Q_1^T$, ordered such that the corresponding eigenvalues satisfy $\lambda_{d1} > \lambda_{d2}$ for all $j = 2, ..., L$. See Appendix P for the derivations of Eqs. (23) and (24).

Notice that the conditions for evolutionary-branching lines (or manifolds) explained above are based on locally approximated invasion-fitness functions. Thus, satisfying those conditions at $S_0$ ensures the existence of an evolutionary-branching line (or manifold) only at the local scale around this point. However, it is easily shown that if $s_0$ satisfies those conditions, some other points slightly deviated from $s_0$ are also expected to satisfy those conditions. By connecting these points, evolutionary-branching lines (or manifolds) can be found at the global scale (Ito and Dieckmann, 2012).

8. Discussion

In this paper, we have analytically obtained the conditions for evolutionary branching when the invasion-fitness function has significant sensitivity differences among directions in bivariate trait spaces, by focusing on evolutionary paths, called MLIPs, composed of invasions each of which has maximum likelihood. The result, called the MLIP condition, is numerically demonstrated to be a useful indicator for the likelihood of evolutionary branching in evolutionary-branching lines calculated under relaxed assumptions of stochastic invasions (OSIPs) and of non-rare mutations (PSIPs). The obtained conditions have been extended to multivariate trait spaces.
The MLIP condition requires stronger disruptive selection than is needed for univariate branching along OSIPs (Metz et al., 1996; Geritz et al., 1997, 1998). The MLIP condition remains unchanged in multivariate trait spaces as long as the sensitive subspace is univariate, because directional evolution in the insensitive subspace can be contracted into a single dimension. Thus, the MLIP condition generalizes the univariate branching conditions to situations in which a population slowly evolves by weak directional selection in other traits. This generalization is important, as real populations feature many evolving traits with a large variation in evolutionary speeds, with the result that the slow traits are likely to keep evolving directionally after the fast traits have converged to an evolutionary singularity.

One of our main assumptions is that mutational step sizes are sufficiently small so that the first- and second-order terms of the invasion-fitness functions (quantifying the strength of directional and stabilizing/disruptive selection pressures, respectively) capture the dominant selection pressures. On one hand, this of course does not require mutational step sizes to be infinitesimally small, but just to be small enough to ensure a good local approximation of the invasion-fitness function. Indeed, for the MLIP condition to hold, mutational step sizes must be finite and exceed a certain threshold. On the other hand, the MLIP condition cannot be applied when the higher-order terms of invasion-fitness functions have a non-negligible influence, which implies selection pressures that are more complex than combinations of directional and stabilizing/disruptive selection. Therefore, as long as we try to understand selection pressures as combinations of directional and stabilizing/disruptive selection, our assumption of small mutational steps is appropriate.

The conditions for evolutionary-branching lines, which are a combination of the condition for significant sensitivity difference, the condition for convergence stability, and the MLIP condition, can be used to examine the likelihood of evolutionary branching that could not be treated by previous branching conditions requiring convergence-stable singular points (Ackermann and Doebeli, 2004; Ito and Shimada, 2007). For example, Leimar (2005) and Ito et al. (2009) have numerically shown that evolutionary branching occurs in bivariate trait spaces that do not contain any evolutionarily singular points that are convergence stable. In these cases, there exists instead an evolutionarily singular point that is convergence stable only in one direction, but unstable in the other direction. By applying the conditions we have presented here, evolutionary-branching lines can be identified in the trait spaces of those models (Ito and Dieckmann, 2012). In such applications, the condition for significant sensitivity-difference might be relaxed further, or be omitted altogether, because this condition partly overlaps with the MLIP condition; and the non-overlapping parts of the sensitivity-difference condition may be required only for enabling the analytical derivation of the MLIP condition. In this sense, the MLIP condition may still work well even when the sensitivity-difference condition does not hold.

As the MLIP condition describes how weak directional selection needs to be in comparison with disruptive selection for evolutionary branching to occur, this information may also be useful for predicting evolutionary branching in the vicinity of evolutionary-branching points. The MLIP condition then describes how close to an evolutionary-branching point a monomorphic population has to come for evolutionary branching to occur when mutational steps are finite. Based on a heuristic modification of the conditions for evolutionary-branching lines, areas possessing high likelihoods of evolutionary branching can thus be identified around evolutionary-branching points. The resultant evolutionary-branching areas (Ito and Dieckmann, 2012) are important not the least because, in reality, invasion-fitness functions are always changing at least slowly, due to environmental changes or due to the evolution of other species, implying slow shifts of the locations of evolutionary branching points in trait spaces. Such shifts may prevent monomorphic populations from sufficiently converging to the points required for the emergence of dimorphism, or they may destroy initial dimorphisms even after those have emerged (Metz et al., 1996; Metz, 2011). In such cases, by examining whether environmental changes are sufficiently slow for monomorphic populations to enter the evolutionary-branching areas, likelihoods of evolutionary branching may be estimated also under such more realistic circumstances.

A focus on MLIPs, treated as typical and deterministic paths among corresponding OSIPs, has enabled our analytical treatment of evolutionary branching in bivariate trait spaces. This analysis of MLIPs is adding to the evolutionary literature a second deterministic description of mutation-limited evolutionary dynamics. The more common alternative is the mean evolutionary path defined by the canonical equation of adaptive dynamics theory (Dieckmann and Law, 1996). Roughly speaking, such a mean evolutionary path is formed by mutant invasions, each of which occurs by the mean mutant phenotype among all mutants that are able to invade, weighted according to their invasion probabilities. It is therefore interesting to consider how these two deterministic descriptions of mutation-limited evolutionary dynamics are related. In particular, an MLIP is identical to the corresponding mean evolutionary path given by the canonical equation, if directional evolution of a single population with a multivariate Gaussian mutation distribution is considered, although the speed along the MLIPs is $\sqrt{2/\pi} \approx 0.798$ times as fast as along the corresponding mean paths (see Appendix Q).

In general, however, MLIPs and mean evolutionary paths are different, because an MLIP is formed by mutants that are the modes of the invasion-event probability distribution at each invasion event, while a mean path is formed by mutants that are the means of this distribution. Thus, differences between the two descriptions can arise, especially when the mutation distribution is discrete, as, e.g., for the univariate fixed-step mutation distribution. As MLIPs are affected only by the global maximum of an invasion-event probability distribution, but not by any other of its features, and as a distribution’s global maximum may abruptly change its role with a local maximum, the mean evolutionary paths may be deemed more robust than MLIPs for describing directional evolution. On the other hand, by construction, the canonical equation is not capable of describing evolutionary branching, while MLIPs can do so. To our knowledge, MLIPs are the only way of deterministically describing evolutionary dynamics that include evolutionary diversifications, without loss of analytical tractability. Therefore, MLIPs may be useful for analyzing other evolutionary-phenomena in multivariate trait spaces.

Our analyses of invasion-event probabilities are related to phylogenetic reconstruction and ancestral-state reconstruction based on empirical data (Wiens, 2000; Barton et al., 2007), and may hint at worthwhile extensions of such methods. The standard methods for ancestral-state reconstruction first reconstruct phylogenetic trees based on DNA sequences, and then reconstruct the ancestral states of focal traits based on those trees, using constraints alternatively given by maximum parsimony, maximum likelihood, Bayesian methods, etc. While our MLIPs maximize likelihoods not at the level of invasion sequences, but at the level of invasion events, numerical methods could be devised to maximize the likelihoods of OSIPs. When those OSIPs are calculated from a given present composition of residents backward to their common ancestor (e.g., using Markov Chain Monte Carlo methods), the past evolutionary dynamics can be reconstructed as a phylogeny in a trait space. In this case, phylogeny and ancestral states are reconstructed at once, based on a given fitness function providing prior information. This alternative kind of phylogenetic reconstruction might be useful for some genera or families, if key quantitative traits and associated ecological settings are known sufficiently well.
for such knowledge to be translated into fitness functions, and if exogenous changes of those functions are expected to be small. Comparing results obtained from this ecology-based reconstruction method with results from standard reconstruction methods might provide new insights.

In our approach, it is important to identify the fast traits, or fast phenotypic directions, along which a considered evolutionary diversification unfolds. Empirical evidence suggests that in some taxonomic groups the directions of observed trait differences among related populations are positively correlated with the trait directions possessing the largest additive genetic variance within the populations. These directions, which have been called “lines of least resistance” (Schluter, 1996), closely resemble the fast traits in our approach. If corresponding slow traits can also be identified, our conditions for evolutionary-branching lines and manifolds may be applied to understand the evolutionary ecology of the underlying diversifications.

Acknowledgement

The authors thank Hans Metz, Géza Meszéna, Michael Doebeli, Yoh Iwasa, Takashi Ikemgami, Masakazu Shimada, two anonymous reviewers and an editor for valuable comments on earlier versions of this manuscript. H.I. also thanks David Murno for inventing a tool for numerical analysis and visualization, named Yorick, and distributing it for free. Figs. 4 and 5 in this study were produced with inventing Yorick. H.I. acknowledges support in the form of a Research Fellowship for Young Scientists by the Japan Society for the Promotion of Science (JSPS), and by IIASA’s Evolution and Ecology Program, facilitating his participation in the Young Scientists Summer Program (YSSP) of the International Institute for Applied Systems Analysis (IIASA). U.D. gratefully acknowledges financial support by the European Science Foundation, the Austrian Science Fund, the Austrian Ministry of Science and Research, and the Vienna Science and Technology Fund, as well as by the European Commission, through the Marie Curie Research Training Network FishACE and the Specific Targeted Research Project FinE.

Appendix A. Derivation of quadratic form of invasion-fitness functions

Here we derive an approximate quadratic form of \( f(s'; s) \), Eq. (2) in Section 3. For convenience, and without any loss of generality, we shift the origin close to the resident phenotype, so that \( |s| = O(\sigma) \) and \( |s'| = O(\sigma) \). We then expand \( f(s'; s) \) about this origin \( s_0 = (0, 0)^T \) as

\[
\begin{align*}
 f(s'; s) &= f_m s' + f_{m'f} s + 0.5 s' f_{mm} s + s' f_{mf} s + 0.5 s' f_{mm} s + O(\sigma^3), \\
 &= f_m s' + f_{m'f} s + 0.5 s' f_{mm} s + s' f_{mf} s + 0.5 s' f_{mm} s + O(\sigma^3), \tag{A.1a}
\end{align*}
\]

with

\[
\begin{align*}
 f_m &= \frac{f_{xx} f_{yy} + f_{xy} f_{yx}}{s = s = s_0}, \\
 f_{m'f} &= \frac{f_{xx} f_{yy} - f_{xy} f_{yx}}{s = s = s_0}, \\
 f_{mm} &= \frac{f_{xx} f_{yy} + f_{xy} f_{yx}}{s = s = s_0}, \\
 f_{mf} &= \frac{f_{xx} f_{yy} - f_{xy} f_{yx}}{s = s = s_0},
\end{align*}
\]

where the subscripts ‘\( m \)’ and ‘\( f \)’ refer to mutants and residents, respectively, and where \( f_{\alpha\beta} = \frac{df(s'; s)}{d\alpha} \) for \( \alpha = x, y, y' \) and \( f_{\alpha\beta} = \frac{d^2f(s'; s)}{d\alpha d\beta} \) for \( \alpha, \beta = x', y', y \) denote the first and second derivatives of \( f(s'; s) \), respectively. We transform Eq. (A.1a) into

\[
\begin{align*}
 f(s'; s) &= f_m (s' - s) + 0.5 (s' - s) s' f_{mm} (s' - s) + s' f_{mm} f_{mf} (s' - s) \tag{A.1b}
\end{align*}
\]

where

\[
\begin{align*}
 s' f_{mm} = \begin{pmatrix} f_{mm} + f_{im} + f_{mm} \end{pmatrix} s + O(\sigma^3), \tag{A.2a}
\end{align*}
\]

and

\[
\begin{align*}
 s' f_{mm} s' = \begin{pmatrix} f_{mm} f_{mm} \end{pmatrix} s^2 + O(\sigma^3), \tag{A.2b}
\end{align*}
\]

Second, we extend this relationship into a general condition for sensitivity difference due to mutational asymmetry

\[
\begin{align*}
 f(s'; s) &= f_m (s' - s) + 0.5 (s' - s) s' f_{mm} (s' - s) + s' f_{mm} f_{mf} (s' - s) + O(\sigma^3) \tag{A.3}
\end{align*}
\]

where the second line uses the notation introduced in Eq. (2b). The substitution \( s_0 = s' - s \) and considering \( s_0 \neq (0, 0)^T \) give rise to Eq. (2a) in Section 3. Notice that the quadratic approximation in Ito and Dieckmann (2012), where \( C \) is multiplied by \( 1/2 \), becomes identical to Eq. (2a) by defining \( C \) as \( C = f_{mm} + f_{mf} \) (the convention used here) instead of as \( C = 2(f_{mm} + f_{mf}) \) (the convention used there).

Appendix B. Condition for significant sensitivity difference

Here we derive the condition for significant sensitivity difference of normalized invasion-fitness functions, Eq. (4) in Section 3. First, we show how sensitivity difference can be caused by the asymmetry of mutational step sizes in the original trait space. Second, we extend this relationship into a general condition for significant sensitivity difference.

Sensitivity difference due to mutational asymmetry

We assume that the \( X \) and \( Y \)-axes of the original trait space \( S \) have been aligned as shown in Fig. 2b, so that \( V_X = 0 \). In this space, the invasion-fitness function \( F(s'; s) \) is expanded similarly to Eq. (2) as

\[
\begin{align*}
 F(s'; s) &= G(s; s') \left[ 2 s' s + C_X Y + 2 s' s + C_Y X + O(\sigma^2) \right], \tag{B.1a}
\end{align*}
\]

where

\[
\begin{align*}
 G &= \begin{pmatrix} C_X & C_Y \end{pmatrix}, \quad C = \begin{pmatrix} C_{XX} & C_{XY} \\ C_{YX} & C_{YY} \end{pmatrix}, \\
 D &= \begin{pmatrix} D_{XX} & D_{XV} \\ D_{VX} & D_{YY} \end{pmatrix}. \tag{B.1b}
\end{align*}
\]

We now consider the case that \( \sigma_X \) is much smaller than \( \sigma_Y \), such that \( \sigma_Y = O(\sigma^2) \). We introduce a new coordinate system \( s' = (x, y)^T \), where \( x = X \) and \( y = \sigma_Y / \sigma_X Y \). This results in isotropic mutation with standard deviation \( \sigma = \sigma_X \) (Fig. 2c). Substituting \( X = x \) and \( Y = \sigma_Y / \sigma_X Y \) into Eq. (B.1a) yields the following normalized invasion-fitness function,

\[
\begin{align*}
 f(s'; s) &= f(x', \sigma_Y / \sigma_X y') \tag{B.2a} \\
 &= G(x' \sigma_Y / \sigma_X y') + C_X x' \sigma_Y / \sigma_X y' + C_Y y' \sigma_Y / \sigma_X y' + O(\sigma^2), \\
 &= G_{x'} x' \sigma_Y / \sigma_X y' + G_{y'} y' \sigma_Y / \sigma_X y' + O(\sigma^2), \\
 &= G_{x' y'} x' y' \sigma_Y / \sigma_X y' + O(\sigma^2), \tag{B.2b}
\end{align*}
\]
Thus, by including all applicable terms in $O(\sigma^3)$, we see that Eq. (B.2a) yields the normal form of invasion-fitnss functions with significant sensitivity difference, Eq. (3) in Section 3.

**Generalization of sensitivity difference**

The normal form in Eq. (3) can also be obtained when the sensitivity of a fitness function to variation in trait $y$ is weak, so that $G_y, D_{xy}, C_{xy}, C_y, \text{ and } C_y$ are all relatively small. To make this notion precise, we proceed as follows. We suppose an arbitrary invasion-fitness function defined in a normalized trait space $s = (x, y)^T$, in which mutation is isotropic with standard deviation $\sigma \approx 1$. This function is then given by Eq. (2),

$$f(s'; s) = Gs' + (s - s_0)TbD_0s + \frac{1}{2}b^TD_0b + O(\sigma^3).$$

Here we assume that the value of invasion fitness is expressed as Eq.(4),

$$h(s') = h(s') + b(s' - s_0)s_0,$$  \hspace{1cm} \text{(B.4)}

where $h(s') = f(s'; s) - g(x', x)$ is kept smooth and finite, i.e., its first and second derivatives are $O(\epsilon)$. Then it is said that the function $h$ has significant sensitivity difference with respect to the $x$- and $y$-directions.

If $f$ in Eq. (B.3) has significant sensitivity difference, then substituting Eq. (B.4) into Eq. (B.4) and assuming $\epsilon = \sigma$ yields

$$g(x'; x) = g(x'; x) + C_{xy}(x - x_0)dx + \frac{1}{2}D_{xx}dx^2 + O(\sigma^3),$$

$$h(s'; s) = \frac{C_{yx}}{\sigma}dy + \frac{C_{yy}}{\sigma}(y - y_0)dy + \frac{C_{xy}}{\sigma}dy + \frac{D}{2\sigma^2}dy + \frac{D}{2\sigma^2}s_0dy + O(\sigma^3).$$

To keep the first and second derivatives of $h(s'; s)$ at $O(\sigma^2)$, $G_y$, $D_{xy}$, $C_{xy}$, $D_{xx}$ and $D_{yy}$ must be $O(\sigma)$, in which case Eq. (2a) in Section 3 can be transformed into Eq. (3). Thus, a sufficient condition for the normalized invasion-fitness function in Eq. (2a) to be transformed into the normal form in Eq. (3) is expressed as Eq. (4),

$$|G_y| + |C_{xy}| + |C_{yy}| + |D_{xy}| + |D_{xx}| = O(\sigma),$$  \hspace{1cm} \text{(B.6)}

where the denominator enables application to normalized invasion-fitness functions that are not yet suitably scaled.

**Appendix C. Approximation of invasion-event rate density**

Here we explain how the invasion-event rate density is approximated in Eq. (8a) in Section 4. The first row of Eq. (8a) is transformed into

$$E(s'; s) = \mu b(s; s)M(s' - s) + f(s'; s) + b(s; s)\mu b(s'; s) = \mu b(s' - s)f(s'; s) + (1 + \phi h(s'; s)).$$  \hspace{1cm} \text{(C.1a)}

where

$$\phi h(s'; s) = \frac{b(s; s) - b(s'; s)}{b(s; s)} = O(\sigma),$$  \hspace{1cm} \text{(C.1b)}

because $b(s'; s)$ converges to $b(s; s)$ as $\sigma \to 0$, which gives $s = s'$. Thus, the approximation in Eq. (8a) applies in the leading order of $s = s'$, as long as $\sigma$ is sufficiently small such that $b(s'; s) - b(s; s)$ is much smaller than $b(s'; s)$, i.e., $[b(s'; s) - b(s; s)]/b(s'; s) = O(\sigma)$.

**Appendix D. MLIP condition**

Here we prove that the conditions for dimorphic emergence and those for dimorphic divergence are both satisfied if and only if the MLIP condition $D > 1/\sqrt{2}$ holds, provided that $C < 0$.

**Main proof**

When the resident is located on the convergence-stable line, MLI mutants $s_{MLI}$, given by Eq. (17) in Section 5, do not satisfy the second condition for dimorphic emergence, inequality (13b), for $D \leq 1/\sqrt{2}$. Thus, $D > 1/\sqrt{2}$ is necessary for dimorphic emergence. Therefore, we only have to examine whether $D > 1/\sqrt{2}$ satisfies the conditions for dimorphic emergence and dimorphic divergence, as follows.

First, we examine the conditions for dimorphic emergence, i.e., inequalities (13). For convenience, instead of the MLI mutant itself, $s_{MLI}$, we use its deviation from its parental resident, $s_{MLI} - s$. For this $s_{MLI}$, the following lemma holds (see the next subsection for the proof).

**Lemma D.1.** If the MLIP condition $D > 1/\sqrt{2}$ holds, then, for any monomorphic resident $s$, the MLI mutation $s_{MLI}$ satisfies

$$\delta_{MLI} > 0 \quad \text{for } x < 0,$$

$$\delta_{MLI} < 0 \quad \text{for } x > 0,$$

$$0 < \delta y_{MLI} < D\delta x_{MLI},$$

$$1 < |\delta x_{MLI}| < \sqrt{2}.$$  \hspace{1cm} \text{(D.1a) - (D.1c)}

Inequality (D.1a) is identical to inequality (13a). Inequality (D.1b) is a sufficient condition for inequality (13b), Thus, Lemma D.1 ensures that conditions for dimorphic emergence hold under the MLIP condition.

Next, we examine the conditions for dimorphic divergence, i.e., inequalities (14). Under dimorphism of $s_1$ and $s_2$, the MLI mutation is given by $s_{MLI} = s_{MLI} - s_{MLI}$ with $s_{MLI} = s_1$ or $s_2$. For this $s_{MLI}$, the following lemma holds (see the last subsection for the proof).

**Lemma D.2.** If the MLIP condition $D > 1/\sqrt{2}$ holds, then, for any dimorphic residents $s_1$ and $s_2$ satisfying $|y_2 - y_1| < D(x_2 - x_1)^2$, the MLI mutation $s_{MLI}$ satisfies

$$\delta_{MLI} > 0 \quad \text{for } x < 0,$$

$$\delta_{MLI} < 0 \quad \text{for } x > 0,$$

$$0 < \delta y_{MLI} < D\delta x_{MLI},$$

$$1 < |\delta x_{MLI}| < \sqrt{2}.$$  \hspace{1cm} \text{(D.2a) - (D.2c)}

where $s((x, y)^T = [(D(2x_{MLI} - x_2 - x_1) - (y_2 - y_1)/(x_2 - x_1))/C_0, 0)^T$.

Below we check whether Lemma D.2 ensures that the conditions for dimorphic divergence, inequalities (14), hold under $D > 1/\sqrt{2}$. As in inequalities (14), $x_1 < x_2$ is assumed without loss of generality. First, we suppose $s_{MLI}$ originates from $s_1$, i.e., $s_{MLI} = s_1$, and $s_{MLI}$, in which case inequality (14a), i.e., $x_{MLI} < x_1$ and $|y_{MLI} - y_1| < D(x_{MLI} - x_1)^2$, hold as explained below. Clearly, the second inequality, $|y_{MLI} - y_1| < D(x_{MLI} - x_1)^2$, is satisfied by inequality (D.2b). The first inequality, $x_{MLI} < x_1$, also holds under
inequalities (D.2a), because $\delta s_{\text{MLI}} = x_{\text{MLI}} - x_1 < 0$ follows from $\dot{x} > 0$:

$$\dot{x} = \frac{1}{C(x_1-x_2)} \left[(D(x_1-x_2) y_2 - y_1) x_2 - x_1 \right] = \frac{1}{C(x_1-x_2)} \left[D(x_1-x_2)^2 + (y_2 - y_1) \right]$$

$$\geq 0 \quad \text{if} \quad \frac{1}{C(x_1-x_2)} \left[D(x_1-x_2)^2 - |y_2 - y_1| \right] \geq 0,$$

$$\leq 0 \quad \text{if} \quad \frac{1}{C(x_1-x_2)} \left[-D(x_1-x_2)^2 + (y_2 - y_1) \right] \leq 0,$$

$$< 0 \quad \text{if} \quad \frac{1}{C(x_1-x_2)} \left[-D(x_1-x_2)^2 + D(x_2-x_1)^2 \right] < 0. \quad (D.3)$$

where $C(x_1-x_2)$ is positive because $C > 0$ and $x_1 - x_2 < 0$.

Second, we suppose $s_{\text{MLI}}$ originates from $s_2$ (i.e., $s_{\text{MLI}} = s_2$ and $\delta s_{\text{MLI}} = s_{\text{MLI}} - s_2$), in which case inequality (14b), i.e., $x_{\text{MLI}} > x_2$ and $|y_{\text{MLI}} - y_1| > D(x_{\text{MLI}} - x_2)^2$, again hold as explained above. Clearly, the second inequality, $|y_{\text{MLI}} - y_1| > D(x_{\text{MLI}} - x_2)^2$, is satisfied by inequality (D.2b). The first inequality, $x_{\text{MLI}} > x_2$, also holds under inequalities (D.2a), because $\delta s_{\text{MLI}} = x_{\text{MLI}} - x_2 > 0$ follows from $\dot{x} < 0$:

$$\dot{x} = \frac{1}{C(x_1-x_2)} \left[(D(x_2-x_1) y_2 - y_1) x_2 - x_1 \right] = \frac{1}{C(x_1-x_2)} \left[-D(x_2-x_1)^2 + (y_2 - y_1) \right]$$

$$\leq 0 \quad \text{if} \quad \frac{1}{C(x_1-x_2)} \left[-D(x_2-x_1)^2 + y_2 - y_1 \right] \leq 0,$$

$$< 0 \quad \text{if} \quad \frac{1}{C(x_1-x_2)} \left[-D(x_2-x_1)^2 + D(x_2-x_1)^2 \right] < 0. \quad (D.4)$$

Thus, the conditions for dimorphic divergence, inequalities (14), as well as those for dimorphic emergence, inequalities (13), hold under $D > 1/\sqrt{2}$. Therefore, conditions for dimorphic emergence and those for dimorphic divergence both hold if and only if the MLIP condition $D > 1/\sqrt{2}$ holds. This completes the proof.

Proof of Lemma D.1

Among the three inequalities (D.1) of Lemma D.1, we first analyze inequalities (D.1a). MLI mutants, which maximize $P(s', s)$ in Eq. (8b), always have positive invasion fitnesses. Thus, the subscript “+” is not needed here. For arbitrary $s'$ and $s$, $P(s', s)$ satisfies

$$P(\mathbf{s} \rightarrow s; s') = T \frac{\mu}{2\pi} \exp\left(-\frac{1}{2}(s'^2 + s^2)\right) C_{\mathbf{s} \rightarrow \mathbf{s}', \mathbf{s}}, \quad (D.5)$$

where $\mathbf{s} = (s, \bar{s})$ is identical to $\mathbf{s} = (s, \bar{s})$ except that the sign of $\mathbf{s}$ is reversed. When $\delta s$ is positive, $\delta s > 0$, then $\mathbf{s} = \mathbf{s} = (s, \bar{s})$. Then, we prove Lemma D.1 holds, substitution of $\mathbf{s}_{\text{MLI}}(\mathbf{s}, \mathbf{s}') = \mathbf{s}_{\text{MLI}}(s, \bar{s})$ and $\mathbf{s}$ into Eq. (D.1) immediately gives Lemma D.2. The proof of $\mathbf{s}_{\text{MLI}}(s_1, s_2) = \mathbf{s}_{\text{MLI}}(s)$ as follows.

The MLIP mutation $\mathbf{s}_{\text{MLI}}(s_1, s_2)$ is given by the $\mathbf{s}$ that maximizes $P(s', s)$ in Eq. (9b) in Section 4.

$$P_{\text{MLI}}(\mathbf{s} \rightarrow s_1, s_2) = T \frac{\mu}{2\pi} \exp\left(-\frac{1}{2}(s_1^2 + s_2^2)\right) C_{\mathbf{s} \rightarrow \mathbf{s}', \mathbf{s}} \quad (D.11)$$

which hold for $\mathbf{s} = \mathbf{s}_{\text{MLI}}$. Eq. (D.6a) is transformed into

$$\delta y = -D(D\delta x^2 - 2) + C(x_1 - x_2) \quad \text{g}(\delta x), \quad (D.7)$$

where the second equation defines $g(\delta x)$. Taking the difference of Eqs. (D.6a) and (D.6b),

$$\frac{1}{\delta x} \frac{\partial}{\partial \delta x} \left[ \frac{\partial \mathbf{s} \rightarrow \mathbf{s}}{\partial \delta y} \right] = 0, \quad (D.8)$$

we find

$$\delta y = \frac{\delta x}{2D\delta x + Cx} = h(\delta x). \quad (D.9)$$

where the second equation defines $h(\delta x)$.

When $\delta x < 0$, in which case $\delta s_{\text{MLI}}$ must be positive, $\delta s_{\text{MLI}}$ is the crossing point of the two curves given by Eqs. (D.7) and (D.9). As $\delta g(\delta x) / \delta x < 0$ and $\delta h(\delta x) / \delta x > 0$ hold for positive $\delta x$ and $D > 0$, $g(\delta x)$ and $h(\delta x)$ are monotonically decreasing and increasing functions of $\delta x$, respectively, for positive $\delta x$ and $D > 0$. Suppose that $D = 1/\sqrt{2}$. Then $h(1) - (1) < 0$ and $h(1/\sqrt{2}) - g(1) > 0$. Since $h(\delta x) - g(\delta x)$ is a strictly increasing function of $\delta x$ and is zero for $\delta x = \delta x_{\text{MLI}}$, it follows that $1 < \delta x_{\text{MLI}} < \sqrt{2}$ and that $\delta x_{\text{MLI}} < \min(1, h(1/\sqrt{2})) = g(1) = D < D\delta x_{\text{MLI}}$. In addition, as $h(\delta x)$ is always positive for positive $\delta x$, $\delta x_{\text{MLI}} > 0$ holds. Thus, inequalities (D.1b) and (D.1c) both hold for $D > 1/\sqrt{2}$. This completes the proof.

Proof of Lemma D.2

For the proof in this subsection, we denote MLI mutants as functions of the resident phenotypes, i.e., $s_{\text{MLI}}(s)$ for monomorphism and $s_{\text{MLI}}(s_1, s_2)$ for dimorphism. Then the MLIP mutations in these cases can be expressed as $s_{\text{MLI}}(s) = s_{\text{MLI}}(s_1) - s_{\text{MLI}}(s_2)$ and $s_{\text{MLI}}(s_1, s_2) = s_{\text{MLI}}(s_1) - s_{\text{MLI}}(s_2)$, respectively, where $s_{\text{MLI}}$ with $s_{\text{MLI}} = s_1$ or $s_2$ is the parental resident of the MLI mutant $s_{\text{MLI}}(s_1, s_2)$.

We prove Lemma D.2 by demonstrating that the MLI mutation under a dimorphism of $s_1$ and $s_2$ is identical to that under monomorphism of an appropriately chosen $s$.

$$f(s', s_1, s_2) = (y' - y_2 + D(x' - x_1)(x' - x_2)) (x' - x_1)^2 D(x' - x_2)^2 \quad (D.12)$$

as long as the dimorphic residents are still close to the base point of the expansion, $s_0 = (0, 0)^T$ (see Appendix E for the derivation). This function can be expressed in the form of a monotonic invasion-fitness function, Eq. (12a),

$$f(s', s_1, s_2) = f(s + s_{\text{MLI}}(s_1, s_2)) \quad (D.13)$$

by choosing $s$ as in Eq. (D.10). With this relationship, Eq. (D.11) yields

$$P_{\text{MLI}}(\mathbf{s} \rightarrow s_1, s_2) = T \frac{\mu}{2\pi} \exp\left(-\frac{1}{2}(s_1^2 + s_2^2)\right) C_{\mathbf{s} \rightarrow \mathbf{s}', \mathbf{s}}, \quad (D.14)$$

where $\hat{n}$ is the equilibrium population size of the monomorphic resident $s$. Since both $\hat{n}_{\text{MLI}}$ and $\hat{n}$ do not depend on $\mathbf{s}$, any fixed $s_1$ and $s_2$ fulfill $P_{\text{MLI}}(\mathbf{s} \rightarrow s_{\text{MLI}}(s_1, s_2)) = P(\mathbf{s} \rightarrow \mathbf{s})$. Therefore, the $\mathbf{s} = \mathbf{s}_{\text{MLI}}(s_1, s_2)$ that maximizes $E_{\text{MLI}}(\mathbf{s} \rightarrow s_{\text{MLI}}(s_1, s_2))$ is
identical to the $\mathbf{s} = \mathbf{s}_{\text{null}}(\mathbf{s}) = \mathbf{s}_{\text{null}}(\hat{\mathbf{s}})$ that maximizes $E(\mathbf{s} + \hat{\mathbf{s}}, \mathbf{s})$. This completes the proof.

### Appendix E. Invariance fitness functions under dimorphism

Here we approximate dimorphic invasion-fitness functions $f(s', s_1, s_2)$ by a form similar to that derived for monomorphic invasion-fitness functions, Eq. (D.12) in Appendix D. First, we assume that $|s_2 - s_1|$ is sufficiently small for the function $f$ to be approximated using its first and second derivatives only. However, the direct Taylor expansion of $f$ with respect to $s', s_1$, and $s_2$ up to second order cannot generally satisfy the consistency condition $f(s_1; s_1, s_2) = f(s_2; s_1, s_2) = 0$.

Under the restrictive assumption that the two residents are near an evolutionarily singular point, this problem can be solved by allowing the invasion-fitness function to have a rational form composed of its first and second derivatives (Durinx et al., 2008). In our study, however, the two residents may be distant from a singular point in the $y$-direction. Here we therefore generalize the result by Durinx et al. (2008) by showing that for arbitrary dimorphic residents, the relative invasion fitness function can be approximated by a rational form composed of its first and second derivatives.

#### Special case

We first consider a special case that arises when the two residents $s_1$ and $s_2$ are located exactly on the $x$-axis, i.e., $s_1 = (x_1, 0)^T$ and $s_2 = (x_2, 0)^T$, in which case the dimorphic invasion-fitness function $f(s', s_1, s_2)$ can be approximated by a quadratic function of $s'$ and $s_2$ as follows. To satisfy $f(s_1; s_1, s_2) = f(s_2; s_1, s_2) = 0$, the quadratic function must be expressed as

$$f(s'; s_1, s_2) = \frac{1}{2}D_{xy}(x' - x_1)(x' - x_2) + \frac{1}{2}D_{yy}y'^2 + \frac{1}{2}D_{xx}x'^2 + C_{xy}x'y + C_{xx}x'^2 + C_{yy}y'^2,$$

(E.1)

with unknown constant parameters $D_{xx}$, $D_{xy}$, $D_{yy}$, $C_{xx}$, $C_{xy}$, and $C_{yy}$ to be specified. For satisfying $f(s'; s_1, s_2) = f(s'; s_2, s_1)$, $C_{xx} = C_{xy} = C_{yy}$ is required. The other parameters can be determined by comparing with the monomorphic invasion-fitness functions

$$f(s'; s_1) = G(s' - s_1) + s_1^T C(s' - s_1) + \frac{1}{2} s_1^T D(s' - s_1),$$

(E.2a)

and

$$f(s'; s_2) = G(s' - s_2) + s_2^T C(s' - s_2) + \frac{1}{2} s_2^T D(s' - s_2),$$

(E.2b)

where $s_0 = (0, 0)^T$ is assumed without loss of generality.

Now we consider a continuous shift in the resident phenotypes $s_1$ and/or $s_2$ in a way that maintains their coexistence (i.e., $f(s_1; s_2) > 0$ and $f(s_2; s_1) > 0$), such that the population size of $s_2$, denoted by $n_2$, converges to zero. Then $f(s'; s_1, s_2)$ has to converge to $f(s'; s_1)$. As derived in the last subsection, $n_2 \rightarrow +0$ while $n_1 > 0$ implies $f(s_2; s_1) = 0$, and $f(s_1; s_2) > 0$. This consideration yields the consistency condition

$$f(s'; s_1, s_2) = f(s'; s_1)$$

(E.3a)

In the same manner, considering the case of $n_1 \rightarrow +0$ while $n_2 > 0$ yields another consistency condition,

$$f(s'; s_1, s_2) = f(s'; s_2)$$

(E.3b)

First, we examine the consistency condition Eq. (E.3a). The condition $f(s_2; s_1) = 0$ is transformed into

$$x_2 = x_1 - \frac{2C_{xx}x_1 + 2C_{x_2}}{D_{xx}}.$$

(E.4)

which upon substitution into $f(s'; s_1, s_2) = f(s'; s_1)$ gives

$$f(s'; s_1, s_2) = \frac{D_{xx}G_x(x' - x_1) + D_{xx}C_{xx}(x' - x_1) + \frac{1}{2} D_{xx}(x' - x_1)^2}{D_{xx}G_x(x' - x_1) + D_{xx}C_{xx}(x' - x_1) + \frac{1}{2} D_{xx}(x' - x_1)^2} + \left[ \frac{D_{xy}(x' - x_1)y + \frac{1}{2} D_{yy}y^2}{D_{xy}(x' - x_1)y + \frac{1}{2} D_{yy}y^2} \right] + \frac{D_{yy}y + \frac{1}{2} D_{xx}x'^2}{D_{xx}G_x(x' - x_1) + D_{xx}C_{xx}(x' - x_1) + \frac{1}{2} D_{xx}(x' - x_1)^2} + \left( \frac{D_{xy}(x' - x_1)y + \frac{1}{2} D_{yy}y^2}{D_{xy}(x' - x_1)y + \frac{1}{2} D_{yy}y^2} \right) + G_{xy}y + C_{xx}x'y = f(s'; s_1).$$

(E.5)

This equation must be satisfied for arbitrary $x' - x_1$, $x'_1$, and $y'$ as long as $f(s_1; s_1) > 0$. Comparing the coefficients for $x' - x_1$, $x'_1$, and $y'$ at each order, we can specify the unknown parameters as $D_{xx} = D_{xy}$, $D_{yy} = D_{xy}$, $\tilde{C}_{xy} = \tilde{C}_{yy}$, and $\tilde{C}_{XX} = \tilde{C}_{YY}$. As the parameters thus specified also satisfy the other consistency condition, Eq. (D.13a), Eq. (D.1), and Eqs. (E.6) is an appropriate quadratic approximation of the dimorphic invasion-fitness function $f(s'; s_1, s_2)$. It can be shown that Eq. (E.1) with Eqs. (E.6) can be further transformed into

$$f(s'; s_1, s_2) = f_2(s_1; s_1) + f_2(s_1; s_2),$$

(E.7)

where $f_1 = f_{12}((/f_{12} + f_{22}))$, $f_2 = f_{22}(/f_{12} + f_{22})$, and $h = f_{12}f_{21}/(f_{12} + f_{21})$, with $f_{12} = f(s_1; s_2)$ and $f_{21} = f(s_2; s_1)$.

#### General case

Next, we consider the general case in which the two residents $s_1$ and $s_2$ are located neither on the $x$-axis (as in the previous subsection) nor near an evolutionarily singular point (as in Durinx et al. [2008]). As the dimorphic invasion-fitness function obtained for the special case above, Eq. (E.7), has a form independent of the coordinate system (i.e., independent of how the $x$- and $y$-axes are chosen), it is expected that even in this general case the function is obtained in a form identical to Eq. (E.7). Below, we confirm this conjecture.

First, to treat this general case analogously to the special case, we introduce a new coordinate system $\mathbf{s} = (x, y)^T$ so that $s_1$ and $s_2$ are both located on the $x$-axis, i.e., $s_1 = (x_1, 0)^T$ and $s_2 = (x_2, 0)^T$, by an affine coordinate transformation

$$\mathbf{s} = \begin{pmatrix} x \\ y \end{pmatrix} = \begin{pmatrix} \alpha & 0 \\ 1 & 1 \end{pmatrix} \begin{pmatrix} x \\ y \end{pmatrix} + \begin{pmatrix} 0 \\ 1 \end{pmatrix} \alpha x_1,$$

(E.8)

with $a = (y_2 - y_1)/(x_2 - x_1)$. Then, in the new coordinate system, $f(s'; s_1, s_2)$ are expressed as

$$\tilde{f}(\tilde{s}' ; \tilde{s}_1) = \tilde{f}(\tilde{s}' ; \tilde{s}_1) + \tilde{f}(\tilde{s}' ; \tilde{s}_1)$$

(E.9a)

$$= \tilde{G}(\tilde{s}' - \tilde{s}_1) + \tilde{s}_1^T \tilde{C}((\tilde{s}' - \tilde{s}_1) + \frac{1}{2} (\tilde{s}' - \tilde{s}_1)^T D(\tilde{s}' - \tilde{s}_1),$$

(E.9a)

and

$$\tilde{f}(\tilde{s}' ; \tilde{s}_2) = \tilde{f}(\tilde{s}' ; \tilde{s}_2) = \tilde{f}(\tilde{s}' ; \tilde{s}_1) + \tilde{s}_2^T \tilde{C}((\tilde{s}' - \tilde{s}_1) + \frac{1}{2} (\tilde{s}' - \tilde{s}_2)^T D(\tilde{s}' - \tilde{s}_2),$$

(E.9b)
to Eq. (E.7),
\[ \tilde{f}(\tilde{s}^1; \tilde{s}^2) = \tilde{w} \tilde{f}(\tilde{s}^1; \tilde{s}^2) + \tilde{h}, \tag{E.10} \]
where \( \tilde{w}_1 = \tilde{f}_{11}/(\tilde{f}_{11} + \tilde{f}_{21}), \) \( \tilde{w}_2 = \tilde{f}_{21}/(\tilde{f}_{11} + \tilde{f}_{21}) \), and \( \tilde{h} = \tilde{f}_{12}/(\tilde{f}_{11} + \tilde{f}_{21}) \), with \( \tilde{f}_{12} = \tilde{f}(\tilde{s}^1; \tilde{s}^2) \) and \( \tilde{f}_{21} = \tilde{f}(\tilde{s}^2; \tilde{s}^1) \).

Here, Eq. (E.10) directly gives the dimorphic invasion-function function \( f(s^1; s^2) \) in the original coordinate system,
\[ f(s^1; s^2) = \tilde{f}(A^{-1}(s^1 - b); A^{-1}(s^2 - b); A^{-1}(s^1 - b)) = \tilde{f}(s^1; s^2). \tag{E.11} \]

In addition, \( \tilde{f}(\tilde{s}^1; \tilde{s}^2) = f(s^1; s^2) \) and \( \tilde{f}(\tilde{s}^2; \tilde{s}^1) = f(s^2; s^1) \) both hold according to Eqs. (E.9a) and (E.9b), which upon substitution into Eq. (E.10) yields Eq. (E.7), as expected. Therefore, in general a dimorphic invasion-function can be approximated by a rational form composed of its first and second derivatives, Eq. (E.7).

Even if the trait space is multivariate with arbitrary dimension \( L \), denoted by \( s = (u_1, ..., u_L)^T \), we can find, for arbitrary \( s^1, s_1, \) and \( s_2, \) a bivariate subspace that contains all three of these phenotypes. In this subspace, Eq. (E.7) holds, where the monomorphic and dimorphic invasion-function functions defined on \( s = (u_1, ..., u_L)^T \) can be directly used as long as we consider \( s^1 \) to be restricted to this subspace. As we can always find such a subspace for any \( s, s_1, \) and \( s_2, \) the dimorphic invasion-function function \( f(s^1; s^2) \) on \( s = (u_1, ..., u_L)^T \) is given by Eq. (E.7), by using the monomorphic function \( f(s^1; s^2) \) on \( s = (u_1, ..., u_L)^T \).

Finally, substituting the monomorphic invasion-function function Eq. (D.12a) in the main text into Eq. (E.7) yields the corresponding dimorphic invasion-function function, Eq. (D.12).

**Derivation of consistency condition**

Here we derive the consistency condition \( f(s^1; s_1, s_2) = f(s^1; s_1) \) for \( f(s^2; s_1) = 0 \) and \( f(s^1; s_2) > 0 \), by proving that \( \hat{n}_2 \to 0 \) holds for \( f(s^2; s_1) \to 0 \) while \( f(s^2; s_2) > 0 \).

First, we assume a protected dimorphism of \( s_1 \) and \( s_2 \), i.e., \( f(s^2; s_1) > 0 \) and \( f(s^1; s_2) > 0 \). This is possible only when the resident phenotypes \( s_1 \) and \( s_2 \) are both in the neighborhood of an evolutionarily singular point in the one-dimensional trait subspace defined by the \( s \)-axis in Eq. (E.8). Then, the population dynamics of \( s_1 \) and \( s_2 \) can be approximated by Lotka–Volterra equations (Durinx et al., 2008), giving a unique interior stable equilibrium \( (\hat{n}_1, \hat{n}_2) \); see also Appendix M. Thus, when \( f(s^2; s_1) \) converges to zero while \( f(s^2; s_2) > 0, \hat{n}_1 \) converges to zero, while \( \hat{n}_2 \to 0 \). In this case, \( f(s^1; s_1, s_2) \) must converge to \( f(s^1; s_1) \). Therefore, \( f(s^1; s_1, s_2) = f(s^1; s_1) \) for \( f(s^2; s_1) = 0 \) and \( f(s^2; s_2) > 0 \).

**Appendix F. Mutational step sizes of MLI mutants**

Here we prove that MLI mutants \( s_{\text{MLI}} \), which maximize Eq. (11) in Section 4, satisfy \( |z_{\text{MLI}}| \leq \sqrt{2} \sigma \), where \( \sigma \) is the parental resident, when invasion-fitmess functions are approximated by quadratic functions of \( s \) (e.g., Eq. (2)) and mutation probability distributions are approximated by multivariate Gaussian functions.

**Main proof**

We consider a multivariate trait space \( s = (u_1, ..., u_L)^T \) with arbitrary dimension \( L \) that is normalized and rescaled so that mutation is isotropic with standard deviation 1. We assume that the mutation probability distribution is approximated by a multivariate Gaussian function
\[ M(\delta s) = \frac{1}{(2\pi)^{L/2}} \exp(-\frac{1}{2} \delta s^T \Sigma \delta s), \tag{F.1} \]

with a standard deviation 1 in all directions.

In a similar manner to Appendix A, we can expand the invasion-fitness function in \( s' \) about \( s_{\text{MLI}} \) as
\[ f(s^1; ..., s_N) = G(\delta s^1 + 1 \delta s^2 + \delta s^3 + \cdots + \delta s^N) + \text{h.o.t.}, \tag{F.2} \]
where \( \delta s^1 = s' - s_{\text{MLI}} \) is an \( L \)-dimensional row vector, and \( D \) is an \( L \times L \) symmetric matrix. Notice that \( G \) and \( D \) are both functions of \( s_1, ..., s_N \) so that \( f(s_1, ..., s_N) = 0 \) holds for each \( i = 1, ..., N \). For example, \( f(s_1, ..., s_0) \) for \( N = 2 \) (i.e., for a dimorphism) is given by Eq. (E.7) in Appendix E. We assume that the higher-order terms in Eq. (F.2) can be neglected.

According to Eq. (11) in Section 4, \( \delta s_{\text{MLI}} = s_{\text{MLI}} - s_{\text{MLI}} \) maximizes
\[ \mu(\delta s^1; s_1, ..., s_N) = \frac{1}{2} \mu\delta s^1 \exp\left(-\frac{1}{2} \delta s^T \Sigma \delta s\right) \tag{F.3} \]
where \( \delta s_{\text{MLI}} = s_{\text{MLI}} - s_{\text{MLI}} \), \( \mu(\delta s^1; s_1, ..., s_N) \) is the \( L \)-dimensional row vector, and \( D \) is an \( L \times L \) symmetric matrix. Notice that \( G(\delta s^1) = 0 \) indicates that \( G \) is a smooth function, with \( G(0) = 0 \), \( G(+\infty) = 0 \), and \( G(0) > 0 \) for \( z < 2G^2/D_z \), \( H(z) \) has a positive maximum for

\[ 0 \leq z_{\text{MLI}} \leq \frac{2G_z}{(-D_z)} \text{ for } D_z < 0. \tag{F.5a} \]

Notice that \( H(z) \) is a smooth function of \( z \). As \( H(0) = 0 \), \( H(+\infty) = 0 \), and \( H(0) > 0 \) for \( z < 2G^2/D_z \), \( H(z) \) has a positive maximum for
Thus, when \( D_2 \geq 0 \), the \( z_{\text{MLI}} \), which satisfies both Eqs. (F.5a) and (F.6), must satisfy \( 1 \leq z_{\text{MLI}} \leq \sqrt{2} \). On the other hand, when \( D_2 < 0 \), Eq. (F.6) can be expressed as

\[
\frac{dH(z)}{dz} = A_0 \exp\left(-\frac{jx^2}{2}\right) \left[G_0 (1-z^2) + \frac{1}{2} D_2 z(2-z^2)\right] = 0.
\]

This completes the proof.

**Appendix G. Proof of Lemma 1 (conditions for dimorphic emergence)**

Here we prove Lemma 1 in Section 5, specifying conditions on MLI mutants to ensure dimorphic emergence. While this lemma refers to MLI mutants, it also holds for non-MLI mutants as long as they satisfy the conditions for dimorphic emergence, inequalities (13). Thus, here we do not distinguish between \( s' \) and \( s_{\text{MLI}} \) and denote mutants simply by \( s \).

We consider a monomorphic resident \( s = (x,y)^T \). A sufficient condition for protected dimorphism of \( s \) and \( s' \), and thus for dimorphic emergence, is given by mutual invasibility,

\[
f(s'; s) = -\dot{y} + D\delta x^2 + Cx\delta y > 0,
\]

\[
f(s; s') = -\dot{y} + D\delta x^2 - C(x+\delta x)\delta y > 0.
\]

These inequalities can be combined into

\[
0 < \dot{y} + D\delta x^2 + Cx\delta y < (2D-C)\delta x^2.
\]

We first suppose that the population is not close to the convergence-stable line \( x = 0 \), so that \( |x| \) is significantly larger than 1. Since the magnitudes of \( \delta x \) and \( \delta y \) are of order 1, \( (2D-C)\delta x^2 \approx |Cx\delta y| \) holds. In this case, inequalities (G.2) cannot hold. This means that invasion by \( s' \) always replaces \( s \), which corresponds to directional evolution. To satisfy inequalities (G.2), the population has to come close to the convergence-stable line through directional evolution, such that \( |x| \) becomes sufficiently small. Such convergence is ensured if all invading mutants satisfy

\[
\begin{align*}
\{ x &> x \quad \text{for} \quad x < 0, \\
&< x \quad \text{for} \quad x > 0.
\}
\end{align*}
\]

This implies \( \dot{\delta} x > 0 \) for \( x < 0 \) and \( \delta x < 0 \) for \( x > 0 \). The population monotonically converges to the convergence-stable line if all invading mutants satisfy this condition, as long as the resident is monomorphic.

On this basis, we now suppose that the population has come close to the convergence-stable line and that a mutant has arisen such that the resident and the mutant straddle the convergence-stable line, \( xx' < 0 \). As \( C \) is negative, both \( Cx\delta x = -C(-xx'+x^2) \) and \( -C(x+\delta x)\delta y = -C(x^2-xx') \) are always positive. Thus, inequalities (G.2) hold if

\[
x'x < 0
\]

and

\[
\frac{|\delta y|}{\delta x} < D.
\]

Since directional evolution proceeds toward the convergence-stable line under inequalities (G.3), the situation \( xx' < 0 \) inevitably occurs unless protected dimorphism emerges even before that. Thus, for an arbitrary initial resident \( s_0 \), if all subsequent invading mutants satisfy inequalities (G.3) and (G.5), then the population monotonically converges to the line \( x = 0 \), until protected dimorphism has emerged, which inevitably occurs once \( s \) and \( s' \) straddle the line. Inequalities (G.3) and (G.5) are identical to inequalities (13) in Section 5. This completes the proof.

**Appendix H. Proof of Lemma 2 (conditions for dimorphic divergence)**

**Preparation**

Here we prove Lemma 2 in Section 5, specifying conditions on MLI mutants to ensure dimorphic divergence. Similarly to Lemma 1, Lemma 2 also holds for non-MLI mutants as long as they satisfy the conditions for dimorphic divergence, inequalities (14). Thus, as in Appendix G, here we do not distinguish between \( s' \) and \( s_{\text{MLI}} \), and denote mutants simply by \( s' \).

After the emergence of a protected dimorphism, composed of two residents denoted by \( s_1 \) and \( s_2 \), the next invasion event occurs based on the dimorphic invasion fitness \( f(s'; s_1, s_2) \). As long as the two residents remain close to the base point of expansion, \( s_0 = (0,0)^T \), this dimorphic invasion fitness is approximately given by Eq. (D.12),

\[
f(s'; s_1, s_2) = \left( y'-y_2 \right) + D(x'-x_1)(x'-x_2) - \frac{y_2-y_1}{x_2-x_1},
\]

as shown in Appendix E. This dimorphic invasion-fitness function \( f(s'; s_1, s_2) \) and the monomorphic invasion-fitness function \( f(s'; s) \) given by Eq. (12a) together determine whether a sequence of invading mutants can bring about dimorphic divergence.

**Conditions for a single step of dimorphic divergence**

We define dimorphic divergence as the directional evolution of two resident morphs in opposite directions along the \( x \)-axis. Such a compound evolutionary process is formed by repetition of a unit process defined as follows.

**Definition of a single step of dimorphic divergence.** An invading mutant replaces only either of the two residents and coexists with the other resident, and the phenotypic distance along the \( x \)-axis between the new residents is larger than that between the old residents.

For a more formal description, we now consider arbitrary phenotypes \( s', s_1 \), and \( s_2 \). The resident that is replaced by \( s' \) is denoted by \( s_p \) (i.e., \( s_p = s_1 \) or \( s_p = s_2 \)). The other resident that coexists with \( s' \) is denoted by \( s_q \) (i.e., \( s_q = s_1 \) or \( s_q = s_2 \)). Although \( s' \) usually replaces its parental resident (i.e., \( s_p \) is its parental resident), \( s' \) may replace the other non-parental resident (i.e., \( s_p \) is the non-parental resident) when both the two residents are close to each other. Then the definition above of a single step of dimorphic divergence is fulfilled under the following conditions.

**Sufficient conditions for a single step of dimorphic divergence.**

(a) \( f(s'; s_p, s_q) > 0 \) (\( s' \) can invade)

(b) \( f(s_0; s') > 0 \) and \( f(s'; s_1) > 0 \) (\( s' \) coexists with \( s_1 \), owing to mutual invasibility)

(c) \( f(s_p; s_q, s') < 0 \) (\( s' \) excludes \( s_p \))

(d) \( |x'-x_1| > |x_2-x_1| \) (phenotypic divergence along the \( x \)-axis becomes larger)

On this basis, we prove the following lemma in Appendix I.

**Lemma H.1.** Sufficient conditions for satisfying conditions (a) to (d) above are given by

\[
|y_2-y_1| < D(x_2-x_1)^2,
\]

(H.2a)
\((x_0 - x_2)dx > 0, \quad (H.2b)\)

and

\[|\delta y| < D\delta x^2, \quad (H.2c)\]

where \((\delta x, \delta y)^T = (x' - x_0, y' - y_0)^T = s' - s_0\).

Therefore, for dimorphic residents \(s_1\) and \(s_2\) satisfying inequality (H.2a), any mutant satisfying inequalities (H.2b) and (H.2c) ensures a single step of dimorphic divergence.

The set of mutants \(s'\) that satisfy inequalities (H.2b) and (H.2c) is illustrated as the white regions in Fig. 3d.

Conditions for the whole process of dimorphic divergence

We suppose that an invading mutant \(s'\) and residents \(s_1\) and \(s_2\) satisfy inequalities (H.2). Without loss of generality, we assume \(s_p = s_2\) and \(s_0 = s_1\), differently from Lemma 2, which assumes \(x_1 < x_2\) instead. Then \(s'\) excludes only \(s_2\), and coexists with \(s_1\). In addition, as proved in Appendix I, the new dimorphism composed of \(s'\) and \(s_1\) satisfies

\[|y' - y_1| < D(x' - x_1)^2. \quad (H.3)\]

When \(s'\) is renamed as the replaced resident \(s_2\), inequality (H.3) gives inequality (H.2a). Thus, for the next step of dimorphic divergence, only inequalities (H.2b) and (H.2c) have to hold, and the same applies for subsequent steps of dimorphic divergence. Therefore, for any initial dimorphic residents satisfying inequality (H.2a), the whole process of subsequent dimorphic divergence is ensured, if all of the subsequent invading mutants satisfy inequalities (H.2b) and (H.2c). In addition, any initial protected dimorphism emerged under the conditions for dimorphic emergence, inequalities (13) in Section 5, clearly satisfies inequality (H.2a). Thus, provided that the initial dimorphism has emerged under the conditions for dimorphic emergence, sufficient conditions on the subsequent invading mutants for dimorphic divergence are given by inequalities (H.2b) and (H.2c). Inequalities (H.2b) and (H.2c) are equivalent to inequalities (14) in Section 5 when \(x_1 < x_2\) is assumed. This completes the proof.

Appendix I. Proof of Lemma H.1 (conditions for a single step of dimorphic divergence)

Here we prove Lemma H.1 in Appendix H. We also show that the conditions for a single step of dimorphic divergence, inequalities (H.2), ensure inequality (H.3).

Main proof

Without loss of generality, we assume that \(s_p = s_2\) and \(s_0 = s_1\), differently from Lemma 2, which assumes \(x_1 < x_2\). Then the conditions (a) to (d) in Appendix H, for a single step of dimorphic divergence, become \(f(s'; s_1, s_2) > 0, f(s_1; s_1) > 0, f(s; s_1, s') < 0, \) and \(|x' - x_1| > |x_1 - x_1|\). We denote the phenotypic differences of the original and the new residents by \(\Delta s = s_0 - s_1 = s_2 - s_1\), i.e., \((\Delta s, \Delta y)^T = (x_1 - x_1, y_1 - y_1)^T\), and \(\Delta s' = s_p - s_1\), i.e., \((\Delta s', \Delta y')^T = (x_1 - x_1, y_1 - y_1)^T\), respectively. We also define \(\tan \phi = \Delta y/\Delta x\) and \(\tan \phi' = \Delta y'/\Delta x'\).

On this basis, we prove the following two lemmas in the subsequent subsections.

**Lemma 1.1.** All inequalities \(f(s'; s_1, s_2) > 0, f(s_1; s_1) > 0, f(s; s_1, s') < 0, \) and \(|x' - x_1| > |x_1 - x_1|\) hold if \(\Delta x > 0\) and \(|\Delta x| < |\Delta x'|\)

and

\[|\tan \phi' - \tan \phi| < D|\Delta x'| - |\Delta x|. \quad (l.1b)\]

**Lemma I.2.** Inequalities (l.1) hold if inequalities (H.2) hold.

By these lemmas, the proof is completed.

**Proof of Lemma 1.1**

We assume that inequalities (l.1) hold. Inequalities (l.1a) immediately give \(|x' - x_1| > |x_2 - x_1|\). As for the signs of the inversion fitnesses, dividing \(f(s'; s_1, s_2)\) by \(\Delta x'^2 - \Delta x\) yields

\[f(s'; s_1, s_2) = \left(\frac{y' - y_2}{x' - x_2} - \frac{y_1 - y_1}{x_1 - x_1}\right) + D \frac{[(x_1 - x_1) - (x_2 - x_1)]}{(x' - x_2 - x_1)}\]

\[= \tan \phi' - \tan \phi + D\Delta x'. \quad (l.2a)\]

Similarly, \(f(s'; s_1), f(s_1; s'), f(s_1; s_1)'\) and \(f(s_1; s, s')\) satisfy the following equations,

\[f(s_1; s') = - \tan \phi' - \tan \phi + D\Delta x' - C(\Delta x' - \Delta x) + f(s_1; s_1'), \quad (l.2b)\]

and

\[f(s_1; s, s') = -(\tan \phi' - \tan \phi - D\Delta x'). \quad (l.2d)\]

When \(\Delta x > 0\), Eq. (l.1a) gives \(\Delta x' > 0\) and \(\Delta x' - \Delta x > 0\). As \(D > 0\) and \(C < 0\), and \(f(s_1; s_1) > 0\) and \(f(s; s_1, s') < 0\), all inequalities \(f(s'; s_1, s_2) > 0, f(s'; s_1) > 0, f(s; s_1, s') > 0, \) and \(f(s_1; s, s') < 0\) hold under inequality (l.1b), according to Eq. (l.2). When \(\Delta x < 0\), Eq. (l.1a) gives \(\Delta x' < 0\) and \(\Delta x' - \Delta x < 0\). Thus, in the same manner, all inequalities \(f(s'; s_1, s_2) > 0, f(s'; s_1) > 0, f(s; s_1, s') > 0, \) and \(f(s_1; s, s') < 0\) hold under inequality (l.1b). This completes the proof.

**Proof of Lemma I.2**

We assume that inequalities (H.2) hold. Inequality (H.2.b) can be expressed as \((x_2 - x_1)\delta x = \Delta x'\Delta x' - \Delta x\Delta x' = 0\), which gives inequalities (1.1a). In addition, inequalities (1.1a) give \(|\delta x| + |\Delta x| = |\Delta x'|\). Thus, inequalities (H.2a) and (H.2c) yield inequality (1.1b).

\[|\tan \phi' - \tan \phi| = \frac{|\delta y + \Delta y|}{\frac{\Delta x}{\Delta x'}} = \frac{|\Delta x\delta y - \Delta x\Delta y|}{\Delta x'\Delta x} \leq \frac{[|\Delta x\delta y| + |\Delta x\Delta y|]}{\Delta x'} \leq \frac{D\Delta x^2 |\Delta x| + D|\Delta x|\Delta x^2}{\Delta x'\Delta x} = D|\Delta x| = D|\Delta x'| = D|\Delta x' - \Delta x|. \quad (1.3)\]

This completes the proof.

**Proof of inequalities (H.2) ensuring inequality (H.3)**

We assume that inequalities (H.2) hold. In this case, inequality (H.2b) gives \(\Delta x\delta x > 0\). Then, inequalities (H.2a) and (H.2b) yield inequality (H.3),

\[|\Delta y'| = |\delta y| + |\Delta y| < D\delta x^2 + D\Delta x^2 = D\delta x + D\Delta x^2 \leq D\delta x + D\Delta x^2 = D\Delta x^2. \quad (1.4)\]

This completes the proof.
Appendix J. Directional evolution sufficient for evolutionary branching

Here we derive Eq. (19) in Section 5, specifying the sufficient amount of directional evolution along \( y \) for the emergence of a dimorphism of \( s_1 \) and \( s_2 \) satisfying \( |\Delta x| = |x_2 - x_1| > \Delta x^* \) for an arbitrary \( \Delta x^* > 0 \). Provided that inequality (18), \( D > 1/\sqrt{2} \), holds, Lemmas D.1 and D.2 in Appendix D ensure that

\[
\begin{align*}
\delta x_{\text{MLI}} &> 0 \quad \text{for} \quad x < 0 \\
\delta y_{\text{MLI}} &< 0 \quad \text{for} \quad x > 0
\end{align*}
\]

(J.1a)

for a monomorphic resident population, and that

\[
\begin{align*}
\delta x_{\text{MLI}} &< 0 \quad \text{for} \quad s_{\text{MLI}} = s_1 \\
\delta y_{\text{MLI}} &> 0 \quad \text{for} \quad s_{\text{MLI}} = s_2
\end{align*}
\]

(J.1b)

for a dimorphic resident population, where \((\delta x_{\text{MLI}}, \delta y_{\text{MLI}})^T = (s_{\text{MLI}} - s_1, s_2 - s_1)\) and \(s_{\text{MLI}}\) is the parental resident of \(s_k\), and \(x_1 < x_2\) is assumed without loss of generality. Also, for both monomorphic and dimorphic populations, Lemmas D.1 and D.2 ensure that

\[
\begin{align*}
1 < \delta x_{\text{MLI}} < \sqrt{2}, \\
0 < \delta y_{\text{MLI}} < \frac{1}{\sqrt{2}}
\end{align*}
\]

(J.2a) and (J.2b)

Then, under inequalities (J.1) and (J.2), the process of evolutionary branching along an MLIP amounts to monotonic monomorphic convergence toward \( x = 0 \), followed by monotonic dimorphic divergence after the emergence of a dimorphism. Thus, if \( x_2 \) is negative, this evolutionary dynamics in the trait space traces the shape of a lower-case letter “y”. By contrast, if \( x_2 \) is positive, the shape is that of a mirrored “y”.

The sum of the lengths of all branches and trunks of such a tree shape is given by

\[
L(I_{\text{MLI}}; s_1) = \sum_{k=1}^{K} |\delta x_{\text{MLI}}(k)| = \sum_{k=1}^{K} |\delta y_{\text{MLI}}(k)|,
\]

where \( s_{\text{MLI}}(k) = (x_{\text{MLI}}(k), y_{\text{MLI}}(k))^T \) is the MLI mutant invading at the \( k \)-th invasion event, \( s_{\text{MLI}}(k) = (x_{\text{MLI}}(k), y_{\text{MLI}}(k))^T \) is its parental resident, \( K \) is the total number of invasion events, \( I_{\text{MLI}} = (s_{\text{MLI}}(1), ..., s_{\text{MLI}}(k), ..., s_{\text{MLI}}(K)) \), and \( s_1 \) is the initial resident phenotype. Eq. (J.3) describes the length of the evolutionary path formed by the mutant-invasion sequence \( I_{\text{MLI}} \) from \( s_1 \). By projecting this evolutionary path onto the \( x \)- and \( y \)-axes, we obtain its lengths along \( x \) and \( y \) as

\[
\begin{align*}
L_x(I_{\text{MLI}}; s_1) &= \sum_{k=1}^{K} |\delta x_{\text{MLI}}(k)| = \sum_{k=1}^{K} |x_{\text{MLI}}(k) - x_{\text{MLI}}(k)|, \\
L_y(I_{\text{MLI}}; s_1) &= \sum_{k=1}^{K} |\delta y_{\text{MLI}}(k)| = \sum_{k=1}^{K} |y_{\text{MLI}}(k) - y_{\text{MLI}}(k)|.
\end{align*}
\]

(J.4)

respectively. We furthermore decompose \( L_x \) into the portions before and after dimorphic emergence,

\[
L_x(I_{\text{MLI}}; s_1) = \sum_{k=1}^{K_x-1} |\delta x_{\text{MLI}}(k)| + \sum_{k=K_x}^{K} |\delta x_{\text{MLI}}(k)| = L_x + l_{x}. \]

(J.5)

where the \( K_x \)-th invasion event is assumed to bring about the emergence of dimorphism.

Under inequalities (J.1) and (J.2), the population monotonically converges toward the evolutionary-branching line until it becomes dimorphic, which inevitably occurs before the resident and mutant straddle the line \( x = 0 \). In other words, the population starts diversification before the length of its evolutionary path along \( x \) exceeds \( |x_2| \). Thus,

\[
L_x \leq |x_2|.
\]

(J.6a)

In addition, monotonic diversification along \( x \) continues after the emergence of dimorphism, with

\[
L_{xD} = |x_2(K) - x_1(K)|.
\]

(J.6b)

Moreover, as \( |\delta y_{\text{MLI}}|/|\delta x_{\text{MLI}}| < 1/\sqrt{2} \) always holds under inequalities (J.2),

\[
L_y(I_{\text{MLI}}; s_1) = \sum_{k=1}^{K} |\delta y_{\text{MLI}}(k)| < \frac{1}{\sqrt{2}} \sum_{k=1}^{K} |\delta x_{\text{MLI}}(k)|
\]

\[
= \frac{1}{\sqrt{2}}(L_x + L_{xD}) \leq \frac{1}{\sqrt{2}}(|x_2| + |x_2(K) - x_1(K)|)
\]

(J.7)

holds. In addition, obviously,

\[
y(K) \leq \max(y_1(K), y_2(K)) \leq L_y(I_{\text{MLI}}; s_1) + y_a,
\]

(J.8)

where \( y(K) \) is the mean value of \( y \) after \( K \) invasion events, given by

\[
y(K) = \frac{1}{K} \sum_{k=1}^{K} y_{\text{MLI}}(k)
\]

and by \( y_1(K) = (K_1y_1(K_1) + K_2y_2(K_2))/(K_1 + K_2) \) for dimorphism. Substituting inequality (J.8) into inequality (J.7) yields

\[
y(K) - y_a < \frac{|x_2| + |x_2(K) - x_1(K)|}{\sqrt{2}}
\]

(J.9)

Thus, for the emergence of dimorphism with \( |x_2(K) - x_1(K)| > \Delta x^* \), Eq. (19) in Section 5,

\[
y - y_a < \frac{|x_2| + \Delta x^*}{\sqrt{2}}
\]

(J.10)

is the sufficient amount of directional evolution, where \( y = y(K) \). Analogously, the sufficient number of invasion events can be derived as \( (|x_2| + \Delta x^*) \).

Appendix K. Procedures for the numerical calculation of evolutionary dynamics

Here we explain the procedures for the numerical calculation of the evolutionary dynamics shown in Section 6. These calculations are conducted in a normalized and rescaled trait space such that mutation is isotropic with standard deviation 1. For calculating MLIPs, the MLI mutant at each invasion event is determined so that it maximizes the invasion-event probability density defined in Eq. (9b) in Section 4. For calculating OSIPs, each invading mutant is stochastically chosen according to the invasion-event probability density (see also Ito and Dieckmann, 2007). See Appendix M for the details of how to calculate invasion-event probability densities. When an invasion has occurred, the coexisting phenotypes at the next population dynamical equilibrium are determined by checking invasion fitnesses among residents and the mutant.

For each calculation of an MLIP or OSIP, the trait \( x_{\text{MLI}} \) of the initial resident is drawn randomly from a uniform distribution with \( x_{\text{MLI}} \leq 10 \), while the trait \( y_a \) of the initial resident is set to 0 without loss of generality. For evaluating the occurrence of evolutionary branching in OSIPs, it is numerically observed that 99.997% of failures (i.e., collapse of protected dimorphisms) occur for \( |x_1| < 10 \), where \( \Delta x = x_2 - x_1 \) describes the phenotypic difference in \( x \) between the two residents. Thus, we conclude that evolutionary branching has occurred when a dimorphism with \( |x_1| > 10 \) has emerged. Then, the sufficient directional evolution in \( y \) along MLIPs is given by \( L_y = L_{\text{MLI}}(x_1, 10) = (|x_1| + 10)/\sqrt{2} \), according to inequality (19) in Section 5.
We calculate PSIPs using the polymorphic stochastic model (Dieckmann and Law, 1996), which describes individual births and deaths as stochastic events. For illustration, we use the birth and death rates defined for the resource-competition model studied by Ito and Dieckmann (2007), which is a linear combination of the MacArthur-Levins resource-competition model (MacArthur, 1972) in x and a constant selection gradient in y. This model (detailed in Appendix N) is a simple but ecologically plausible realization of the normal form for invasion-fitness functions with significant sensitivity difference considered in this study, as given by Eq. (3). The initial monomorphic phenotype is assigned as described for OSIPs above. To examine the process of evolutionary branching, phenotypes whose phenotypic distance is less than 2 are clustered together. When an initial single cluster splits into two clusters, |Δx| is calculated as the phenotypic distance between the averages of x within the two clusters. We conclude that evolutionary branching has occurred when |Δx| exceeds 10, analogous to the criterion used for OSIPs, as described above.

Appendix L. Mutation distributions

Here we specify three additional mutation distributions used in our numerical calculations, which are the bivariate fixed-step distribution, the univariate Gaussian distribution, and the univariate fixed-step distribution.

The bivariate fixed-step distribution describes mutations that are limited to an ellipse \( \delta X^2 / \sigma_X^2 + \delta Y^2 / \sigma_Y^2 = 1 \), as illustrated in Fig. 5b. Along this ellipse, mutations are distributed uniformly. The standard deviations of the mutational step sizes, calculated according to Eq. (1) in Section 3, are \( \sigma_X = \sqrt{V_{XX} - \delta X/2} \) and \( \sigma_Y = \sqrt{V_{YY} - \delta Y/2} \), with \( V_{XY} = 0 \).

The univariate Gaussian distribution describes separate mutations in X and Y, with the corresponding relative mutation rates given by \( \mu_X \) and \( \mu_Y \), respectively, as illustrated in Fig. 5c. The mutational step sizes follow Gaussian distributions with standard deviations \( \delta X \) and \( \delta Y \), respectively. The standard deviations of mutational step sizes, calculated according to Eq. (1) in Section 3, are \( \sigma_X = \sqrt{\mu_X^2 / \delta X_X} \) and \( \sigma_Y = \sqrt{\mu_Y^2 / \delta Y_Y} \), with \( V_{XY} = 0 \).

The univariate fixed-step distribution describes separate mutations in X and Y, with the corresponding relative mutation rates given by \( \mu_X \) and \( \mu_Y \) respectively (\( \mu_X + \mu_Y = 1 \)). The only possible mutations are \( \pm \delta X \) and \( \pm \delta Y \). The only possible mutations are \( \pm \delta X \) and \( \pm \delta Y \), which occur with probabilities \( \mu_X / \mu_Y \), \( \mu_Y / \mu_X \), and \( \mu_X / \mu_Y \), as illustrated in Fig. 5d. The standard deviations of mutational step sizes, calculated according to Eq. (1) in Section 3, are \( \sigma_X = \sqrt{\mu_X^2 / \delta X_X} \) and \( \sigma_Y = \sqrt{\mu_Y^2 / \delta Y_Y} \), with \( V_{XY} = 0 \).

After normalizing and rescaling the trait space \( S = (X, Y) \) into \( s = (x, y) \) such that the standard deviations of mutational step sizes become equal to 1 in all directions, the MLIP condition is applied. MLIP conditions for the additional three mutation distributions described above are applied by using the \( \sigma = \sigma_X = \sigma_Y \) for these distributions in Eqs. (12c) and (18). For clarity, here we refer to these conditions as approximate MLIP conditions (as they were derived for bivariate Gaussian distributions, but are now applied to other distributions).

Alternatively, MLIP conditions for the three other distributions can be derived directly by using these mutation distributions in the invasion-event probability function, analogously to the derivation for bivariate Gaussian distribution, as defined by Eq. (15) in Section 5. If the X- and Y-axes correspond to the sensitive and insensitive directions, respectively, then the resultant exact conditions are obtained as \( \sqrt{\sigma_X^2 / \mu_X^2 + \sigma_Y^2 / \mu_Y^2} > \sqrt{e} / 2 \), and \( \mu_X \sigma_Y / \sqrt{2 \mu_X^2 \sigma_Y^2} > \sqrt{e} / 2 \). These results can be compared with \( D = (\sigma_X^2 / \mu_X^2 + \sigma_Y^2 / \mu_Y^2) > 1 / \sqrt{2} \) for the bivariate Gaussian distribution.

As it turns out, the exact MLIP conditions tend to overestimate the likelihood of evolutionary branching in OSIPs and PSIPs when univariate fixed-step or univariate Gaussian mutations with large \( \mu_X \) or \( \mu_Y \) are considered (results not shown). Thus, using the approximate MLIP conditions seems to be more robust than using the exact ones.

Appendix M. Calculation of invasion-event probability densities in MLIPs and OSIPs

 Necessary elements for calculation

Here we explain how invasion-event probability densities, as defined by Eq. (9b) in Section 4, are determined in the calculation of MLIPs and OSIPs. For any given composition of resident phenotypes, their equilibrium frequencies and the invasion fitness of possible mutants are required for the calculation (absolute population sizes are needed only for determining the waiting time for each invasion event, which is not needed for the numerical results we present in this study).

As for invasion-fitness functions, we have those for monomorphism, Eq. (12a) in Section 5, and for dimorphism, Eq. (D12) in Appendix D. Invasion-fitness functions for higher degrees of polymorphism are not needed, because trimorphism is impossible under the dimorphic invasion-fitness function, Eq. (D12). Thus, the monomorphic and dimorphic invasion-fitness functions are sufficient for the calculation of invasion-event probability densities.

For equilibrium frequencies of resident phenotypes, the frequency of a monomorphic resident is of course always 1. For a dimorphism of \( s_1 \) and \( s_2 \), the corresponding frequencies are approximately given by \( q_1 = f(s_1, s_2) / f(s_1, s_1) + f(s_2, s_1) \) and \( q_2 = 1 - q_1 \), as explained in the next subsection.

Equilibrium frequencies of dimorphic phenotypes

Here we approximate the equilibrium frequencies of the dimorphic residents \( s_1 \) and \( s_2 \). Without loss of generality, we consider a normalized but non-scaled trait space, so that mutation is isotropic with standard deviation \( \sigma = 1 \).

As explained in Appendix E, the population dynamics of \( s_1 \) and \( s_2 \) can be approximated by Lotka–Volterra equations,

\[
\frac{1}{n_1} \frac{dn_1}{dt} = r_1 \left( 1 - \frac{n_1 + \alpha_1 n_2}{K_1} \right),
\]

\[
\frac{1}{n_2} \frac{dn_2}{dt} = r_2 \left( 1 - \frac{n_2 + \alpha_2 n_1}{K_2} \right). \tag{M.1}
\]

where \( r_2 - r_1, \alpha_1 - \alpha_2, \) and \( K_2 - K_1 \) are \( O(s_2 - s_1) \). The corresponding equilibrium population sizes are given by

\[
\hat{n}_1 = K_1 / (1 - \alpha_2 K_2), \quad \hat{n}_2 = K_2 / (1 - \alpha_1 K_1). \tag{M.2}
\]

while \( f(s_1, s_1) \) and \( f(s_1, s_2) \) are given by

\[
f(s_1, s_2) = \frac{r_1}{K_1} \left[ 1 - \alpha_2 K_2 \right], \quad f(s_1, s_1) = \frac{r_2}{K_2} [1 - \alpha_1 K_1]. \tag{M.3}
\]

Then, the following relationship holds,

\[
\frac{\hat{n}_1 + \hat{n}_2}{f(s_1, s_2) + \frac{r_1}{K_1} f(s_1, s_1)} = \frac{f(s_1, s_2)}{f(s_1, s_1)} \left( 1 + \epsilon \frac{n_2}{n_1} \right) = \frac{1}{1 + \epsilon n_2 / n_1}, \quad \epsilon = \frac{w_1 - \epsilon w_1 w_2 + O(\epsilon^2)}{w_1 (1 - \epsilon w_2 + O(\epsilon^2))}, \tag{M.4}
\]
where $w_1 = (f(s; s_1) - f(s; s_2) - d(s; s_1))(f(s; s_1) + f(s; s_2))$, $w_2 = 1 - w_1$, and $\varepsilon = (r_1K_2)/(r_1K_2 - 1) = \Omega(s_2 - s_1)$. Notice that $f(s; s_2) > 0$ and $f(s; s_1) > 0$ both hold for a protected dimorphism of $s_1$ and $s_2$. Thus, as long as $|s_2 - s_1| = O(\varepsilon)$,

$$
\hat{n}_1 + \hat{n}_2 = w_1 + O(\varepsilon)
$$

is a good approximation. Therefore, after rescaling this trait space such that the standard deviation of mutational step sizes is equal to 1, Eq. (M.5) is a good approximation as long as $|s_2 - s_1|$ is of order 1.

### Appendix N. Specific model for calculation of PSIPs

Here we explain the model used for the calculation of PSIPs in Section 6. We consider a normalized bivariate trait space $s = (x, y)^T$, in which mutation is isotropic with standard deviation $\sigma = 1$. We define individual birth and death rates following Ito and Dieckmann (2007), as explained below. The trait $x$ affects the death rate through resource competition (as, e.g., when beak size in birds determines the size of seeds they compete for). The death rate $d(s; s_1, ..., s_n)$ of phenotype $s = (x, y)^T$ depends on the trait values $x_i$ and $y_i$, as well as on the abundances $n_i$ of extant phenotypes $i = 1, ..., N$.

$$
d(s; s_1, ..., s_N) = \sum_{j=1}^{N} \frac{\alpha(x_i - x_j) n_j}{K(x_j)}.
$$

Here,

$$
K(x) = K_0 \exp\left(-\frac{1}{2}\sigma_x^2 / \sigma_0^2\right)
$$

is the carrying capacity of phenotype $s$, given by a Gaussian function with variance $\sigma_0^2$ and mean $x_0 = 0$. The function $\alpha(x_i - x_j) = \exp\left(-\frac{1}{2}(x_i - x_j)^2 / \sigma_x^2\right)$ describes the strength of competition between phenotype $x_i$ and phenotype $x_j$; it is also given by a Gaussian function, with variance $\sigma_x^2$ and mean $x_i - x_j = 0$. Accordingly, the strength of competition is maximal between identical phenotypes and monotonically declines with phenotypic distance. If the birth rate $b(s; s_1, ..., s_N)$ is assumed to be constant and equal to 1, the birth and death rates imply the MacArthur–Levins resource competition model (MacArthur, 1972).

$$
\frac{\delta n_i}{\delta t} = [b(s; s_1, ..., s_N) - d(s; s_1, ..., s_N)] n_i
$$

for $i = 1, ..., N$, in the limit of infinite population size.

Directional selection on $y$ can be seen as any ecological interaction (e.g., competition, predator-prey interaction, or mutualism) and may act on any morphological, physiological, or life-history trait $y$. A simple way of introducing a fitness gradient in $y$ is

$$
b(s; s_1, ..., s_N) = 1 + b_1(y_i - \bar{y}),
$$

where $\bar{y}$ denotes the population average of trait value $y$, $\bar{y} = \frac{\sum y_i n_j}{\sum n_i}$, and $b_1$ is a constant describing the constant directional selection pressure on $y$.

We now consider a monomorphic resident $s$ with sufficiently large population size. The invasion fitness of $s'$ with respect to $s$ is given by

$$
f(s'; s) = b(s'; s) - d(s'; s) = 1 - \frac{\alpha(x - x')(\hat{n}(x'))}{K(x')} + b_1(y' - y)
$$

$$
= 1 - \frac{\alpha(x - x')K(x)}{K(x')} + b_1(y' - y),
$$

where $\hat{n} = K(x)$ is the equilibrium population size of $s$. Since trait $y$ contributes to invasion fitness only through the linear term $b_1(y' - y)$, the condition for significant sensitivity difference, Eq. (4) in Section 3, is immediately satisfied whenever $b_1$ is sufficiently small such that $C_y = b_1 = O(\varepsilon)$. In this case, the invasion fitness can be expanded about $s_0 = (0, y_0)^T$ with arbitrary $y_0$, in the form of Eq. (3) in Section 3,

$$
f(s'; s) = C_x \delta x + C_y \delta y + \frac{1}{2} D_{xx} \delta^2 x + \frac{1}{2} D_{yy} \delta^2 y,
$$

where

$$
C_x = \frac{\partial^2 f(s'; s)}{\partial x^2} |_{s = s_0} = 0,
$$

$$
C_y = \frac{\partial^2 f(s'; s)}{\partial y^2} |_{s = s_0} = b_1,
$$

and

$$
C_{xy} = \frac{\partial^2 f(s'; s)}{\partial x \partial y} |_{s = s_0} = \frac{1}{\sigma_y^2}
$$

$$
D_{xx} = \frac{\partial^2 f(s'; s)}{\partial x^2} |_{s = s_0} = \frac{1}{\sigma_x^2}
$$

Since $C_y = 0$ and $C_{xy} < 0$ hold, $x = 0$ is a convergence-stable line. Then $C$ and $D$, in the rescaled trait space, are given by

$$
C = \frac{\sigma_{C_y}}{\sigma_{C_x}} = \frac{\sigma_y}{\sigma_x}
$$

$$
D = \frac{\sigma_{D_{xx}}}{\sigma_{D_{yy}}} = \frac{1}{\sigma_x^2 (1 - \sigma_x^2 / \sigma_y^2)}.
$$

As $C$ is always negative, the convergence-stable line is an evolutionary-branching line if the MLIP condition, $D > 1 / \sqrt{2}$, holds.

### Appendix O. Extension of conditions for evolutionary-branching lines to multivariate trait spaces

Here we extend the conditions for evolutionary-branching lines – Eqs. (4), (6a), (6b), and (18) – to multivariate trait spaces. We consider an arbitrary $L$-variate trait space $S = (U_1, ..., U_L)^T$ with a mutational variance–covariance matrix $\Sigma$, which has real and non-negative eigenvalues, $\sigma_1^2, ..., \sigma_L^2$. The maximum eigenvalue of $\Sigma$, denoted by $\sigma_1^2$, gives the maximum standard deviation $\sigma_1$ of mutational step sizes along all directions. The trait space is normalized by an affine coordinate transformation of $S$ into $S' = (u_1, ..., u_L)^T$ with isotropic mutation $\sigma = \sigma_1$ (see Appendix P).

The invasion fitness in the normalized space can be written analogously to the bivariate case (Appendix A) as

$$
f(s'; s) = G s' + (s - s_0)^T G s' + \frac{1}{2} \sum_{i=1}^L \sigma_i^2 \delta^2 s_i + O(\sigma^3),
$$

where $G$ is a row vector of length $L$, and $C$ and $D$ are $L \times L$ matrices, with $D$ being symmetric. In a manner similar to the bivariate case, we can define

**Significant sensitivity difference in multivariate trait spaces.**

After normalization to make mutation isotropic, the normalized invasion-fitness function, Eq. (O.1), can be made to satisfy

$$
\frac{\partial^2 f(s'; s)}{\partial U_i^2} + \frac{1}{\sigma_i^2} + \frac{\partial^2 f(s'; s)}{\partial U_j^2} + \frac{1}{\sigma_j^2} + \frac{\partial^2 f(s'; s)}{\partial U_i \partial U_j} = O(\sigma)
$$

for all $i = 1, ..., L$ and all $j = L + 1, ..., L$, by rotating the axes, where $\tilde{L} < L$, $G_i$ is the $j$th component of $G$, and $C_{ij}$ and $D_{ij}$ are the $(i, j)$ components of $C$ and $D$, respectively.

If the invasion-fitness function, Eq. (O.1), has significant sensitivity difference, the trait space can be decomposed into an $L$-variate sensitive subspace $x = (x_1, ..., x_L)^T = (u_{L+1}, ..., u_L)^T$ and an $\tilde{L}$-variate insensitive subspace $y = (y_1, ..., y_{\tilde{L}})^T = (u_1, ..., u_{\tilde{L}})^T$. Notice that
Eq. (O.2) allows decomposition of Eq. (O.1) into \( f(s'; s) = g(x'; x) + e h(s'; s) \) with a small \( e = O(\sigma) \), while \( h(s'; s) \) is kept smooth and finite, as in the bivariate case (Appendix B). In this case, Eq. (O.1) is transformed into

\[
f(s'; s) = \sum_{i=1}^{\lambda} G_i \delta x_i + \sum_{i=1}^{\lambda} \sum_{i=1}^{\lambda} C_{ij} (x_i - x_{0j}) \delta x_i
\]

In this case, it is expected that the evolutionary branching (Dieckmann and Metz, in preparation). In this case, it is expected that the evolutionary branching, if the following conditions hold. In this case, it is expected that the evolutionary branching, if the following conditions hold.

\[
\frac{x_0}{x_{0j}} = \frac{1}{\sqrt{j}} \sum_{j=1}^{\lambda} \frac{C_{ij} (x_i - x_{0j})}{G_i} \delta x_i + O(\sigma^3)
\]

where \( \delta y = (G_i / G_j) \delta y_j \) is the element of \( \delta y \) parallel to the fitness gradient \( G_j \) in the insensitive subspace. Notice that the insensitive subspace contributes to the invasion fitness above only through this element \( \delta y \). Thus, the dimensionality of the local evolutionary dynamics can be contracted to \( L + 1 \). If the sensitive subspace is univariate, then \( G_i \), \( C_{ix} \), and \( D_{ax} \) become scalars, in which case Eq. (O.3) yields

\[
f(s'; s) = G_i \delta x_i + C_{ix} (x_i - x_{0j}) \delta x_i + \frac{1}{2} D_{ax} \delta x_i \delta x_i + |G_i| |\delta y_j| + O(\sigma^3).
\]

Thus, the local evolutionary dynamics around \( x_0 \) can be contracted into that in a bivariate trait space \((x_i, x_j)\). Then, by denoting \( G_i \) by \( |G_j| \), the conditions for evolutionary-branching lines in bivariate trait species can be applied directly, yielding

\[
\lambda_{1i} < 0,
\]

for all \( i = 1, \ldots, L \), where \( \lambda_{1i}, \ldots, \lambda_{L} \) are the eigenvalues of \( C_{xx} \) (Leimar, in press). Second, \( x_0 \) is also evolutionarily unstable, i.e.,

\[
\lambda_{L_{max}} > 0.
\]

Here \( \lambda_{L_{max}} \) is the maximum eigenvalue of \( D_{ax} \), with eigenvector \( v_{0_{max}} \). This \( \lambda_{L_{max}} \) is always real, as \( D_{ax} \) is a symmetric matrix. The inequality above means that the fitness landscape has a positive curvature (i.e., second derivative) in the direction of \( v_{0_{max}} \) with selection favoring evolutionary diversification mainly in this direction. As mutual invasibility in this direction is also ensured in this case, i.e., \( v_{0_{max}} = D_{ax} - C_{xx} v_{0_{max}} > 0 \) always holds under inequalities (0.5) and (0.6), dimorphic divergence may proceed without collapse, resulting in evolutionary branching (Dieckmann and Metz, in preparation). In this case, it is expected that the diversifying residents stay in the neighborhood of the line \( x_i = x_0 + \alpha v_{0_{max}} \) (with real parameter \( \alpha \)). Thus, if the \( x \)-axis is chosen as \( x = \alpha v_{0_{max}} (x_i - x_0) \), then the condition

\[
D = \sigma_{L_{max}} > \frac{1}{\sqrt{2}}
\]

and inequalities (0.5) together may be a useful indicator of the likelihood of evolutionary branching.

In summary, Eq. (O.2) allows a multivariate trait space to be decomposed into a sensitive subspace and an insensitive one; if the sensitive subspace is univariate, the situation is reduced to a bivariate one, in which case the MLIP condition is applicable. On the other hand, if the sensitive subspace is multivariate, there is no assurance of the validity of the MLIP condition. Yet, inequalities (0.5) and (0.7) may still be useful.

### Appendix P. Checking conditions for evolutionary-branching lines without prior normalization

Here we explain how conditions for evolutionary-branching lines can be checked without the prior normalization of trait spaces. We consider a not-normalized \( L \)-variate trait space \( S = (U_1, \ldots, U_L)^T \) that has a mutation variance–covariance matrix \( \Lambda \), which is diagonalized as

\[
\Lambda = P_{\Lambda} \begin{pmatrix} \sigma_1^2 & 0 & 0 \\ 0 & \ldots & 0 \\ 0 & 0 & \sigma_L^2 \end{pmatrix} P_{\Lambda}^T.
\]

where \( P_\Lambda \) is an orthogonal matrix, \( P_{\Lambda}^T = P_\Lambda \), and the square roots of all eigenvalues \( \sigma_1, \ldots, \sigma_L \) are positive. Without loss of generality, \( \sigma_1 \) is assumed to be the largest eigenvalue.

In this trait space, the invasion–fitness function can be written analogously to the bivariate case (Appendix A) as

\[
F(s'; S) = G_{s} S + (S - S_0)^T C_{s} S + \frac{1}{2} (S - S_0)^T D_{s} (S - S_0) + O(\sigma^2),
\]

where \( \tilde{G} = F_{mm}, \tilde{C} = F_{mnm} + F_{mm}, \) and \( D = F_{nnm} \), with the subscripts ‘m’ and ‘r’ corresponding to the mutant \( S \) and the resident \( S \), respectively, and where \( F_{mn} = \partial F(S; S) / \partial S |_{S = S_0} \) is the gradient row vector at \( S_0 \), while \( F_{mm} = \partial^2 F(S; S) / \partial S \partial S |_{S = S_0} \) and \( F_{nn} = \partial^2 F(S; S) / \partial S \partial S |_{S = S_S} \) are Hessian matrices.

This trait space can first be normalized to yield isotropic mutation with standard deviation \( \sigma < 1 \), where \( \sigma \) otherwise is arbitrary, by an affine transformation to coordinates \( s \).

\[
S = Q^T s,
\]

with

\[
Q = \frac{1}{\sigma} \begin{pmatrix} \sigma_1 & 0 & 0 \\ 0 & \ldots & 0 \\ 0 & 0 & \sigma_L \end{pmatrix} P_{\Lambda}^T.
\]

because \( \Lambda = (\sigma Q)^T (\sigma Q) \) gives \( \delta S^T \Lambda^{-1} \delta S = |Q^T \delta s|^2 \sigma^2 |Q|^T Q^{-1} \delta S^T \delta S / \sigma^2 \) (see also Appendix Q). (Notice that the matrix \( Q \) can be interpreted as the Cholesky decomposition of \( \Lambda / \sigma^2 \).)

Next, to adjust the axes of \( s = (u_1, \ldots, u_L)^T \) so as to maximize the sensitivity difference, we add a coordinate rotation to Eq. (P.3).

\[
S = Q^T B s,
\]

with an orthogonal matrix \( B \) for describing the rotation. Substituting this equation into Eq. (P.2) yields

\[
f(s'; s) = F(Q B s'; Q B s) = G_{s} S + (S - S_0)^T C_{s} S + \frac{1}{2} (S - S_0)^T D_{s} (S - S_0) + O(\sigma^2),
\]

where

\[
G = \tilde{G} Q^T B,
\]

\[
C = B^T \tilde{C} Q^T B,
\]

\[
D = B^T \tilde{D} Q D B^T.
\]

which are Eqs. (23) in Section 7.

Finally, we explain how to rotate the axes of \( s = (u_1, \ldots, u_L)^T \) such that the condition for significant sensitivity difference, Eq. (21) with \( L = 1 \),

\[
\frac{|G_1| + |C_{11}| + |C_{1j}| + |D_{1j}| + |D_{jj}|}{|G_1| + |C_{11}| + |D_{1j}|} = O(\sigma)
\]

(0.6)
for all \( j = 2, \ldots, L \), may hold \((L = 2\) gives the condition for significant sensitivity difference for the bivariate case, Eq. (4)). The exact approach is to find the matrix \( B^* \) that minimizes the left-hand side of Eq. (P6). However, this \( B^* \) may not be easy to determine. Fortunately, as explained below, \( B^* \) can be obtained approximately without such minimization when the MLIP condition, inequality (18) in Section 5, holds. In particular, when the condition for significant sensitivity difference and the MLIP condition both hold, all components of \( D \) are \( O(\sigma) \), except for \( D_{11} \), which is of order \( \sigma^0 \). In this case, when the trait space is bivariate \((L = 2)\), each of the two eigenvectors of \( D \) is almost parallel to one axis of the trait space, with the corresponding eigenvalues of \( D \) satisfying \( \lambda_{11} = D_{11} + O(\sigma) \) and \( \lambda_{22} = O(\sigma) \). In other words, \( B^* \) approximately diagonalizes the symmetric matrix \( QDQ^T \). In this case, \( B^* \) is approximately obtained by requiring that \( B^*QDQ^TB^* \) becomes diagonal

\[
B^* \approx (\tilde{B}D_1, \tilde{B}D_2),
\]

where \( \tilde{B}D_1 \) and \( \tilde{B}D_2 \) are the eigenvectors of \( QDQ^T \).

 Analogously, when the trait space is more than bivariate \((L > 2)\), \( B^* \) is approximately obtained as in Eq. (24) in Section 7,

\[
B^* \approx (\tilde{B}D_1, \ldots, \tilde{B}D_L),
\]

where \( \tilde{B}D_1, \ldots, \tilde{B}D_L \) are the eigenvectors of \( QDQ^T \), corresponding to the eigenvalues \( \tilde{\lambda}_{11}, \ldots, \tilde{\lambda}_{L} \), where the eigenvalues are assumed to be ordered such that \( \tilde{\lambda}_{11} > \tilde{\lambda}_{22} \) for all \( j = 2, \ldots, L \), without loss of generality. (When \( \tilde{B}D_1, \ldots, \tilde{B}D_L \) are difficult to obtain, possibly because of too small \( \tilde{\lambda}_{22}, \ldots, \tilde{\lambda}_{L} \), those vectors can be chosen arbitrarily as long as \( \tilde{B}D_1, \ldots, \tilde{B}D_L \) form an orthogonal coordinate system.) If conditions for evolutionary-branching lines hold for this \( B^* \), an evolutionary-branching line passes through \( S_0 \).

**Appendix Q. Directional evolution in MLIPs**

Here we derive MLIPs of monomorphic populations when directional selection is the dominant selection pressure. We consider an arbitrary \( L \)-variate trait space \( S = (U_1, \ldots, U_L)^T \) and a monomorphic population with phenotype \( S \). The invasion-fitness function is written as Eq. (P2) in Appendix P,

\[
F(S; S) = \mathcal{L}S + (S - S_0)^2 + \lambda S - \lambda' T S + \text{h.o.t.}
\]

We assume a mutation distribution given by a multivariate Gaussian function,

\[
M(\delta S) = \frac{1}{(2\pi)^{L/2}|\Lambda|^{1/2}} \exp\left( -\frac{1}{2} \delta S^T \Lambda^{-1} \delta S \right),
\]

where \( \Lambda \) is an \( L \times L \) variance–covariance matrix. By normalizing this trait space using \( S = Q^T s \) in Eq. (P5a) in Appendix P, the argument of the exponential function in Eq. (Q2) is transformed into

\[
\frac{1}{2} \delta S^T \Lambda^{-1} \delta S = \frac{1}{2} \delta S^T Q \Lambda^{-1} Q^T \delta S = \frac{1}{2} \sigma^2 \delta S^T Q Q^{-1} (Q^T)^{-1} Q^T \delta S = \frac{1}{2} \sigma^2 \delta S^2.
\]

Then the mutation distribution in the normalized trait space is given by

\[
M(\delta S) = A_0 M(Q(\delta S)) = \frac{A_0}{(2\pi)^{L/2}|\Lambda|^{1/2}} \exp\left( -\frac{1}{2} \delta S^2 / \sigma^2 \right),
\]

where \( A_0 \) is determined by \( \int M(\delta S) d\delta S = 1 \). In addition, substituting \( S = Q^T s \) into Eq. (Q1) gives the normalized invasion-fitness function

\[
f(s^T; s) = Q^T \delta S + (s - s_0)^T Q^T \delta S + \frac{1}{2} \sigma^2 \delta S^2 Q Q^T \delta S + O(\sigma^2).
\]

Without loss of generality, we chose the base point \( s_0 \) of the expansion as the resident phenotype \( s_0 = s \) after each invasion event. If the gradient \( GQ^T \) is the dominant component of the invasion fitness, such that we can neglect the higher-order terms, the MLI mutant \( s_{MLI} \) is obtained as

\[
s_{MLI} = s + \sigma GQ^T / \mu h \quad \text{(Q6)}
\]

by substituting Eqs. (Q4) and (Q5) into Eq. (8a) in Section 4. The expected waiting time for the next invasion event is given by

\[
T = \frac{\sqrt{2\pi} \sigma}{\mu h} Q^T \mu h \quad \text{(Q7)}
\]

Then the directional change of the resident \( s \) per unit time is given by

\[
s_{MLI} - s = \frac{2}{\sqrt{2\pi} \sigma} \mu h \sigma Q^T \mu h \quad \text{(Q8)}
\]

Substituting \( s = (Q^T)^{-1} S \) yields

\[
dS/dt = Q^T (s_{MLI} - s) / \mu h + \frac{\mu h}{2} Q^T \mu h \quad \text{(Q9)}
\]

As the canonical equation of adaptive dynamics theory (Dieckmann and Law, 1996) is given by

\[
dS/dt = \mu h Q^T H \quad \text{(Q10)}
\]

we see that Eq. (Q9) differs from the canonical equation only in speed, by a factor of \( \sqrt{2/\pi} \approx 0.798 \).

**References**


