The transition from evolutionary stability to branching: A catastrophic evolutionary shift

Fabio Dercole¹, Fabio Della Rossa¹ & Pietro Landi²,³

Evolutionary branching—resident-mutant coexistence under disruptive selection—is one of the main contributions of Adaptive Dynamics (AD), the mathematical framework introduced by S.A.H. Geritz, J.A.J. Metz, and coauthors to model the long-term evolution of coevolving multi-species communities. It has been shown to be the basic mechanism for sympatric and parapatric speciation, despite the essential asexual nature of AD. After 20 years from its introduction, we unfold the transition from evolutionary stability (ESS) to branching, along with gradual change in environmental, control, or exploitation parameters. The transition is a catastrophic evolutionary shift, the branching dynamics driving the system to a nonlocal evolutionary attractor that is viable before the transition, but unreachable from the ESS. Weak evolutionary stability hence qualifies as an early-warning signal for branching and a testable measure of the community’s resilience against biodiversity. We clarify a controversial theoretical question about the smoothness of the mutant invasion fitness at incipient branching. While a supposed nonsmoothness at third order long prevented the analysis of the ESS-branching transition, we argue that smoothness is generally expected and derive a local canonical model in terms of the geometry of the invasion fitness before branching. Any generic AD model undergoing the transition qualitatively behaves like our canonical model.

Twenty years ago, S.A.H. Geritz, J.A.J. Metz, and coauthors¹–³ introduced a mathematical framework—referred to as Adaptive Dynamics (AD)—for modeling long-term evolutionary dynamics, with special emphasis on the generation of diversity through evolutionary branching. AD has been successfully developed and applied in and outside biology (see ref. 4 and refs therein) as well as debated, mainly due to its asexual nature (see vol. 18 of the J. Evol. Bio.). Evolutionary branching takes off when a resident and a similar mutant type coexist in the same environment and natural selection is disruptive, i.e., favors outer rather than intermediate phenotypes. In the restricted (but classical) formulation in which resident individuals are characterized by the same value \( x \) of a one-dimensional strategy, Geritz, Metz et al.¹–³ derived explicit conditions for evolutionary branching in terms of the invasion fitness—namely the exponential rate of growth \( s(x) \) initially shown by a mutant strategy \( y \) appeared when the resident is at its ecological regime.

Subsequently, analogous conditions have been derived for models including sexual characters⁶–⁷, spatial distribution⁸, and multi-dimensional phenotypes⁹, establishing evolutionary branching as the prerequisite step for sympatric and parapatric speciation¹⁰.

Evolutionary branching requires, first, the resident strategy \( x \) to be in the vicinity of a convergence stable singular strategy \( x^* \)—a strategy making neutral the selection pressure measured by the fitness gradient \( \partial_y s_x(y) \big|_{y=x} \), i.e.,

\[
\partial_y s_x(y) \big|_{y=x} = 0 \quad \text{(singularity),}
\]

and locally attracting the evolutionary dynamics driven by rare and small strategy mutations¹–³,¹¹,¹² (convergence stability).

Second, there should be pairs \( (x_1, x_2) \) close to the singular point \( (x^*, x^*) \) for which a resident \( x = x_1 \) and a mutant \( y = x_2 \) can coexist. For this, a sufficient condition is a negative fitness cross-derivative, i.e.,
that implies the existence, locally to \((x^*, x^*)\), of a conical region of coexistence in the strategy plane (Fig. 1). Geritz, Metz et al.\textsuperscript{1–3} showed that mutual invasibility occurs in the region, i.e., that (2) implies the invasion fitness of strategy \(x_2\) against the resident \(x_1\), and vice versa, \(s_{x_1}(x_2) > 0\), and vanishing each on one of the region boundaries. The existence of a unique ecological attractor of coexistence was shown more recently\textsuperscript{13–15} for the class of unstructured ecological models—no individual distinction w.r.t. age, size, location, etc.—under stationary coexistence. (See Fig. 1c, where \(n_{x_1}(x_2)\) is the equilibrium density of the resident before mutant invasion and \(n_{x_2}(x_1)\) and \(n_{x_2}(x_1)\) are the equilibrium densities of coexistence).

Third, selection must be disruptive, a sufficient condition for which being

\[
\partial y^* := \left. \partial y s_y \right|_{y=y^*} < 0, \tag{2}
\]

i.e., the opposite of the evolutionary stability of the singular strategy\textsuperscript{16,17} (see Fig. 1e,f).

Condition (3) was obtained\textsuperscript{1–3} based on the twice differentiability of the dimorphic fitness \(s_{y_1}(x_1)\) — the invasion fitness of a mutant \(y_1\) invading an environment set by the two residents \(x_1\) and \(x_2\) at their ecological regime—and by expanding it around \((x_1^*, x_2^*)\) (see Appx. 1 in ref. 3 for details). Exploiting the consistency relations

C1: \(s_{x_1}(x_2) = s_{x_2}(x_1)\), the link between the dimorphic and monomorphic fitness functions at the singular point,

C2: \(s_{x_1}(x_2) = s_{x_2}(x_1)\), the order irrelevance of the two residents, and

C3: \(s_{y_1}(x_1) = 0\) (a) and \(s_{y_2}(x_2) = 0\) (b), the selective neutrality of the two residents,

Geritz, Metz et al.\textsuperscript{1–3} arrived to the following expansion

\[
\partial y s^* := \left. \partial y s_y \right|_{y=y^*} > 0, \tag{3}
\]

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\[
\partial y s^* := \left. \partial y s_y \right|_{y=y^*} > 0, \tag{3}
\]
\[ s_{x_i}(y) = \frac{1}{2} \partial_\cdot\cdot s^*(y - x)(y - x) + \cdots \]  

(4)

It says that the selection pressures on strategies \( x_i \) and \( x_j \), as measured by the fitness gradients

\[ \partial_j s_{x_i}(y) = \frac{1}{2} \partial_\cdot\cdot s^*(2x_i - x_j + x_j) + \cdots, \quad i = 1, 2, \]  

(5)

are opposite (\( x_i, x_j \)) in the region of coexistence locally to (\( x^*, x^* \)), meaning that dimorphic evolution points away or toward (\( x^*, x^* \)) if \( \partial_j s^* \geq 0 \) (see Fig. 1f,e, respectively).

The smoothness of the dimorphic fitness \( s_{x_i}(y) \) at the singular point \((x_i, x_j) = (x^*, x^*)\) is a controversial open problem of AD. In ref. 1 (Sect. 6.3.2), the twice differentiability is shown geometrically (see Methods for an algebraic proof) and the authors argue that higher-order derivatives may fail to exist at the singular point. The source of the expected nonsmoothness is the ecological attractor of coexistence being undefined when \( x = x_i \). Think, e.g., of the equilibrium densities \( \eta_1(x_i, x_j) \) and \( \eta_2(x_i, x_j) \) in Fig. 1. They are evidently discontinuous at \((x^*, x^*)\), respectively being equal to zero and \( \pi(x_i) \) along the extinction boundary 1 (blue) and to \( \pi(x_j) \) and zero along the extinction boundary 2 (red). The nonsmoothness of the dimorphic fitness arising at 3rd order remained a mystery of AD that long prevented the analysis of the ESS-branching transition.

The nonsmoothness conjecture was recently disproved (exploiting a new structural ecological property) for the class of unstructured ecological models under stationary coexistence. We have indeed shown that the smoothness of the dimorphic fitness up to 3rd order. Moreover, our methodology is general up to any order and there is thus no reason to expect nonsmoothness to arise at some higher order.

Here we lift the above result to the generality of any class of ecological models, based on the newly discovered smoothness of the dimorphic fitness. Only assuming a three-times differentiable dimorphic fitness, we show that the consistency relations C1–C3 (with C1 considered along the extinction boundaries delimiting the coexistence region, see Methods) determine the expansion of the dimorphic fitness up to 3rd order, and the result is the same as obtained in ref. 20. That is, the 1st order is null, the 2nd coincides with (4), and the 3rd is

\[ \left\{-\frac{1}{4} \frac{\partial_j s^* s_i s_j s^*}{s_i s^*} (x_1 + x_2 - 2x^*) + \frac{1}{6} \partial_j s^* (x_1 + x_2 + y - 3x^*) \right\} (y - x_1)(y - x_2). \]  

(6)

Note that the expansion is given in terms of the monomorphic fitness derivatives \( \left( \partial_j s^* := \frac{\partial_j s^*}{s_i s^*} \right|_{y=x, s^=} \) and \( \partial_j s^* := \frac{\partial_j s^*}{s_i s^*} \) are the involved third derivatives)—that are determined by the ecological dynamics before branching—in contrast to what preliminarily expected based on the analysis of Lotka-Volterra models. Unfortunately, the consistency arguments offered by relations C1–C3 do not determine the fitness expansion at higher orders, so whether the local approximation of the dimorphic fitness is fully controlled by the geometry of the monomorphic fitness remains an open question. Any answer necessarily requires specifying the class of underlying ecological models, so that one can proceed with the direct computation of the fitness expansion, as in ref. 20.

Our analysis allows to go beyond 2nd order in the analysis of the branching dynamics, thus solving the 20-yrs-lasting impasse. In particular, the analysis at 3rd order is of fundamental importance, because necessary to unfold the mechanisms underlying the ESS-branching transition \( (\partial_j s^* = 0) \) under the coexistence condition (2), see Fig. 2, close to which the branching dynamics is dominated by the 3rd-order terms in the approximation of the dimorphic fitness. Note that close to the transition \( (\partial_j s^* \approx 0) \) \( \partial_j s^* \) is the leading 3rd-order coefficient in (6), so we perform our analysis assuming the genericity condition

\[ \partial_j s^* := \partial_j s^* \big|_{y=x, s^=} = 0. \]  

(7)

The analyses at 4th and higher orders will be possible only under specific ecological assumptions and are interesting for the understanding of degenerate branching scenarios, where some of the fitness derivatives vanish at the singularity due to model-specific properties, like symmetries in the phenotypic dependence of demographic parameters around optimal values.

Based on the 3rd-order expansion (4, 6), we derive the following canonical model for the ESS-branching transition:

\[ \dot{x}_i = -\frac{1}{\partial_j s^*} \eta_i (\Delta x_i, \Delta x_j) s_j (\Delta x_i, \Delta x_j), \quad \Delta x_i := x_i - x^*, \quad i = 1, 2, \]  

(8a)

\[ \eta_1(\Delta x_i, \Delta x_j) := \partial_j s^* \Delta x_i + \frac{1}{2} \partial_j s^* (\Delta x_i + \Delta x_j) + \frac{1}{2} \partial_j s^* \Delta x_j^2 \]

\[ + \frac{1}{2} \partial_j s^* \Delta x_i (\Delta x_i + \Delta x_j) + \frac{1}{6} \partial_j s^* (\Delta x_j^2 + \Delta x_i \Delta x_j + \Delta x_j^2), \]  

(8b)

\[ \eta_2(\Delta x_i, \Delta x_j) := \eta_1(\Delta x_j, \Delta x_i), \]  

(8c)
Figure 2. Unfolding of the ESS-branching transition. The flow of model (8) (black trajectories with dashed $x_1$-blue- and $x_2$-red-nullclines) is shown in the resident-mutant coexistence region delimited by the extinction boundaries $1$ ($\eta_1(\Delta x_1, \Delta x_2) = 0$, see Eq. (8b), blue) and $2$ ($\eta_2(\Delta x_1, \Delta x_2) = 0$, see Eq. (8c), red). The panels are symmetric w.r.t. the diagonal $\Delta x_1 = \Delta x_2$ (due to property C2). The cubic approximation (4, 6) of the dimorphic fitness gives only a quadratic approximation of the extinction boundaries and a linear approximation of the nullclines w.r.t. their fully nonlinear versions. The approximation of the dimorphic flow is also quadratic in the case of stationary ecological coexistence (see Methods). The figure is obtained for $\partial_j x^* s = \partial_j^2 x^* s = -1$, $\partial_j x^* s = 2$, $\partial_j x^* s = 10$. The case with $\partial_j x^* s < 0$ can be obtained through a local symmetry w.r.t. the anti-diagonal $\Delta x_1 + \Delta x_2 = 0$.

\[
s_j(\Delta x_1, \Delta x_2) := \frac{1}{2} \partial_j x^* s - \frac{1}{4} \partial_j x^* s \partial x_j x^* s + \frac{1}{6} \partial_j x^* s (2\Delta x_1 + \Delta x_2),
\]

(8d)

\[
s_2(\Delta x_1, \Delta x_2) := s_j(\Delta x_2, \Delta x_1).
\]

(8e)

The evolutionary dynamics of model (8) are depicted (under $\partial_j x^* s > 0$) in Fig. 2 in the coexistence region delimited by the extinction boundaries $1$ ($\eta_1(\Delta x_1, \Delta x_2) = 0$ (blue) and $\eta_2(\Delta x_1, \Delta x_2) = 0$ (red). Within the coexistence region, $(-1/\partial_j x^* s) \eta_j(\Delta x_1, \Delta x_2)/\eta_i(\Delta x_1, \Delta x_2)$ represents the birth output of populations at the ecological regime and $s_j(\Delta x_1, \Delta x_2)(\Delta x_i - \Delta x_j)$, $i \neq j$, approximates the fitness gradient (5) (see Methods). The (black) trajectories describe the continuous dynamics obtained in the limit of infinitesimally small mutational steps. The dashed (blue and red) lines represent $s_j(\Delta x_1, \Delta x_2) = 0$ and $s_2(\Delta x_1, \Delta x_2) = 0$ and therefore approximate the $x_j$ and $x_2$ nullclines of the evolutionary dynamics (the lines on which selection is neutral in strategy $x_j$ and $x_2$, respectively, so that evolution points vertically and horizontally). The nullclines and the extinction boundaries (the latter also behaving as nullclines with same color code) are known to connect at special points—the boundary equilibrium with one strategy at the singularity $x^*$, the other being absent (connections with different colors, half-filled points), and the points where the absent strategy is (potentially) singular (same color connections, occurring at the horizontal and vertical extremal points of the extinction boundaries). Model (8) shows such connections while approaching the ESS-branching transition.

The unfolding of Fig. 2 has been observed in previous analyses of specific models and conjectured to be general based on the above described properties of the evolutionary nullclines. Our contribution at last gives the full theoretical support. We have shown that any single or multi-species community undergoing the ESS-branching transition due to gradual changes in, e.g., climate, nutrients, habitat fragmentation, or biotic control and exploitation qualitatively behaves like model (8) near the singular point ($x^*$).

Figure 2 clearly shows the mechanisms underlying the ESS-branching transition. Restricting the attention to the coexistence region above the diagonal (since relation C2 implies the symmetry w.r.t. the diagonal), the boundary equilibria (half-filled) at $x_1 = x^*$ (left) and at $x_2 = x^*$ (right) behave as saddles for the dimorphic evolutionary dynamics, attracting along one trajectory—the stable manifold of the saddle—and repelling for all other nearby initial conditions. In the ESS case (left), the saddle’s stable manifold separates the initial conditions leading to the ESS—a dimorphic phase up to the extinction of one of the two populations followed by the monomorphic convergence (represented on the diagonal) to the ESS—from those leading away, typically to a nonlocal evolutionary attractor not involved in the transition. In the branching case (right), the initial conditions near the singular point ($x^*$) all lead away. Approaching the transition (left-to-right) along with gradual changes in the model parameters, the boundary saddle approaches the ESS, so do the initial conditions leading away. Note that
branching is possible at the transition too (for mutant strategy \(x^*\) larger than \(x^\circ\), central panel; branching at the transition symmetrically requires \(x^* < x^\circ\) under \(\partial_1 s^\circ < 0\), not shown, see Methods).

In real systems, where mutational steps are small but finite, the monomorphic dynamics converging to the singular strategy \(x^\circ\) eventually jump into the dimorphic region. In a “phase transition” case far from the transition (see Fig. 2, left, panel 2), the short periods of rapid morphological change observed by paleontologists in the fossil record, along with gradual environmental changes on the macro scale—the so-called punctuated equilibria—could correspond to the branching dynamics triggered by the transition and developing on the meso scale.

This work fully explains one of the most important “phase transition” in Nature.

Methods Summary

We consider two similar populations, with densities \(n_1\) and \(n_2\) and one-dimensional strategies \(x_1\) and \(x_2\). In the monomorphic phase populations 1 and 2 are resident and mutant, respectively, whereas they are both residents in the dimorphic phase. We do not specify a particular class of ecological models, but only assume that the resulting monomorphic and dimorphic invasion fitnesses are smooth (only differentiability up to \(3^\text{rd}\) order is actually required). In particular, for the dimorphic fitness \(s_1(x_1, x_2)\), smoothness at \((x_1, x_2, y) = (x^\circ, x^\circ, x^\circ)\) means that a polynomial local expansion in \(\Delta x_i := x_i - x^\circ\), \(i = 1, 2\), \(\Delta y := y - x^\circ\) holds good in the resident-mutant coexistence region—the error of a truncated expansion of order \(k\) is \(O((\Delta x_1, \Delta x_2, \Delta y)^{k+1})\) when \((\Delta x_1, \Delta x_2)\) vanish along a path in the coexistence region.

The above smoothness assumption is currently proved for the class of unstructured ecological models under stationary coexistence, though we expect it to hold with wide generality. Specifically, smoothness is generically expected radially from the singular point \((x^\circ, x^\circ, x^\circ)\), with \(\theta\) parameterizing the extinction boundary 2 in the strategy plane \((x_1, x_2)\). Then, showing the smoothness of the dimorphic fitness at \((x_1, x_2, y) = (x^\circ, x^\circ, x^\circ)\) reduces to showing that the directional expansion along rays \((x_1, x_2) = (x^\circ + \varepsilon \cos \theta, x^\circ + \varepsilon \sin \theta)\) at given \(\theta\) and around \(y = x^\circ\) is polynomial in the direction components \((\cos \theta, \sin \theta)\). However, without specific assumptions on the underlying ecological model, this is possible only up to order 2 (see SI, Sect. 1).

Using a \(3^\text{rd}\)-order expansion of the dimorphic fitness and imposing the consistency relations C1–C3, with C1 replaced by

\[ C1': s_{x_1x_2}(y) = s_{x_1}(y), \]

along the extinction boundary 2 on which only strategy \(x_1\) is present, we derive the expansion (4, 6) under the coexistence condition (2) (note that imposing the monomorphic–dimorphic link on the extinction boundary 1 is redundant due to the diagonal symmetry between the two boundaries and property C2, see SI, Sects. 2 and 3).

Specifically, to impose C1’, we use the polar coordinates \((\varepsilon, \theta)\) and \(\varepsilon\)-parameterize the extinction boundary 2 as \(\theta = \theta_1(\varepsilon), \theta_2(\varepsilon)\) being the function that gives the angle \(\theta\) of the boundary point at distance \(\varepsilon\) from the singular point \((x^\circ, x^\circ)\). Then, C1’ becomes

\[ C1': s_{x_1 + \varepsilon \cos \theta_1(\varepsilon)}(x^* + \varepsilon \sin \theta_1(\varepsilon)) (x^* + \Delta y) = s_{x_1 + \varepsilon \cos \theta_2(\varepsilon)}(x^* + \Delta y), \]

to be imposed together with its \((\varepsilon, \Delta y)\)-derivatives at \((\varepsilon, \Delta y) = (0, 0)\) up to order 3. This involves the angle \(\theta(0)\)—the tangent direction to the extinction boundary 2 at \((x^\circ, x^\circ)\) and the first derivative \(\partial_1 s(x^*, x^\circ)\)—the local curvature of the boundary (whether \(\theta\) increases or decreases while moving away from \((x^\circ, x^\circ)\), see Fig. 1e,f). The two quantities are determined by the second and third derivatives of the monomorphic fitness. They are obtained by imposing the second and third \(\varepsilon\)-derivatives of the boundary definition

\[ s_{x_1}(x_2) = s_{x_1 + \varepsilon \cos \theta_2(\varepsilon)}(x^* + \varepsilon \sin \theta_2(\varepsilon)) = 0 \]

(the first derivative being uninformative).
Unfortunately, the above procedure does not apply at order 4, as the conditions imposed by relations C1′–C3 are less than the number of coefficients in the expansion of the dimorphic fitness. The missing conditions must then come from specific assumptions to be made at the ecological level.

The derivation and the analysis of the canonical model (8) are detailed in the Supplementary Information (Sects. 4 and 5).

An example

We illustrate the developed body of theory on a well-known example: a single species AD Lotka-Volterra model of asymmetric competition. The resident-mutant ecological model reads:

\[
\dot{n}_1 = n_1(\rho(x_1) - \alpha(0)n_1 - \alpha(x_1 - x_2)n_2),
\]

\[
\dot{n}_2 = n_2(\rho(x_2) - \alpha(x_2 - x_1)n_1 - \alpha(0)n_2),
\]

where \(\dot{n}_i\) here stands for the time-derivative of \(n_i\) on the ecological time scale, \(i = 1, 2\), and the strategy \(x\) scales (from \(-\infty\) to \(+\infty\) ) with competitive ability. The intrinsic growth rate \(\rho\) and the competition function \(\alpha\) are strategy-dependent, with Gaussian \(\rho = \sigma^2 \exp(-x^2/2\sigma^2)\), \(\sigma > 0\), that identifies an optimal strategy for a single population, and sigmoidal \(\alpha(x) = 1 - \frac{1}{1 + \exp(-x)}\), \(v > 0\), that gives a competitive advantage to larger strategies.

The model is simple enough that we can solve analytically for all the relevant quantities: the monomorphic and dimorphic resident equilibrium densities (at which \(\dot{n}_i = 0\) in the absence of population \(2\) and \(\dot{n}_1 = \dot{n}_2 = 0\) in the presence of both populations)

\[
\pi = \frac{\rho(\alpha)}{\alpha}, \quad \eta_1(x_1, x_2) = \frac{\rho(x_1)\alpha(0) - \rho(x_2)\alpha(x_1 - x_2)}{\alpha^2 - \alpha(x_1 - x_2)\alpha(x_2 - x_1)}, \quad \eta_2(x_1, x_2) = \eta_1(x_2, x_1),
\]

the monomorphic and dimorphic fitnesses

\[
s(x) = \rho(\alpha)/\alpha, \quad s_{1}(x_1, x_2) = \rho(x_1)\alpha(0) - \rho(x_2)\alpha(x_1 - x_2)/\alpha^2 - \alpha(x_1 - x_2)\alpha(x_2 - x_1),
\]

the monomorphic and dimorphic fitness gradients

\[
\frac{\partial s_{1}(y)}{\partial x} = \rho'(x) - \frac{\alpha'(0)}{\alpha}\rho(x) = \frac{\sigma^2 - \alpha\sigma^2}{\sigma^2 + 2\sigma\exp(x^2/2\sigma^2)},
\]

\[
\frac{\partial s_{2}(y)}{\partial x} = \rho'(x) - \frac{\alpha'(0)}{\alpha}\rho(x) = \frac{\sigma^2 - \alpha\sigma^2}{\sigma^2 + 2\sigma\exp(x^2/2\sigma^2)}.
\]

The singular strategy making the monomorphic fitness gradient zero

\[
x^* = \sigma^2/(1 + \nu),
\]
the fitness second derivatives ruling branching at $x^*$

$$\partial_2 x^* = \frac{\alpha''(0)}{\alpha(0)} \rho(x^*) - \frac{\alpha'(0)}{\alpha(0)} \rho'(x^*) = -\frac{\nu}{(1 + \nu)^2 \exp[(\sigma^2/2(1 + \nu)^2)]}, \quad$$

$$\partial_3 x^* = \frac{\rho''(x^*)}{\alpha(0)} \rho(x^*) = -\frac{1}{\sigma^2(1 + \nu)^2 \exp[(\sigma^2/2(1 + \nu)^2)]},$$

and the third derivatives entering our approximations

$$\partial_2^2 x^* = \frac{\alpha''(0)}{\alpha(0)} \rho(x^*) + \frac{2\alpha''''(0)}{\alpha(0)} \rho''(x^*) - \frac{\alpha'(0)}{\alpha(0)} \rho'(x^*) = \frac{\nu^2 \sigma^2 - \nu^2 - 2\nu \sigma^2 - 2\nu - 1}{\sigma^2(1 + \nu)^3 \exp[(\sigma^2/2(1 + \nu)^2)],}$$

$$\partial_3 x^* = \frac{\alpha''(0)}{\alpha(0)} \rho'(x^*) - \frac{\alpha'(0)}{\alpha(0)} \rho''(x^*) = \frac{\nu(3 - \nu)}{(1 + \nu)^3 \exp[(\sigma^2/2(1 + \nu)^2)]},$$

$$\partial_2^3 x^* = \frac{\rho''(x^*)}{\alpha(0)} \rho(x^*) = \frac{\nu^2 \sigma^2 + 3\nu + 4\nu \sigma^2 + 6\nu + 3}{\sigma^2(1 + \nu)^3 \exp[(\sigma^2/2(1 + \nu)^2)].}$$

It is easy to verify that the singular strategy $x^*$ is attracting the monomorphic evolutionary dynamics for any positive $(\sigma, \nu)$ (the eigenvalue of the continuous dynamics is $\partial_2 x^* + \partial_3 x^* = -(\sigma^2 \exp[(\sigma^2/2(1 + \nu)^2)])^{-1} < 0$, that coexistence in its vicinity is always possible ($\partial_3 x^* < 0$ for $\nu > 0$, and that branching ($\partial_3 x^* > 0$) occurs if $\sigma^2 > (1 + \nu)^2/\nu$. At $\sigma^2 = (1 + \nu)^2/\nu$ the system undergoes the ESS-branching transition. Increasing the value of $\sigma$, the ESS turns into a branching point.

Figure 3 compares our (approximating) canonical model (8) and extinction boundaries $\eta_i(\Delta x_1, \Delta x_2) = 0$. $i = 1, 2$, with the fully nonlinear versions. As in Fig. 2, the extinction boundary 1 and the $x_i$-nullcline of the dimorphic evolutionary dynamics are plotted in blue (solid and dashed); red for boundary 2 and the $x_i$-nullcline. Lighter colors are used for the fully nonlinear versions (recall that the approximation is quadratic for the extinction boundaries and linear for the $x_i$-nullclines, locally to the singular point $(x^*, x^*)$).

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Author Contributions
All authors were involved in the design of the research and in the analysis. F.D.R. and P.L. performed the numerical analysis and produced the graphics, F.D. wrote the paper.

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