Organism-induced habitat restoration leads to bi-stability in metapopulations

Feng Zhang, Yi Tao, Cang Hui

1. Introduction

Habitat destruction and fragmentation is the prime process causing current mass biodiversity loss [1]. As a result, metapopulation theories have been widely applied in conservation and spatial ecology [2]. The classic metapopulation model is developed by Levins [3,4] and has contributed critical insights into the conservation of species on fragmented landscapes [2–4]. Levins’ patch occupancy model can be further modified by incorporating habitat destruction [5–9]. Such a modification has led to the Levins rule in conservation biology, estimating the minimum amount of suitable habitat necessary for the long-term persistence of metapopulations based on the colonization and extinction rates of focal species.

The patch occupancy model, however, assumes that the fraction of suitable habitat remains a constant in the landscape, which is often not the case in real systems. In reality, the size of suitable habitat is constantly fluctuating due to the habitat destruction and restoration. As an alternative to human restoration, organisms themselves could also induce habitat restoration. All organisms affect their environments on various temporal and spatial scales [10,11]. The modification and creation of environment by organisms could potentially facilitate habitat restoration and thus alter the available amount of suitable habitat to themselves or/and other species [12]. For example, soil microorganisms can form biological soil crusts and protect soil surface from erosion, preserving nitrogen and carbon to the soil [13]. Ants, termites and earthworms can also shift the soil profile towards their favorite conditions [10]. Dune plants can reinforce and stabilize their habitat on drift sand [14]; desert plants can accumulate soil content around them and facilitate their future recruitment [15]. These organism-environment feedbacks can cause complex spatial patterns of species distributions [16–19]. Therefore, when evaluating metapopulation persistence, we should include not only habitat destruction (or degradation) and human restoration in the population dynamics, but also the fine-scale organism-induced habitat restoration.

Following Levins’ [3,4] patch occupancy model, we presented a differential-equation model, in which both the metapopulation dynamics and the dynamics of the fraction of suitable patches in the habitat are characterized. Habitat restoration induced by organism itself (internal restoration) and by other organisms or/and abiotic causes (external restoration) were incorporated in the model, together with habitat destruction. Stability analysis revealed the existence of alternative equilibriums (i.e., bi-stability) in the system. The internal restoration of habitat was identified as the trigger for the bi-stability, whereas the external restoration, in contrast, can eliminate the bi-stability from the system. The results, thus, emphasize the important role of the organism-environment feedback in biological conservation.

2. Model

Metapopulations describe a group of local populations inhabiting a patch network linked by migration between patches. The metapopulation theory explains that, even if local populations are
subject to extinction, the long-term persistence of the species can still be achieved at regional level [2]. The classical metapopulation framework is based on the Levins [3,4] patch occupancy model:

$$\frac{dp}{dt} = cp(h - p) - ep.$$  

(1)

where $p$ is the fraction of patches occupied by the species (or called the metapopulation size); parameter $c$ and $e$ are the colonization and extinction rates, respectively; parameter $h$ denotes the fraction of suitable patches (thus $1 - h$ indicates the fraction of unsuitable patches, or the fraction of habitat loss). The non-trivial equilibrium of Eq. (1), $p^* = h - e/c$, is globally stable if $h > e/c$. Therefore, the minimum fraction of suitable patches necessary for metapopulation persistence is defined by the ratio of the extinction rate to the colonization rate, and the fraction of empty suitable patches remains constant (i.e. $h - p^* = e/c$) unless the metapopulation becomes extinct, known as the Levins rule [20].

In the following, we assume that the suitable patches are destructed at a certain rate $(d)$ due to human activities or natural causes (e.g. wild fires and hurricanes). The unsuitable patches can be restored at a certain rate $(\lambda)$ through external habitat restoration by the activities of other organisms (e.g. their metabolism), abiotic causes (e.g. self recovering) and human endeavor (e.g. conservation effort). Moreover, the unsuitable patches can also be restored by the organism itself at a certain rate $(\mu)$, i.e. organism-induced internal habitat restoration. Accordingly, we have the following model by extending Eq. (1) (see Appendix A for a derivation from the Markov process):

$$\frac{dp}{dt} = cp(h - p) - (e + d)p,$$

$$\frac{dh}{dt} = (\lambda p + \mu)(1 - h) - dh.$$  

(2)

Notice that all parameters $(c, e, d, \lambda$ and $\mu$) are non-negative, and that $p \leq h$. We define $\alpha = \lambda/d$ and $\beta = \mu/d$ in the following analysis as the intensities of the internal and external processes of habitat restoration, relative to habitat destruction (at a rate of $d$). Let $\delta = (e + d)/c$ indicate the decreasing rate of the fraction of occupied patches (including the natural extinction of local populations and the patch loss due to habitat destruction) relative to the colonization rate (see Table 1 for details). Eq. (2) implies that the habitat loss is completely random, meaning that (i) all suitable patches including occupied and empty patches may be destroyed at the same rate, and (ii) when an occupied patch is destroyed the local population within the patch goes to extinction simultaneously.

### 3. Analysis

Eq. (2) has a unique boundary equilibrium $(0, \beta/(\beta + 1))$. Obviously, this boundary equilibrium corresponds to the extinction of metapopulation. We have the following two theorems for its stability.

**Theorem 1.** The boundary equilibrium $(0, \beta/(\beta + 1))$ is locally asymptotically stable if $\delta > \beta/\beta + 1$, and is unstable if $\delta < \beta/\beta + 1$.

**Proof.** The Jacobian matrix of Eq. (2) at the boundary equilibrium $(0, \beta/(\beta + 1))$ is:

$$J_{\text{boundary}} = \begin{pmatrix} \frac{cp}{\beta + 1} - (e + d) & 0 \\ \frac{cp}{\beta + 1} - (\lambda + d) & \lambda \end{pmatrix}.$$  

(3)

with two eigenvalues $\frac{cp}{\beta + 1} - (e + d)$ and $- (\lambda + d)$. Thus, the boundary equilibrium $(0, \beta/(\beta + 1))$ is unstable if $\delta < \beta/\beta + 1$ (Fig. 1A) and locally asymptotically stable if $\delta > \beta/\beta + 1$ (Fig. 1D–F). □

### Table 1

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
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<tbody>
<tr>
<td>$c$</td>
<td>Colonization rate to empty suitable patches</td>
</tr>
<tr>
<td>$e$</td>
<td>Extinction rate of local populations</td>
</tr>
<tr>
<td>$d$</td>
<td>Habitat destruction rate</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>Internal habitat restoration rate</td>
</tr>
<tr>
<td>$\mu$</td>
<td>External habitat restoration rate</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Relative intensity of internal restoration to habitat destruction</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Relative intensity of external restoration to habitat destruction</td>
</tr>
<tr>
<td>$\delta$</td>
<td>Relative decrease rate of occupied patch fraction to colonization rate</td>
</tr>
</tbody>
</table>

**Theorem 2.** If $\delta = \beta/(\beta + 1)$, then the boundary equilibrium $(0, \beta/(\beta + 1))$ is locally asymptotically stable if and only if $\alpha \leq \beta/\beta + 1$.

There are at most two interior equilibriums for the Eq. (2) (see Appendix C for details and Table 2 for a summary). For convenience, let $(p^*, h^*)$ denotes the interior equilibrium of Eq. (2). On the $p$-$h$ plane phase, the zero isocline for $dp/dt = 0$, denoted by $L_1$, is determined by the equation $c(h - p) - (e + d) = 0$, and the zero isocline for $dh/dt = 0$, denoted by $L_2$, is determined by the equation $(\lambda p + \mu)(1 - h) - dh = 0$. It is easy to see that the slope of $L_1$ is one, and the slope of $L_2$ at interior equilibrium $(p^*, h^*)$ is $\alpha(1 - h^*)/(\lambda p^* + \mu + d)$. Subsequently, for the stability of $(p^*, h^*)$, we have the following theorem.

**Theorem 3.** An interior equilibrium of Eq. (2), $(p^*, h^*)$, is locally asymptotically stable if and only if the slope of $L_2$ at the interior equilibrium satisfies $\alpha(1 - h^*/(\lambda p^* + \mu + d) < 1$.

**Proof.** The Jacobian matrix of Eq. (2) at $(p^*, h^*)$ is

$$J_{p^*, h^*} = \begin{pmatrix} -cp^* & cp^* \\ \lambda(1 - h^*) & - (\lambda p^* + \mu + d) \end{pmatrix}.$$  

(4)

Notice that the eigenvalues of the matrix $J_{p^*, h^*}$ are

$$-\frac{1}{2} \left( \frac{cp^* + \alpha^*}{(cp^* + \alpha^*)^2} \pm \frac{1}{4(c p^* + \alpha^*)^2} - 4cp^* \alpha^* \left( 1 - \frac{\alpha(1 - h^*)}{\lambda p^* + \mu + d} \right) \right),$$  

(5)

where $\alpha^* = \lambda p^* + \mu + d$. Thus, $(p^*, h^*)$ is locally asymptotically stable if $\alpha(1 - h^*/(\lambda p^* + \mu + d) < 1$ (Fig. 1A, B, D), and unstable if $\alpha(1 - h^*/(\lambda p^* + \mu + d) > 1$ (Fig. 1D). For $\alpha(1 - h^*/(\lambda p^* + \mu + d) = 1$, one of the two eigenvalues is zero, and the other is negative. Eq. (2) can be equivalently rewritten as (similarly to the proof in Theorem 2, see Appendix B)

$$\frac{dp}{dt} = -\frac{\lambda p^*}{cp^* + \alpha^*} p^2 + \lambda(\lambda p^* + \alpha^*) h + \lambda(\lambda p^* + \alpha^*) h^2,$$

(6)

where $\lambda p^* = \alpha p^* + \lambda p^* + \mu + d$ and $\lambda h = \alpha p^* + \lambda p^* + \mu + d$. The function $\lambda(\lambda p^* + \alpha^*) h$ and $\lambda(\lambda p^* + \alpha^*) h^2$ are polynomials with respect to $p$ and $h$ (the first-degree terms are excluded). Therefore, in term of the center manifold theory [21], there exist a local center manifold $h = \phi(p)$ with $\phi(0) = 0$ and $d \phi(0)/dt = 0$, which can be obtained by solving the equation

$$\frac{d \phi(p)}{dp} = -\frac{\lambda p^*}{(cp^* + \alpha^*)^2} p^2 + \lambda(\lambda p^* + \alpha^*) \phi(p) + \lambda(\lambda p^* + \alpha^*) \phi(p).$$  

(7)

Thus, the stability of system (2) is determined by the reduced system...
Fig. 1. Dynamical behaviors of the system are plotted on the $p$-$h$ phase plane (the metapopulation size versus the fraction of suitable patches): (A) a globally asymptotically stable interior equilibrium with $d < b/(1 + b)$; (B) a globally asymptotically stable interior equilibrium with $d = b/(1 + b)$ and $a > (1 + b)^2$; (C) if $d = b/(1 + b)$ and $a \leq (1 + b)^2$, then the boundary equilibrium $(0, b/(1 + b))$ is stable, and no interior equilibrium can exist; (D) for $b/(1 + b) < d < \min\left(\frac{a}{C_0}, \frac{b}{C_0}\right)$, the boundary $(0, b/(1 + b))$ is stable, and there are two interior equilibriums, in which one is stable and the other is unstable; (E) the boundary equilibrium $(0, b/(1 + b))$ is stable, and the unique interior equilibrium is unstable; (F) for the other situations, the boundary equilibrium $(0, b/(1 + b))$ is stable, with no other interior equilibriums. The solid circles represent the stable equilibriums, and the hollow circle the unstable equilibrium. Gray lines indicate zero isoclines (straight lines denote $L_1$, i.e., $\frac{dp}{dt} = 0$, and curves $L_2$, i.e., $\frac{dh}{dt} = 0$), and other lines are trajectories from various initial values (directions indicated by arrows). For the parameters, we take: $c = 0.5$, $e = 0.1$, and $d = 0.1$ in (A)–(F); $\lambda = 0.8$ and $\mu = 0.12$ in (A); $\lambda = 0.8$ and $\mu = 2/30$ in (B); $\lambda = 0.2$ and $\mu = 2/30$ in (C); $\lambda = 0.8$ and $\mu = 0.03$ in (D); $\lambda = 0.6$ and $\mu = 0.0299$ in (E); $\lambda = 0.4$ and $\mu = 0.03$ in (F).

Table 2
The equilibrium and stability of Eq. (2) and the metapopulation persistence.

<table>
<thead>
<tr>
<th>Parameter condition</th>
<th>Boundary equilibrium</th>
<th>Interior equilibrium</th>
<th>Metapopulation persistence</th>
</tr>
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<tbody>
<tr>
<td>$\delta &lt; \frac{1}{1+b}$</td>
<td>Unstable</td>
<td>(One) Stable</td>
<td>Persistence (Fig. 1A)</td>
</tr>
<tr>
<td>$\delta = \frac{1}{1+b}$ and $a &gt; (\beta + 1)^2$</td>
<td>Unstable</td>
<td>(One) Stable</td>
<td>Persistence (Fig. 1B)</td>
</tr>
<tr>
<td>$\frac{a}{1+b} &lt; \delta &lt; \min\left(\frac{a}{1+b}, \frac{1}{1+b}\right)$</td>
<td>Stable</td>
<td>(Two) Unstable stable</td>
<td>Persistence depends on the initial state (Fig. 1D)</td>
</tr>
<tr>
<td>$\delta = \sqrt{2} - 1$ and $a &gt; (\beta + 1)^2$</td>
<td>Stable</td>
<td>(One) Unstable</td>
<td>Extinction (Fig. 1E)</td>
</tr>
<tr>
<td>Others</td>
<td>Stable</td>
<td>(None)</td>
<td>Extinction (Fig. 1C&amp;F)</td>
</tr>
</tbody>
</table>

* Parentheses indicate the number of interior equilibrium.
\[
\frac{dp}{dt} = \frac{\lambda p^r}{(cp^r + \alpha)} - b p^2 + o(p^2)
\]  
(8)

i.e., \((p^*, h^*)\) is unstable if \(\alpha(1 - h^*)(xp^* + \beta + 1) = 1\) (Fig. 1E). □

Subsequently, a straightforward question here is whether the existence of periodic solution is possible. The following theorem shows that no periodic solution exists in Eq. (2).

**Theorem 4.** For Eq. (2), the existence of periodic solution is impossible.

**Proof.** The Dulac function is defined as \(\psi(p, h) = 1/p\), and notice that

\[
\frac{\partial(\psi P)}{\partial p} + \frac{\partial(\psi Q)}{\partial h} = -\left(c + \frac{\lambda p + \mu + d}{p}\right) < 0
\]  
(9)

for all positive \(p\) and \(h\), where \(P(p, h) = dp/dt\) and \(Q(p, h) = dh/dt\). Thus, from The Bendixson–Dulac theorem, no periodic solution can exist in Eq. (2). □

Theorem 4 implies that, if there is a unique interior equilibrium and it is also locally asymptotically stable, then it must be globally asymptotically stable, because, in this situation, the unique boundary equilibrium is unstable (see Table 2 and Fig. 1A, B).

According to the above analysis, the dynamical properties of Eq. (2) can be summarized as: (i) if the boundary equilibrium \((0, \beta)\) (\(\beta + 1\)) is unstable, then there exists a unique interior equilibrium that is globally asymptotically stable (Fig. 1A, B); (ii) if the boundary equilibrium \((0, \beta + 1)\) is stable, then there are at most two interior equilibriums; (iii) no periodic solution can exist in the system. The situation (ii) further includes three scenarios: (a) the situation with no interior equilibrium is possible (Fig. 1C, F); (b) if there exists only one interior equilibrium, then it must be unstable saddle point (Fig. 1E); (c) if there are two interior equilibriums, the one is locally asymptotically stable but the other unstable (Fig. 1D).

All of these results are summarized in Table 2. The case with two interior equilibriums indicates a bi-stable system, and thereby the fate of metapopulation depends on its initial size, i.e., a threshold phenomenon.

The relationship between the equilibrium of metapopulation size and the internal restoration intensity \(\alpha(= \lambda/d)\) is plotted in Fig. 2. If the value of \(\beta(=\mu/d)\); the external restoration intensity) is small \((\beta < \delta/(1 - \delta))\), the bi-stability will occur when the internal restoration intensity \(\alpha\) is high (Fig. 2A, B). No bi-stability exists if \(\beta \geq \delta/(1 - \delta)\) (Fig. 2C, D). Similarly, the relationship between the equilibrium of metapopulation size and the external restoration intensity \(\beta\) is plotted in Fig. 3. If \(\alpha\) equals zero (or small), the system of Eq. (2) is essentially similar to the Levins model, with \(\beta(=\beta + 1)\) analogous to \(h\) as in Eq. (1) (Fig. 3A, B). If \(\alpha\) is large (determined by Eq. (C6) in Appendix C), the bi-stability will occur when \(\beta\) is moderate (Fig. 3C, D). Therefore, strong habitat restoration induced by the organism itself plays a crucial role to trigger the bi-stability, whereas the intensity of external restoration is a key to eliminate the bi-stability.

4. Discussion

In this paper, following Levins’ [3,4] patch occupancy model of metapopulations, we developed a theoretical model that describes not only metapopulation dynamics but also habitat dynamics induced by two processes of habitat destruction and restoration. The internal habitat restoration induced by the organism itself is separated from the external restoration by other biotic and abiotic causes. The positive feedback relationship between the habitat and metapopulation dynamics has been revealed by a full stability analysis of the system. The outcome implies that the organism-environment interaction can affect the persistence of metapopulation significantly to a certain degree (reflected by the relative strength of habitat restoration, \(\alpha\) and \(\beta\)). If the habitat destruction is inevitable, then the habitat restoration possibly driven by the organism itself and other conservation efforts are crucial for the metapopulation persistence (see Table 2 for details).

The evaluation of the persistence of a species inhabiting fragmented habitat is important in ecology [22]. Multiple factors, such as habitat destruction [23], environmental stochasticity [24], demographic stochasticity [5,25], dynamical complexity [26], behavior evolution [27,28] and rapid evolution [29], have been identified to be able to affect the persistence of metapopulations. In addition, our results suggested that the internal habitat restora-

![Fig. 2](https://example.com/image)
tion by the organisms can also promote the persistence of their metapopulations and thus increases the metapopulation size. However, whether a biological system can restore to its initial status depends not only on its capacity of habitat restoration but also on the level of habitat destruction. A highly destructed habitat could have potentially altered the system equilibrium from positive to extinction, making small-scale restoration in vain.

Internal restoration induced by the organism itself is the trigger of the bi-stability (or the threshold phenomenon) of the system, whereas the external restoration can eliminate this bi-stability (Fig. 3). This threshold phenomenon could arise from the density-dependent negative growth rate at low metapopulation size, indicating a sufficient number of suitable patches or metapopulation size for persistence. Other mechanisms that can cause such negative growth rate and threshold phenomenon have been identified, such as the Allee effect [30,31] and the rescue effect [2]. The organism-environment feedback is a crucial interaction in ecosystems [10,32], yet often neglected in biological conservation. This positive feedback (i.e. the capacity of habitat restoration) can strongly affect the distribution and the abundance of species [32], the consequence of competition [16], the structure and functioning of ecosystem [33–35], evolutionary directions [36], the maintenance of polymorphism [17,18], and population dynamics [37–39]. Our study here suggests that organism-induced habitat restoration can profoundly affect the persistence of metapopulations, change stable systems into bi-stable ones, cause the threshold phenomenon, and thus should merit further investigations.

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Appendix A. Derivation of Eq. (2) from the Markov stochastic process

Consider a total number of z patches in the landscape, including a number of x occupied patches and a number of y suitable patches \((0 \leq x \leq y \leq z)\); we can thus define a continuous-time Markov process with the following non-zero transition rates:

<table>
<thead>
<tr>
<th>Event</th>
<th>Rate</th>
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<tbody>
<tr>
<td>((x, y) \rightarrow (x + 1, y))</td>
<td>(c'nx(y - x)/z)</td>
</tr>
<tr>
<td>((x, y) \rightarrow (x - 1, y))</td>
<td>(ex)</td>
</tr>
<tr>
<td>((x, y) \rightarrow (x, y + 1))</td>
<td>((\mu + rmx/z)(z - y))</td>
</tr>
<tr>
<td>((x, y) \rightarrow (x, y - 1))</td>
<td>(d(y - x))</td>
</tr>
<tr>
<td>((x, y) \rightarrow (x - 1, y - 1))</td>
<td>(dx)</td>
</tr>
</tbody>
</table>

where \(m\) is the number of emigrants from each occupied patch, \(c'\) the colonization coefficient and \(e\) the extinction rate. It describes the following Markov process: emigrants randomly land into patches regardless of the patch status, and thus each gets \(mx/z\) immigrants; for an empty suitable patch, the probability of successful colonization is \(c'mx/z\); for an occupied patch, the patch status remains; for an unsuitable patch, the immigrants contribute to restoring the patch by organism-induced way, and the restoration rate of an unsuitable patch is thus \(\mu + rmx/z\). The above Markov process can also be approximated for large z using the following two-dimensional ordinary differential equations [40]:

\[
\frac{dx}{dt} = c'm(y - x) - ex - dx \\
\frac{dy}{dt} = (r\frac{mx}{z} + \mu)(z - y) - d(y - x) - dx.
\] (A2)

If we let \(c = c'm\), \(\lambda = rm\), \(p = x/z\) (the fraction of occupied patches) and \(h = y/z\) (the fraction of suitable patches), Eq. (A2) becomes Eq (2) in the main text.

Appendix B. Proof of Theorem 2

If \(\delta = \beta/(\beta + 1)\), then one of the eigenvalues of the Jacobian matrix \(J_{\text{boundary}}\) must be zero. Let \(\bar{p} = p\) and \(\bar{h} = h - \beta'y - x\gamma^2\), where \(\gamma = 1/(\beta + 1)\), then Eq. (2) can be equivalently expressed as:

\[
\frac{dp}{dt} = c(x\gamma^2 - 1)p^2 + cp\bar{h} \\
\frac{d\bar{h}}{dt} = -(\mu + d)\bar{h} - \pi(\bar{p}, \bar{h}).
\] (B1)

where \(\pi(\bar{p}, \bar{h}) = (\lambda + cx\gamma^2)p\bar{h} + |x\gamma^2(\lambda + c) - cx\gamma^2|p^2\). From the center manifold theory [21], there exists a local center manifold.
\[ \dot{h} = \varphi(p) \] with \( \varphi(0) = 0 \) and \( d\varphi(0)/dp = 0 \). Notice that \( \dot{h} = \varphi(p) \) satisfies

\[ \frac{d\varphi(p)}{dp} [c(x^2 - 1)p^2 + c\varphi(p)] = -(a + d\varphi(p) - \pi(p, \varphi(p))). \]  

(B2)

and that \( \varphi(p) \) can be solvable. Thus, the stability of \( (0, \beta/(\beta + 1)) \) is completely determined by the reduced system:

\[ \frac{dp}{dt} = c(x^2 - 1)p^2 + c\varphi(p). \]  

(B3)

Since \( c\varphi(p) = o(p^2) \), the boundary equilibrium \( (0, \beta/(\beta + 1)) \) is a saddle point if \( x^2 \neq 1 \). Specifically, \( (0, \beta/(\beta + 1)) \) is unstable for positive trajectories if \( x^2 > 1 \) (Fig. 1B), and it is a local attractor for positive trajectories if \( x^2 < 1 \) (Fig. 1C). For the case with \( x^2 = 1 \), the solution of Eq. (B3) can be expressed as \( \varphi(p) = -x^2/p^2 + o(p^2) \), and Eq. (B3) can be rewritten as

\[ \frac{dp}{dt} = -cx^2/p \]  

(B4)

Thus, \( (0, \beta/(\beta + 1)) \) is locally asymptotically stable if \( x^2 = 1 \). ☐

Appendix C. Interior equilibrium of Eq. (2)

The interior equilibrium of Eq. (2) must be the solution of the following equations

\[ c(h - p) - (e + d) = 0 \]

\[ (\lambda + \mu)(1 - h) - dh = 0. \]  

(C1)

Notice that these equations can be equivalently expressed as

\[ x^2 - (\alpha(1 - \delta) - \beta - 1)p + (\beta + 1)\delta - \beta = 0. \]  

(C2)

Thus, Eq. (2) has at most two interior equilibriums, and the necessary condition for the existence of an interior equilibrium is \( (\alpha(1 - \delta) + \beta + 1)^2 - 4\alpha \geq 0 \). In order to determine whether there exists an interior equilibrium, let

\[ f(p) = xp^2 - (\alpha(1 - \delta) - \beta - 1)p + (\beta + 1)\delta - \beta. \]  

(C3)

It is straightforward to see that

\[ f(0) = (\beta + 1)\delta - \beta \]

\[ f(1) = (\alpha - \beta) + 1, \]  

and that the minimum point of \( f(p) \) is

\[ \hat{p} = \frac{\alpha(1 - \delta) - \beta - 1}{2\alpha}. \]  

(C5)

Since a necessary condition for the existence of a positive equilibrium is \( e + d < 0 \) (i.e., \( \delta < 1 \)) [41,42], we have \( f(1) > 0 \). This implies that (i) if \( f(0) < 0 \), i.e. \( \delta > \beta/(\beta + 1) \), then there could be only one interior equilibrium (Fig. 1A); (ii) under the condition \( |\alpha(1 - \delta) + \beta + 1|^2 - 4\alpha \geq 0 \), if \( f(0) > 0 \) and \( 0 < \beta < 1 \), i.e.,

\[ \frac{\beta}{\beta + 1} < \delta < \min \left( \frac{\beta - 1}{\alpha}, \frac{\sqrt{(\alpha - 1)^2 + \beta}}{\alpha} \right), \]  

(C6)

two interior equilibria can occur (Fig. 1D); (iii) for \( f(0) = 0 \), i.e., \( \delta = \beta/(\beta + 1) \), only one interior equilibrium occurs when \( x > (\beta + 1)^2 \) (Fig. 1B), and no interior equilibrium exists when \( x < (\beta + 1)^2 \) (Fig. 1C); (iv) for \( |\alpha(1 - \delta) + \beta + 1|^2 - 4\alpha = 0 \), i.e. \( \delta = ((\alpha - 1)^2 + \beta)/\alpha \), there is only one interior equilibrium when \( x > (\beta + 1)^2 \) (Fig. 1E); and (v) for other situations, there is no any interior equilibrium (Fig. 1F). These results were summarized in Table 2.