Patterns of invasive species spread in a landscape with a complex geometry

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**ABSTRACT**

Patterns and rates of invasive species spread have been a focus of attention for several decades. Majority of studies focused on the species proliferation in a relatively uniform “open space” thus leaving aside the effects of the landscape geometry as given by size and shape of inaccessible areas. In this paper, we address this issue by considering the spatiotemporal dynamics of an alien species in a domain where two large uniform habitats are connected by a narrow corridor. We consider the case where the species is originally introduced into one of the habitats but not to the other. The alien species is assumed to be affected by a predator, so that mathematically our system consists of two coupled diffusion–reaction equations. We show that the corridor tends to slow down the spread; it takes the alien population an extra time to penetrate through the corridor, and this delay time can be significant in the case of patchy spread. We also show that a sufficiently narrow corridor blocks the spread; simple analytical estimates for the critical width of the corridor are obtained. Finally, we show that the corridor can become a refuge for the alien population. If considered on a longer timescale that includes species adaptation and/or climate change, the corridor may then become a source of a secondary invasion.

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1. Introduction

Biological invasion is a phenomenon that has a variety of important implications for ecology (in particular, through species extinctions and biodiversity loss), environment, human wellbeing, agriculture/aquaculture, and sometimes industries beyond agriculture (Keller et al., 2006; Pimentel, 2002; Sandlund et al., 2001). For these reasons, it has been a focus of intense empirical and theoretical research for several decades (Drake et al., 1989; Richardson, 2011; Williamson, 1996). Mathematical modelling is widely accepted as an efficient tool to study biological invasion (Hengeveld, 1989; Lewis et al., 2016; Shigesada and Kawasaki, 1997). Indeed, in the situation when replicated studies are hardly possible, mathematical models along with computer simulations provide a virtual laboratory where the effect of various factors can be refined and analyzed and different hypotheses can be tested safely and at a relatively low cost.

In studies on biological invasion, a central question is how the alien species proliferates into space away from the place of its introduction. In its turn, it evokes the question as to what is the pattern of spread. For several decades, a paradigm of invasive species spread was a travelling population front. A large amount of theoretical work has been done in order to evaluate the speed of the front propagation (Aronson and Weinberger, 1978; Fisher, 1937; Kolmogorov et al., 1937; Skellam, 1951) and to reveal how the propagation can be affected by various factors (Bell et al., 2009; Lewis and Kareiva, 1993; Owen and Lewis, 2001), in particular, by environmental heterogeneity (Andow et al., 1990; Keitt et al., 2001; Petrovskii, 1998; Shigesada et al., 1986). The travelling front paradigm was eventually complemented by an alternative pattern of spread known as patchy invasion (Jankovic and Petrovskii, 2013; Petrovskii et al., 2002, 2005b; Morozov et al., 2006) which is often observed in nature (Davis et al., 1998; Liebhold et al., 1992; Mack, 1981) and predicted by various dispersal-growth models (Lewis, 2000; Mistro et al., 2012; Rodrigues et al., 2015; Shigesada et al., 1995). In the patchy invasion scenario, there is no population front and species spread into space occurs through the dynamics of separate disconnected population patches.

Whether it is a population travelling front or a patchy spread, previous studies on biological invasions have overwhelmingly focused on an idealized case of species spread in an unbounded space. (In case of a simulation study, the domain is bounded but...
can be chosen ‘sufficiently large’ in order to imitate an unbounded space.) Even that the effect of environmental heterogeneity was considered in a number of studies, e.g. see the references above, the role of inaccessible areas and hence the effect of landscape geometry on the rate of spread have remained poorly understood. Meanwhile, such a role is likely to be very important. One real world example is the invasion of grey squirrels in the UK. Whilst they successfully colonized most of England and Wales, the rate of their spread being in a good agreement with predictions of relevant mathematical models (Bell et al., 2009; Okubo et al., 1989), they have largely failed to spill over to central and northern Scotland, and the landscape geometry – as given by large areas that are not accessible to squirrels such as swamps or high mountain ranges – is thought to be a reason for that (White, 2013). As another example, it was found in a study on the invasion of nutria in Iran that about one third of the country could potentially be colonized (Farashi and Najafabadi, 2015). In reality, however, having been introduced about one hundred years ago to the sub-Caspian region, this species remains confined to largely the same area. A closer look reveals that different parts of the potential range are poorly connected (e.g. due to the effect of mountain ranges inaccessible to nutria), which is likely to limit species dispersal. In this paper, we consider the effect of the landscape geometry on the invasive species spread by means of mathematical modelling and computer simulations. The “complexity” of the landscape geometry (as is relevant in the context of this study) is represented by a dumbbell-like, “H-shaped” spatial domain consisting of two large habitats connected by a narrow passage or corridor. The initial conditions correspond to the situation where the alien species is present in

![Fig. 1. Snapshots of the spatial distribution of prey (i.e. the alien species) illustrating different scenarios of invasive spread as predicted by Eqs. (5) and (6) obtained in a square domain 200 x 200 for parameters (a) δ = 0.63, (b) δ = 0.51, (c) δ = 0.43 and (d) δ = 0.37. Other parameters are given in the text. Here and below yellow color stands for high population density, green and light blue for intermediate values, dark blue for zero density. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image)
one of the habitats but not in the other. We are especially interested in the following questions:

- How much a narrow passage can slow down the spread? Can a sufficiently narrow passage block the spread completely so that the invading population would never spill over to the other side?
- Depending on species traits, invasive spread into open space is known to follow a few different scenarios (see Fig. 1). Can the effect of the passage change the invasion scenario, e.g., turning a propagating population front into patchy spread (or visa versa)?
- Can the effect of the passage depend on the invasion scenario (e.g., slowing down the spread in one case but not having much effect in another)?

It has been shown in our earlier work (Alharbi and Petrovskii, 2016) that the success of the species invasion into the second habitat depends on the width and length of the corridor as well as demographic parameters such as the maximum growth rate and the strength of the Allee effect. However, the effect of the dynamical complexity on the invasive species spread, in particular due to interspecific interactions and pattern formation, has remained obscure. Here we show that the dynamical complexity can affect the invasion success significantly. In particular, we show that the effect of the corridor on the invasion success is significantly different depending on whether the invasive species spreads following the travelling population front scenario or the patchy invasion scenario. We also show that a corridor that blocks the species spread can become a refuge for the alien species and can subsequently become a source of a secondary invasion.

2. Model

The choice of the mathematical model is a somewhat subtle issue (Maynard Smith, 1974; Pascual, 2005; Petrovskii and Petrovskaia, 2012). From a biological standpoint, it might be tempting to account in detail for a variety of biological interactions that the invasive species is experiencing during its spread, e.g., interactions with all species of the native community. A corresponding model is likely to be very complicated. Such a model would be difficult to parameterize as the information about the feedbacks and functional responses as well as parameter values is usually meagre and incomplete. Moreover, the higher the complexity of the model is, the more difficult it may be to reveal the general properties of the phenomenon as they may become obscure by specific details. Since in this paper our goal is to look into general properties of the dynamics rather than to study a particular case, the model should not be too complicated. However, it should not be too simple either as some important features of the invasion dynamics – such as, for instance, pattern formation (Jankovic and Petrovskii, 2013; Malchow et al., 2008; Petrovskii et al., 1995, 1997; Shigesada et al., 1995) – can be missing altogether due to the insufficient complexity of the model.

We therefore look for a model of an intermediate complexity. Arguably, an appropriate model is a prey-predator system. On the one hand, it obviously takes into account only some common, basic interactions, i.e. how the population dynamics of the alien species (regarded in this context as a prey) is affected by its predator where the predator can be a native species or can be introduced as a biological control agent, cf. (Kenis and Vaamonde, 1998; Petrovskii and McKay, 2010; Petrovskii et al., 2005b). On the other hand, a spatially explicit prey-predator model allowing for the population dispersal is known to have a complicated spatiotemporal dynamics (Lewis et al., 2016; Malchow et al., 2008), its “realistic” properties being consistent with at least some field observations (Jankovic and Petrovskii, 2013; Owen and Lewis, 2001).

A general prey–predator system is described by the following coupled diffusion–reaction equations (Murray, 1989; Owen and Lewis, 2001; Sherratt et al., 1995):

$$\frac{\partial u(r, t)}{\partial t} = D_u \nabla^2 u(r, t) + f(u) - f(u, v),$$  \hspace{1cm} (1)

$$\frac{\partial v(r, t)}{\partial t} = D_v \nabla^2 v(r, t) + \kappa f(u, v) - MV,$$  \hspace{1cm} (2)

where $u(r, t)$ and $v(r, t)$ are, respectively, the prey and predator population densities at time $t$ and position $r = (X, Y)$. $D_u$ and $D_v$ are the corresponding diffusion coefficients, $\kappa$ is the food utilization coefficient. Function $f(u, v)$ describes predation, and $M$ is the predator mortality rate. A particular choice of functions $f(u, v)$ may vary depending on the species traits. We assume that the predator trophic response is of the Holling type II and choose the parametrization accordingly (Murray, 1989):

$$f(u, v) = \frac{AUV}{U + H},$$  \hspace{1cm} (3)

where $A$ quantifies the predation rate and $H$ is the half-saturation prey density.

With regard to the prey growth, there is considerable evidence that the Allee effect is common in the population dynamics (Courchamp et al., 1999, 2008). Correspondingly, we assume that the alien species is affected by the strong Allee effect and consider the following parametrization for its growth rate (Lewis and Kareiva, 1993):

$$F(u) = G(U - U_A)/(K - U),$$  \hspace{1cm} (4)

where $G$ is a coefficient related to the maximum per capita growth rate $g$ as $G = 4g/(K(1 - U_A))$ (Lewis and Kareiva, 1993), $K$ is the carrying capacity, and the Allee threshold density $U_A (0 < U_A < K)$ is a parameter quantifying the strength of the Allee effect (Lewis and Kareiva, 2001; Wang and Kot, 2001).

Eqs. (1) and (2) with (3) and (4) contain a large number of parameters, which makes their numerical investigation cumbersome. However, if we choose new, dimensionless variables as $u = U/K$, $v = V/(\kappa K)$, $t = \alpha t$ where $\alpha = \kappa K[H, \gamma = \lambda(\alpha D_1)1/2]$ and $y = \lambda(\alpha D_1)1/2$, Eqs. (1) and (2) turn to the following:

$$\frac{\partial u(x, y, t)}{\partial t} = \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) + \gamma u(u - \beta)(1 - u) - \frac{uv}{1 + \alpha u},$$  \hspace{1cm} (5)

$$\frac{\partial v(x, y, t)}{\partial t} = \varepsilon \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) + \frac{uv}{1 + \alpha u} - \delta v,$$  \hspace{1cm} (6)

where $\alpha = K[H, \beta = U_A/K, \gamma = \lambda(\alpha K)]$, $\delta = M/\alpha$, and $\varepsilon = D_2/D_1$ are new (dimensionless) parameters. Properties of dimensionless solutions $u$ and $v$ therefore depend on five dimensionless combinations rather than on each of the nine original parameters separately.

Biological invasion usually starts with an introduction of a (small) number of individuals of an alien species into a certain location inside a new ecosystem (Shigesada and Kawasaki, 1997). Immediately after the introduction, the population density of the alien species is only non-zero inside a certain (small) area. Therefore, in order to account for the biological invasion, Eqs. (5) and (6) must be supplemented with initial conditions described by functions of a finite support. Here we consider the initial conditions as follows:

$$u(x, y, 0) = u_0$$ if $x_{11} < x < x_{12}$ and $y_{11} < y < y_{12}$ otherwise $u(x, y, 0) = 0.$  \hspace{1cm} (7)
\[ v(x, y, 0) = v_0 \text{ if } x_{21} < x < x_{22} \text{ and } y_{21} < y < y_{22} \]
where \( v_0, v_0 \) are the initial prey and predator densities, and parameters \( x_i \) and \( y_i \) (\( i = 1, 2 \)) determine the initially occupied area.

3. Simulations and results

Eqs. (5) and (6) with the initial conditions (7) and (8) were solved numerically by finite-differences. In most cases we used the simple explicit scheme. The mesh steps were chosen as \( \Delta x = \Delta y = 0.4 \) and \( \Delta t = 0.03 \); it is readily seen that they satisfy the Courant-Friedrichs-Lewy stability condition (Richtmyer and Morton, 1967).

Having performed simulations for a few other values of the mesh steps, we checked that the above values are sufficiently small to avoid numerical artifacts.

Although our main goal in this paper is to study the effects of a complex geometry, we begin with a simple case when Eqs. (5) and (6) are considered in a square domain. The domain has size \( L \) so that \( 0 < x < L \) and \( 0 < y < L \). At the boundary of the domain the Neumann-type no-flux conditions are imposed. For any given parameter set, the simulation is stopped either just before or immediately after the spreading population reaches the domain boundary. In this way, the effect of the boundary conditions on the population dynamics inside the domain is minimized; correspondingly, the population spread imitates the spread in an open (unbounded) space.

Depending on the parameter values such as the predator mortality rate \( \delta \) and/or the Allee threshold \( \beta \), the prey–predator system (5) and (6) is known to exhibit different patterns of the population spread (Malchow et al., 2008; Morozov et al., 2006; Volpert and Petrovskii, 2009). Having chosen \( \delta \) as the controlling parameter, a brief visual overview of the patterns is shown in Fig. 1 where the results were obtained in a square domain \( 200 \times 200 \) for parameters \( \epsilon = 1, \gamma = 3.9, \alpha = 0.1, \beta = 0.2 \), the initial conditions (7) and (8) with \( v_0 = 1, v_0 = 0.2, x_{11} = 85, x_{12} = 105, x_{21} = 85, x_{22} = 95, y_{11} = 100, y_{12} = 105, y_{21} = 95 \) and \( y_{22} = 115 \), and different values of \( \delta \).

Here and below, only the distribution of prey is shown as the distribution of predator exhibits similar features. Unless the initially inhabited domain is very small so that the alien population goes extinct due to the strong Allee effect (Lewis and Kareiva, 1993), a somewhat trivial case that we do not consider here, the alien population spreads into the empty space away from the place of its introduction. This spread can follow the population front propagation scenario (Fig. 1a–c) with or without pattern formation in the wake or the patchy invasion scenario (Petrovskii et al., 2002, 2005b) (Fig. 1d) where the population spread is not preceded by a travelling front propagation.

A practically important question is how high is the rate of invasive spread. It can be rephrased as follows: for a given location of the alien species introduction, how fast the invading species will arrive at a given location elsewhere? In a spatially uniform system, this question is linked to the question about the speed of the propagation of the travelling population front2: indeed, once the speed is known, a common sense estimate is \( \text{time} = \text{distance} / \text{speed} \). However, it is intuitively clear that this simple argument is unlikely to remain valid in a case where the invasive spread takes place in a landscape of a complicated geometry where the spread can possibly be slowed down or block altogether by, for instance, the existence of narrow valleys or ‘corridors’.

In order to make an insight into possible effects of the corridors, we now consider a domain of a more complex shape, namely, H-shaped where two habitats (subdomains) of a rectangular shape are connected by a narrow passage or corridor. For the initial conditions, we consider the situation where the alien species is introduced to a certain location close to the center of the left-hand side habitat (which we for convenience also call Habitat 1); see Fig. 2. For the boundary conditions, we consider the Neumann-type zero-flux condition along the whole boundary. One can expect that during the first stage of invasion, i.e. before the invading population reaches the boundary of Habitat 1, the dynamics should be essentially the same as in a domain of a simple shape or in an infinite space. Once the invading population reaches the entrance to the corridor, however, it becomes a new, different problem and the question arises as to whether the population spills over to the right-hand side habitat (which we call Habitat 2) for given length and width of the corridor and for a given pattern of spread, cf. Fig. 1.

The case of the population spread with a uniform spatial population distribution behind the invasive front (Fig. 1a) is similar to the invasive spread in a single species model considered in Alharbi and Petrovskii (2016); we do not reconsider it here. We therefore begin with a more interesting case where invasion into a homogeneous space takes place via the propagation of the population front with spatiotemporal pattern formation in the wake (cf. Fig. 1b). The results shown in Fig. 3 are obtained for the parameters of the corridor \( \alpha_1(\lambda, \nu) = (10, 2.8) \) where \( \lambda \) is the length and \( \nu \) is the width. In the first stage of invasion, the population front propagates fast, apparently with the same speed as in an infinite space (Lewis et al., 2016). Once it reaches the entrance to the corridor, the spread slows down considerably; it takes the population about 100 extra time units to penetrate through the corridor, cf. the top-right and middle-right panels in Fig. 3. Eventually, the population spills over to Habitat 2 and invade over the rest of the domain, see the bottom row in Fig. 3.

Interestingly, the corridor affects the pattern of spread. The spread of the invading population in Habitat 1 is accompanied by the formation of irregular asymmetric patterns in the wake of the propagating front, which apparently is a result of the asymmetry in the initial conditions. However, the population distribution behind

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2 The situation is somewhat more complicated in the case of patchy invasion or in the case where the invasive species employs a long-distance dispersal. In the former case there is no front and hence the definition of the speed is somewhat ambiguous (Morozov et al., 2006), in the latter case the front can have variable speed (Kot et al., 1996).

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Fig. 2. A sketch of the H-shaped domain consisting of Habitat 1 at the left and Habitat 2 at the right connected by a narrow passage (corridor). The alien species is introduced in Habitat 1, the initially inhabited area is shown by a small yellow rectangle. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
the front in Habitat 2 exhibits a perfect circular symmetry, e.g. see the bottom-left panel in Fig. 3. However, spilling over after a delay caused by the corridor is not the only possibility. The system can also exhibit a qualitatively different dynamics resulting in an invasion failure. Simulation results (not shown here for the sake of brevity) obtained for different parameters of the corridor reveal that the dynamics depend strongly on the size of the corridor, in particular on its width. An increase of the corridor width will not change the dynamics much resulting only in a decrease of the time required for the population to penetrate through. However, a decrease of the corridor width will eventually result in invasion blocking: as soon as the width is smaller than a certain critical value, the invading population is blocked in the passage and never spills over to Habitat 2.

In order to check whether the above situation is general or pattern-specific, we then performed simulations for a value of $\delta$ where the species spread in an open space takes place via the
propagation of a solitary ring (see Fig. 1c). The results are shown in Fig. 4. It is readily seen that the population dynamics follows a similar scenario. In the first stage, the population spreads away from the place of its original introduction in the same manner as in an open space. Once the front reaches the entrance to the passage, the spread slows down; it takes it about 40 time units to penetrate through to enter Habitat 2. (Interestingly, this time appears to about twice shorter than in the previous case.) Once it spills over, the population spreads over Habitat 2 in a similar way as before. One difference from the corresponding pattern of spread in an open space is that a fraction of the population settles down inside the passage (this is well seen in the middle row of Fig. 4) and works as a source to generate a sequence of population fronts (spreading rings) into both habitats. As well as above, a successful spill-over through the corridor only happens if its width is not too narrow. Once the width becomes less than a certain critical value, the population is blocked in the passage and never penetrates to the other side.

Finally, we performed simulations for the value of $\delta$ where the alien population spreads into space following the patchy invasion scenario (see Fig. 1d). The results are shown in Fig. 5. The dynamics exhibit features similar to those observed in Figs. 3 and 4. The
population first spreads from the place of its original location in exactly the same manner as it would happen in a domain of a simpler shape or in an open space. Once it approaches the entrance to the passage, it is delayed for a considerable time before it spills over to Habitat 2. Note that the delay is two orders of magnitude longer than it was in the previous cases. An explanation of this lies in the absence of the population front: because of the chaotic movement of the patches in space (Morozov et al., 2006), it takes an extra time for a patch to hit the entrance to the passage compared to the directional propagation of the front. Eventually, if the corridor's width is greater than a certain critical value, the population spreads through the passage and invades over Habitat 2 (as in the bottom row of Fig. 5). However, in case the corridor is not wide enough the population is blocked and never penetrates to the other side (not shown here).

3.1. Propagation blocking

In order to make a more detailed insight into the existence of the corridor's critical width and how its value depends on the
predator mortality $\delta$ (considered as a controlling parameter) and hence on the type of species spread, we performed extensive computer simulations with different values of $w$ and $\delta$ keeping other parameters the same as above. For each parameter set, the code was run up to $t = 4000$. The results are shown in Fig. 6. We therefore observe that the critical width exhibits a non-monotone dependence on the predator mortality but tends to increase with a decrease in $\delta$.

An analytical estimation of the critical width appears to be possible. Using the comparison principle for nonlinear partial differential equations (Protter and Weinberger, 1984; Volpert and Khudyaev, 1985), it is readily seen that the following single-species model:

$$\frac{\partial u(x,y,t)}{\partial t} = \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) + \gamma u(u - \beta)(1 - u),$$

i.e. Eq. (5) without the last term in the right-hand side, is the upper bound for the prey–predator system (5) and (6). However, for the single-species model a simple analytical estimate of the critical width can be easily obtained. Indeed, once the front arrives to the end of the passage and starts spilling into Habitat 2, it is of approximately circular shape (consistent with the numerical simulations above, e.g. see the middle–right panel in Fig. 3 and the middle–right panel in Fig. 5). For a circular propagating front of the radius $R$ in a 2D system, its speed $c(R)$ is known to be related to the speed $c$ of the corresponding 1D front as

$$c(R) = c - \frac{D}{R},$$

(10)

e.g. see Volpert and Petrovskii (2009). For the front confined at the exit of the passage, $R = w/2$. Since the front is blocked, $c(R) = 0$. Eq. (10) then turns into the following:

$$c - \frac{2D}{w} = 0.$$  

(11)

For a scalar diffusion–reaction equation with the population growth parameterized by a cubic polynomial, the analytical expression for the speed of the front propagation is well-known:

$$c = \sqrt{\frac{DC}{2}(K - 2U_A)},$$

(12)

e.g. see Lewis et al. (2016). From (11) and (12), we therefore obtain:

$$w_{cr} = \frac{1}{C_0} \left( \frac{K - 2U_A}{C_0} \right),$$

(13)

or, in dimensionless variables,

$$w_{cr} = \frac{2\sqrt{2}}{(1 - 2\beta)\sqrt{\gamma}}.$$  

(14)

For the parameters of Fig. 6, the estimate (14) gives $w_{cr} \approx 2.3$ which agrees well with the simulation results.

Now, a question can arise here as to how sensitive the above results are to the boundary conditions along the corridor. Indeed, it is well known that the critical behaviour of diffusion–reaction systems is generally sensitive to the boundary conditions; for instance, the solution of the critical size problem is essentially different in case of the Neumann (zero-flux) and Dirichlet (zero-density) conditions. In order to make an insight into this issue, we now consider the same H-shaped domain as above but with the Dirichlet zero-density conditions along the passage boundary; see Fig. 7. We mention here that, compared to the zero-flux condition, the zero-density boundary condition corresponds to a somewhat different ecological situation. Instead of a narrow canyon (connecting two large valleys) where the population density can build up in the vicinity of the canyon wall, one can think, for instance, of a narrow ridge (e.g. connecting two mountain plateaus) where a casual step over the cliff would result in death – hence, the zero population density.

Having performed extensive computer simulations in the domain with the zero-density condition in the passage (Fig. 7), we obtain that the dynamics of the system is qualitatively the same as in the previous case of the zero-flux conditions. In particular, we obtained that for all invasion regimes (as in Fig. 1) the invading population spills over to Habitat 2 if the corridor width is large enough and is blocked if the corridor is too narrow. Dependence of the critical width on the predator mortality is shown in Fig. 8. Therefore, as well as in the previous case we observe non-monotone dependence of $w_{cr}$ on $\delta$ with a tendency for the critical width to increase with a decrease in $\delta$.

As well as above, it appears possible to obtain a simple analytical estimate of the critical corridor width, although the argument and calculations are different. We recall that the single-species model is an upper bound for the prey–predator model, so a

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**Fig. 6.** Parameter plane (w, $\delta$), other parameters are the same as above: For the parameters from above the green curve (obtained in simulations) the population spills over and invade Habitat 2, for the parameters from below the curve the population is blocked in the passage. Red line shows the analytical estimate (14). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

**Fig. 7.** Sketch of an H-shaped habitat with Dirichlet boundary conditions at the sides of the corridor. (For interpretation of the references to color in the text, the reader is referred to the web version of this article.)
and in the dimensionless variables:
\[ \text{w}_{cr} = \frac{2\pi}{(1 - \beta)\sqrt{\gamma}}. \]  \hfill (17)

For the parameters used in simulations, the estimate (17) gives the value \( \text{w}_{cr} \approx 4 \), which appears to be in good agreement with numerical results; see Fig. 8.

Note that the mechanism of spread blocking is somewhat different in the two cases. For the zero-density conditions at the boundary of the passage, the population is blocked inside the passage (in a manner similar to how a subcritical size of the inhabited domain results in the population extinction). For the zero-flux conditions, however, the population cannot be blocked inside (except for the trivial case where \( \beta > 0.5 \)); instead, its propagation is blocked at the exit of the passage.

Finally, we want to emphasize here that both (14) and (17) are sufficient conditions of propagation blocking but not criteria. Once \( w \leq \text{w}_{cr} \), the propagation will necessarily be blocked by the passage; however, \( w > \text{w}_{cr} \) does not guarantee that the population will spread into Habitat 2. In terms of the diagrams shown in Figs. 6 and 8, it means that the whole of the red line lies in the parameter range where the population spread is blocked in the prey–predator system (i.e. in the area marked by crosses).

### 3.2. Sensitivity of the patchy spread to the initial conditions

From the four invasion scenarios shown in Fig. 1, perhaps the patchy invasion poses the biggest challenge. The absence of a continuous propagating population front leads to a certain ambiguity in the definition of the extent of the invaded area (Morozov et al., 2006). Similarly, it introduces an uncertainty as to when exactly, for a given set of the initial conditions, the population reaches the entrance to the passage. It is not enough for the expanding ‘envelope’ (e.g. the location of the foremost patch, cf. (Morozov et al., 2006)) to reach the entrance to the corridor; in order for the population to start penetrating through the corridor, there must be one of the patches that actually hits the entrance. However, since the spatiotemporal population dynamics during the patchy invasion is known to be chaotic, the movement of the patches is effectively random. In its turn, the exact location of the patches depends on the initial population distribution. Therefore, one can expect that, altogether, the time that takes the population to arrive at the passage entrance and to penetrate to the other side can vary significantly.

In order to make a more quantitative insight into this issue, we therefore performed simulations for a few different initial conditions choosing the demographic parameters in the range where the patchy invasion occurs (Jankovic and Petrovskii, 2013; Morozov et al., 2006; Petrovskii et al., 2005b). Specifically, we use \( \gamma = 3.9, \alpha = 0.1, \beta = 0.2 \) and \( \delta = 0.37 \). The parameters of the corridor conditions of population propagation blocking in the model (9) will be a sufficient condition of the propagation blocking in model (5) and (6). Consider the case of a long corridor. Consider the situation where the invasive species has penetrated through the corridor and successfully spilled over to Habitat 2. In this case, in the large time limit the population density profile across the corridor does not depend on its position along the corridor as far as the position is sufficiently far away from the corridor ends. The problem then becomes one-dimensional as the population density depend only on the coordinate across the corridor but not along the corridor. However, if considered in the 1D settings, this turns into the classical critical size problem (e.g. see Section 3.1 in Lewis et al. (2016)), that is, how large the domain must be to ensure the survival of the population inside. For the growth rate described by a cubic polynomial, its analytical solution is unknown. However, if we consider an upper bound of the growth rate as
\[ yu(u - \beta)(1 - u) \leq gu \text{ for any } u \geq 0, \]  \hfill (15)
where \( g \) is the maximum per capita growth rate, see the lines below Eq. (4), the expression for the critical size in the corresponding linear problem is well known:
\[ \text{w}_{cr} = \pi \sqrt{\frac{D}{g}}. \]  \hfill (16)

Fig. 9. Snapshots of different initial predator population distributions used in simulations: (a) (IC-1) \( y_{11} = 55, y_{12} = 60, y_{21} = 50 \) and \( y_{22} = 70 \); (b) (IC-2) \( y_{11} = 32.8, y_{12} = 35.2, y_{21} = 30 \) and \( y_{22} = 40 \); (c) (IC-3) \( y_{11} = 27.8, y_{12} = 30, y_{21} = 25.2 \) and \( y_{22} = 35.2 \). The initial conditions for the prey and other parameters are given in the text.
are chosen as $(\lambda, w) = (10, 9.2)$. For the initial conditions, we fix the initial population densities as $u_0 = 1$ and $v_0 = 0.2$, and the initial distribution of prey as $x_{11} = 85.2, x_{12} = 105.2, x_{21} = 85.2, x_{22} = 95.2$, and consider a few different initial distribution of the predator; see Fig. 9.

For the above parameters and for each of the three initial distributions, the population spread from the place of its introduction generally follows the scenario shown in Fig. 5 (we do not show the results for the sake of brevity). We first consider the case with the zero-flux conditions on the passage boundary. In each simulation run, we pick the time required for the population to reach Habitat 2, e.g. the ‘moment’ when the population has penetrated through the passage and starts spilling over into the open space on the other side, e.g. as shown in the middle-right panel in Fig. 5. The results are presented in Table 1. It is readily seen that the time varies quite significantly; in fact, by an order of magnitude. We then repeat simulations for the same parameters and the same sets of initial conditions but imposing the zero-density conditions at the passage boundary. As well as in the

<table>
<thead>
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<th>Boundary conditions in the passage</th>
<th>IC-1</th>
<th>IC-2</th>
<th>IC-3</th>
</tr>
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<tbody>
<tr>
<td>Zero-flux (Neumann)</td>
<td>2340</td>
<td>460</td>
<td>5090</td>
</tr>
<tr>
<td>Zero-density (Dirichlet)</td>
<td>4450</td>
<td>3970</td>
<td>1240</td>
</tr>
</tbody>
</table>

For the above parameters and for each of the three initial distributions, the population spread from the place of its introduction generally follows the scenario shown in Fig. 5 (we do not show the results for the sake of brevity). We first consider the case with the zero-flux conditions on the passage boundary. In each simulation run, we pick the time required for the population to reach Habitat 2, e.g. the ‘moment’ when the population has penetrated through the passage and starts spilling over into the open space on the other side, e.g. as shown in the middle-right panel in Fig. 5. The results are presented in Table 1. It is readily seen that the time varies quite significantly; in fact, by an order of magnitude. We then repeat simulations for the same parameters and the same sets of initial conditions but imposing the zero-density conditions at the passage boundary. As well as in the

Fig. 10. Dynamics of the system resulting in the formation of a stable steady state distribution of the prey species inside the passage that survives after the invading population goes extinct in the rest of the domain. Parameters and the initial conditions are given in the text.
previous case, the spill-over time shows a considerable variation between different initial conditions.

3.3. Corridor as a refuge

Finally, we report on a curious invasion scenario that occurs in a certain range of the parameters of the corridor. Fig. 10 shows snapshots of the prey distribution obtained for the demographic parameters chosen as $\gamma = 6.25$, $\alpha = 0.1$, $\beta = 0.2$ and $\delta = 0.5$. The parameters of the corridor are $(\lambda, w) = (10, 2.8)$ and the initial conditions are given by (7) and (8) with $u_0 = 1.0$, $v_0 = 0.2$, $x_{11} = 85.2$, $x_{12} = 105.2$, $x_{21} = 85.2$, $x_{22} = 95.2$, $y_{11} = 55.2$, $y_{12} = 60$, $y_{21} = 50$ and $y_{22} = 70$. The boundary conditions are zero-flux along the whole boundary of the domain (including the passage). Apparently, for these parameters the population spreads into an open space via ring-shaped travelling fronts; see the top-left panel in Fig. 10. After the population has propagated through the passage, a number of travelling concentric fronts is produced on both sides of the passage; see the middle row in Fig. 10. Eventually, the fronts go away (cf. the bottom row in Fig. 10) and the alien population eventually goes extinct in the most of the domain – except for the passage. A close inspection reveals that there is a residual population that settled down inside the corridor; this is readily seen from Fig. 11 that also gives a magnified view of the passage. Since this residual population is still persistent in the passage after a considerable time ($t = 9000$ in Fig. 11), we assume that it corresponds to a stationary solution of Eqs. (5) and (6).

A question arises here as to what can be a response of this residual population to a change in the demographic parameters. Such a change can be a result of the alien species adaptation to their new environment, or it can be a result of some exogenous process such as the global climate change. In either case, the question is whether the population will remain confined inside the passage or it may start spreading.

Looking for an answer to the above question, we performed simulations using the residual population as the initial condition for another simulation run with a different value $\beta_1$ for the Allee threshold density. We obtain that for $\beta_1$ just slightly larger than $\beta$, e.g. $0.2 < \beta_1 < 0.23$, the population remains confined inside the passage and at any moment of time the spatial population distribution of the prey is similar to the one shown in Fig. 11. In case $\beta_1$ is sufficiently larger than $\beta$, e.g. $0.24 < \beta_1 < 0.3$, the residual population goes extinct. However, in case $\beta_1$ is sufficiently smaller than $\beta$, e.g. $0.1 < \beta_1 < 0.16$ the prey can start spreading away from the passage into both habitats 1 and 2. As just one example, Fig. 12 show the results obtained in the case where $\beta_1 = 0.1$. Note that the spread takes place following a different scenario compared to the original one; instead of the propagating ring (cf. top-left panel in Fig. 10), it now spreads through a propagating circular front with pattern formation in the wake. This is not surprising as $\beta$ is, along with the predator mortality, a parameter that controls the pattern of invasive spread (Morozov et al., 2006; Petrovskii et al., 2005a).

4. Discussion and conclusions

Factors affecting the patterns and rates of alien species spread have been a major focus of research for several decades. A vast majority of studies were concerned with the species proliferation into an “open space” – a relatively uniform environment where all areas, although not necessarily being equally favorable, are accessible to the invading population. However, a realistic landscape can often be strongly heterogeneous combining areas accessible for the invader with inaccessible ones. The effect of the landscape geometry (as given by the size and shape of the inaccessible areas) remains poorly understood. In this paper, we addressed this problem by considering a mathematical model where the population dynamics of an alien species takes place in an H-shaped domain consisting of two large uniform habitats connected by a narrow passage or corridor. For the initial conditions, we considered that the alien species is introduced in a (small) area inside one of the habitats. Our goal was to reveal the
effect of the corridor on the species spread, in particular whether the spread from the one habitat to the other could be slowed down or blocked. In terms of the real-world landscape, such a system could account for an alien species spread through a narrow valley or an istmus. To account for interspecific interactions, we considered a generic prey-predator model which is known to exhibit a rather complex, ‘realistic’ spatiotemporal dynamics and predicts a variety of different invasion scenarios (Lewis et al., 2016; Malchow et al., 2008).

In reply to the questions bullet-pointed in the introduction, we obtained the following answers:

- The corridor does slow down the spread. It takes the spreading population an extra time to penetrate through the passage. A sufficiently narrow corridor, i.e. its width being less than a certain critical value, blocks the spread so that the invasive population never spills over to the other habitat. Analytical estimates for the critical width are obtained, see Eqs. (14) and (17), that are in good agreement with numerical results;
- The corridor can modify the pattern of spread. An example of this is shown in Fig. 3 where the irregular spatiotemporal pattern emerging behind the propagating front in Habitat 1 changes to a regular (symmetric) spatial population distribution behind the front in Habitat 2. Interestingly, this is exactly opposite to the results of some earlier work (Smith et al., 2008) where the existence of inaccessible areas – but with a different geometry – changed a regular spatiotemporal pattern to chaos. The regularizing effect of the corridor in our study can be readily explained: having propagated through the narrow passage, the population distribution at the exit to Habitat 2 become symmetric and effectively works as a symmetric initial condition for the population spread in Habitat 2;
- The delay of the spread caused by the corridor is significantly different depending on the pattern of spread. In particular, we obtained that it can be about two orders of magnitude longer in case of the patchy spread compared to the spread by the propagating continuous front.

Our results seem to send a potentially important message to the invasion management specialists. Since the corridor tends to impede the invasive species spread, it might be regarded as a possible control strategy. Although it apparently implies a significant alteration of the environment (e.g. to make a natural corridor more narrow), which is likely to be very expensive, we mention here that this strategy is not unrealistic and some examples of large scale environment intervention with the goal to slow down the spread are well known (Sharov et al., 2002). Importantly, however, based on our results (especially see the last item from the above list) this cannot be a universal approach as its efficiency depends on the pattern of spread: whilst it slows down the invasion considerably in the case of patchy spread, the delay might be rather insignificant in the case of spread via a continuous front. In its turn, this observation highlights the need for a better understanding of the invasion patterns and their more efficient identification including advance adaptive monitoring (Petrovskii et al., 2014).

Interestingly, although the corridor tends to impede the invasive species spread, it also can promote the invasion if considered on a longer time scale. Once the time is long enough to include an environmental change or species adaptation (e.g. through phenotypic evolution), the corridor can become a source of a secondary invasion (see Section 3.3) hence bringing the alien species back even it may have already disappeared from the rest of the domain. If considered in the context of invasion management, this observation restricts the potential use of the corridor as a factor controlling invasion to a small-to-intermediate timescale.

From the point of alien species monitoring, it also means that a special attention should be paid to landscape features that can act as corridors, because they may become a refuge for the alien pest.

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