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The Characteristics of Epidemics and Invasions with Thresholds

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Received April 7, 1998

In this paper we report the development of a highly efficient numerical method for determining the principal characteristics (velocity, leading edge width, and peak height) of spatial invasions or epidemics described by deterministic one-dimensional reaction-diffusion models whose dynamics include a threshold or Allee effect. We prove that this methodology produces the correct results for single-component models which are generalizations of the Fisher model, and then demonstrate by numerical experimentation that analogous methods work for a wide class of epidemic and invasion models including the S-I and S-E-I epidemic models and the Rosenzweig–McArthur predator–prey model. As exemplary application of this approach we consider the atto-fox effect in the classic reaction–diffusion model of rabies in the European fox population and show that the appropriate threshold for this model is within an order of magnitude of the peak disease incidence and thus has potentially significant effects on epidemic properties. We then make a careful re-parameterisation of the model and show that the velocities calculated with realistic thresholds differ surprisingly little from those calculated from threshold-free models. We conclude that an appropriately thresholded reaction–diffusion model provides a robust representation of the initial epidemic wave and thus provides a sound basis on which to begin a properly mechanistic modelling enterprise aimed at understanding the long-term persistence of the disease.

1. INTRODUCTION

The spatial spread of newly introduced species or diseases is a subject of continuing interest to both theoreticians and empiricists. One strand of theoretical developments (e.g., Skellam, 1951; Kendall, 1965; Mollison, 1972; Murray, 1989) built on the pioneering work of Fisher (1937), who used a logistic-based reaction–diffusion model to investigate the spread of an advantageous gene in a spatially extended population. With initial conditions corresponding to a spatially localised introduction, such models predict the eventual establishment of a well-defined invasion front which divides the invaded and uninvaded regions and moves into the uninvaded region with constant velocity.

Provided that very small populations grow in the same way as finite ones, the velocity at which an invasion front moves is set by the rate of divergence from the (unstable) invader-free state, and can thus be determined by linear methods (e.g., Murray, 1989). These techniques have been refined by Thieme (1979), van den Bosch et al. (1990), and Kot et al. (1996), who used a closely related renewal equation formalism to facilitate the inclusion of latent periods and more general and realistic transport models. All the analytic work and most of the simulations reported in these papers depend on the assumption that the invaders do not exhibit any special dynamics at very low densities. Where this assumption is not justified, for example when there is an Allee effect (Allee, 1931) the wave speed is determined by the flux of individuals into the region of negative growth ahead of the main front. Although numerical work (e.g., Kot et al., 1996) has
shown that such effects always slow down the front relative to the threshold-free case, analytic treatment of this intrinsically non-linear class of problems has proved elusive, the only counter-example being Lewis and Kareiva's (1993) treatment of a one-component system with cubic growth dynamics.

Despite their popularity, deterministic representations of the spatial dynamics of invasions and epidemics which assume that finite and infinitesimal populations have indistinguishable dynamics, suffer from a generic difficulty first identified by Mollison (1972, 1991). Far behind the invasion or epidemic front, most epidemic models settle to a spatially homogeneous equilibrium state in which all the players co-exist at finite abundances. In many cases, however, the passage from the epidemic front to co-existence passes through conditions where the local abundances of some or all of the players drops to truly microscopic levels. Worse, local rekindling of the disease usually takes place not because of the immigration of infectives but by in situ infections produced by the non-biological remnants of previous populations. Mollison (1991) refers to this, not inconsiderable, infelicity as “the atto-fox problem.”

A related difficulty is observed in spatial predator–prey models. Sherrat et al. (1995, 1997) have made careful observations of the behaviour of a variety of deterministic predator–prey models in the wake of invasion fronts. Where the local dynamics are strongly damped, the system passes uncontroversially to spatially homogeneous co-existence. However, where the local dynamics are weakly damped or locally unstable, they observe persistent complex wakers which are frequently chaotic. They note that such wakes invariably contain local regions in which the abundance of one or more of the players falls to microscopic levels and that recovery from these events involves in situ (and thus unbiological) regrowth rather than immigration from more highly populated regions.

Gurney and Nisbet (1998) show that preventing unbiological infections stemming from microscopic remnant populations in the SEI epidemic model has a relatively minor effect on the propagation of the initial epidemic front but causes a dramatic change in the wake following that front. The complex wake caused by unbiological re-infection is replaced by simple extinction depletion, leading to predator extinction and eventual prey recovery to carrying capacity.

Gurney et al. (1998) show that this change in post-invasion behaviour has very significant consequences for long-term population persistence. In essence, the initial invasion (or epidemic) front is followed by a region of prey (or susceptible) depletion which the predators (or infectives) cannot cross. Long-term persistence of the predator or disease thus requires either very long range dispersal of viable groups of individuals or the formation of spatially ordered structures such as spiral waves. Gurney et al. (1998) develop a series of low threshold approximations for the principal characteristics of the invasions predicted by the Rosenzweig–McArthur model and show that these are adequate predictors of the properties of one- and two-dimensional soliton and spiral-wave solutions obtained by direct simulation.

Although the characteristics which facilitate the development of these low threshold approximations are common in predator–prey systems, they are seldom found in realistic epidemic models. Although in situ regrowth can often be prevented by the ad hoc choice of a very small threshold, the biologically appropriate threshold value for many epidemic systems is large enough to cause a significant change in propagation velocity.

In this paper, we address the problem of determining those characteristics of epidemics and invasions which are robustly predicted by deterministic models with thresholds. Our first target was the velocity of propagation of the main epidemic front. Although we have been unable to improve on the linear zero-threshold result as an analytic predictor, we have been able to develop an extremely efficient numerical method with essentially arbitrary accuracy. For one-species models we can prove the correctness of this method. The extension to multi-component systems such as the SEI model is essentially heuristic, but a considerable quantity of numerical cross-checking has failed to unearth a single instance where its results are inconsistent with full simulations taking three to four orders of magnitude longer to perform. We have also been able to extend the method to provide values for the spatial extent of the initial invasion front and the peak density of predators (or infectives) contained in it.

Our paper concludes by extending the model of European fox rabies developed by Anderson et al. (1981) and Murray (1989) to include a threshold effect. We chose this system as an exemplary application of the methodology developed in this paper because its parameters are well known, and because the likely threshold is a substantial proportion of the peak incidence—leading us to expect a significant reduction in
epidemic velocity. However, because the transmission coefficient is estimated by observing that the epidemic dies out at a given (low) carrying capacity, the parameter revisions which accompany the introduction of a threshold return our velocity estimate to a value close to the zero-threshold result. Thus, the only effect of adding a realistic threshold to the fox-rabies epidemic model is to eliminate the erroneous behaviour in the wake of the primary epidemic wave—that is, the atto-fox effect.

2. ONE-COMPONENT SYSTEMS

2.1. General Treatment

In this section we consider a generalisation of the model introduced by Fisher (1937) to describe the invasive spread of a diffusively dispersing organism with logistic local dynamics. If we represent the density of the organism at time \( T \) and position \( X \) by \( N(X,T) \), then the model is

\[
\frac{\partial N}{\partial T} = Q N \left( 1 - \frac{N}{K} \right) + \Psi \frac{\partial^2 N}{\partial X^2},
\]

(1)

where \( Q \) is the intrinsic growth rate, \( K \) is the carrying capacity, and \( \Psi \) is the diffusion coefficient. We recognise the inverse of the intrinsic growth rate \( Q^{-1} \), the carrying capacity \( K \), and the diffusion length \( \sqrt{\Psi/Q} \) as the natural scales of time, population, and length, respectively. We define scaled population, time, and distance as \( n = N/K, \ t = QT, \) and \( x = X \sqrt{Q/\Psi} \), respectively. Thus, without any loss of generality, we restate the model as

\[
\frac{\partial n}{\partial t} = n(1-n) + \frac{\partial^2 n}{\partial x^2}.
\]

(2)

The properties of this model are well known (Skellam, 1951; Hadeler and Rothe, 1975; Murray, 1989). If a propagule of individuals is introduced into a previously empty system, the local population grows towards the carrying capacity and spreads into the surrounding empty territory. Eventually, the invaded and uninvaded regions are separated by a narrow “invasion front,” which moves into the uninvaded territory at constant speed. Although the front velocity formally depends on the initial condition, all initial conditions with compact support excite a front traveling at the minimum possible scaled velocity, which can be shown to be \( v = 2 \sqrt{Q/\Psi} \).

To generalise this model, we define \( g(n) \) to represent the (scaled) local population growth rate, and so write

\[
\frac{\partial n}{\partial t} = g(n) + \frac{\partial^2 n}{\partial x^2}.
\]

(3)

For the Fisher model, \( g(n) = n(1-n) \), implying positive growth for any population between zero and the carrying capacity \( (n = 1) \). We want to represent a situation where very small populations cannot grow, so we assume that \( g(n) \) has the form typified by Fig. 1a: negative for \( n \) below a threshold value \( n_T \), positive for \( n_T < n < 1 \), negative for \( n > 1 \), and zero at \( n = 0, n_T, \) and 1.

We want to study an invasion front moving from right to left as shown in Fig. 1b. We therefore transform the problem into a frame of reference moving leftwards at speed \( v_R \), by defining new variables \( z \equiv x + v_R t \) and \( \phi(z) \equiv dn/dz \). For a front travelling with constant shape, \( n \) depends only on \( z \), so we can re-express Eq. (3) as

\[
\frac{dn}{dz} = \phi, \quad \frac{d\phi}{dz} = v_R \phi - g(n).
\]

(4)

If we place the zero of \( z \) at a point where \( n = n_0 \ll n_T < 1 \) then Eq. (4) can be approximated for \( z \ll 0 \) by

\[
\frac{d^2n}{dz^2} - v_R \frac{dn}{dz} + g'(0)n = 0 \quad \text{where} \ g'(0) \equiv \left[ \frac{dg}{dn} \right]_{n=0} < 0.
\]

(5)
This has two solutions of the form \( n = n_0 e^{xz} \). Any solution with \( \lambda > 0 \) goes to zero as \( z \to -\infty \) and is thus of the form which we require. Hence we can deduce that

\[
\dot{\lambda} = \frac{1}{2}(v_R + \sqrt{v_R^2 - 4g'(0)}).
\]  

(6)

Hence, for \( z \geq 0 \) we solve Eq. (4) subject to the initial conditions

\[
n(0) = n_0, \quad \phi(0) = \dot{n}n_0.
\]

(7)

We can deduce the characteristics of the resulting solutions from a phase-plane analysis. Inspection of Eq. (4) shows that the condition for \( n \) to increase with \( z \) is just \( \phi > 0 \), while the condition for \( \phi \) to increase with \( z \) is

\[
\phi > \frac{g(n)}{v_R}.
\]

(8)

The implications of these conditions, and especially the boundary between the regions in which \( \phi \) increases and decreases, are illustrated in Fig. 1c.

We first consider solutions with a given value of \( v_R \), \( n(0) = n_0 \ll n_T < 1 \), and arbitrary (positive) values of \( \dot{n} \). For large enough values of \( v_R \), all such trajectories tend to \( n = \infty \) as \( z \to \infty \). At lower values of \( v_R \) there are three possibilities. For a single value of \( \phi_0 \), which we denote by \( \phi_0(v_R) \), the trajectory hits the steady state \((1, 0)\). For \( \phi_0 > \phi_0(v_R) \), \( n \to \infty \) as \( z \to \infty \), while for \( \phi_0 < \phi_0(v_R) \), the trajectory rises to a value above \( n = n_T \) and below 1 before eventually falling back below \( n = n_T \). In Appendix A we show that \( \phi_0 \) is a non-increasing function of \( v_R \).

We now consider solutions with the initial value of \( \phi \) linked to the frame of reference velocity \( v_R \) by setting \( \phi_0 = \dot{n}n_0 \) with \( \dot{n} \) given by Eq. (6). We shall restrict ourselves to growth functions which permit the formation of an invasion front. Thus, if we set \( v_R \) to the exact value of the front velocity \( v \), the trajectory must be that which hits \((1, 0)\), and we know that \( \lambda(v) n_0 = \phi_0(v) \). If we now consider a trajectory generated with \( v_R < v \) then we know from Eq. (6) that \( \phi_0 = \lambda(v) n_0 < \lambda(v) n_0 \). But, since \( \phi_0 \) is a non-increasing function of \( v_R \), we know that \( \phi_0 < \phi_0(v_R) \) and thus that the trajectory falls into the class which rise to a value between \( n_T \) and unity before eventually falling back below zero. An essentially parallel argument shows that if we set \( v_R > v \) the solution always tends monotonically to \( n = \infty \) as \( z \to \infty \).

As we illustrate in Fig. 1d, the behaviour implied by this argument appears highly inconvenient, because it requires us to set the frame of reference velocity exactly equal to the front velocity if we wish our solution to represent the wave-front correctly. However, viewed from a different perspective it provides us with a way of determining whether the frame of reference velocity is greater than or less than the (unique) front velocity, and thus of determining the front velocity by a simple bisection search.

### 2.2. The Fisher Equation with an Allee Effect

As an example of the class of systems discussed above, we extend the basic Fisher model to include an Allee effect (Allee, 1937). Specifically, we imagine an organism with a linearly density dependent per capita death rate, \( D \), and a per capita fecundity rate, \( B \), which is constant \((B_0)\) at high densities, but increases linearly with slope \( B_1 \) at low densities. That is, we write

\[
D = D_0 + D_1 n, \quad B = \begin{cases} B_0 & n > (B_0/B_1) \\ B_1 n & n \leq (B_0/B_1) \end{cases}
\]

(9)

If we define \( Q = B_0 - D_0 \) and \( K = Q/D_1 \) then these assumptions imply that the unscaled net growth function \( G(N) \) is

\[
G(N) = \begin{cases} QN(1 - N/K) & N > (B_0/B_1) \\ -D_0 N + (B_1 - D_1) N^2 & n \leq (B_0/B_1) \end{cases}
\]

(10)

Adopting the scaling which led to Eq. (2), and defining \( a = D_0/(B_0 - D_0) \) and \( n_T = aD_1/(B_1 - D_1) \) we find that the scaled equivalent of this expression is

\[
g(n) = \begin{cases} \frac{n(1 - n)}{a(n/n_T)(n - n_T)} & n > n_s \\ \frac{n}{1 + a/n_T} & \text{otherwise} \end{cases}
\]

(11)

where

\[
n_s = \frac{1 + a}{1 + (a/n_T)}.
\]

(12)

In this parameterisation, \( n_T \) represents the scaled threshold value and \(-a\) is the slope of the growth function at \( n = 0 \). To determine the initial condition for Eq. (4) we set \( n_0 = \min(0.001, n_T/100) \), use Eq. (6) to relate \( \lambda \) to our choice of \( v_R \), and thus determine \( n(0) \) and \( \phi(0) \) from Eq. (7). We then conduct a bisection search, starting with \( v_R = -1 \) and \( v_R = 2 \), which we know must bracket the actual velocity. The centre point of the interval is tested by solving Eq. (4) and determining if the solution passes above unity or falls back below threshold. In the first case the test velocity is above the front velocity.
and the centre point becomes the high end of the new bracketing pair. In the second, it is below and the centre point becomes the low end of the next bracket. Twenty repetitions of this procedure, which takes about 0.1 s on a medium power workstation, gives the scaled front velocity to 1 in $10^6$.

In Fig. 2a we show the relation between scaled front velocity and threshold for three values of $a$. In all cases the velocity converges slowly to the non-threshold system value, $v = 2$, as $n_T \rightarrow 0$. As we would expect, finite thresholds slow up the front. At $n_T \approx 10^{-4}$ the velocity is reduced by about 10%, while at $n_T \approx 10^{-2}$ the velocity is reduced by between 20% and 40%. As the threshold approaches unity, forward propagation stops at a critical threshold which depends strongly on $a$. For thresholds in excess of the critical value, the dynamic equations formally possess a solution of the same form but with a negative velocity. However, with any biologically realistic initial invasion, this solution corresponds to the extinction of the invading propagule.

A further possibility is suggested by Fig. 1d which shows the behaviour of this system for $a = 1.5$, $n_T = 0.3$. As $v_R$ converges towards the exact front velocity $v$, the solution of Eq. (4) follows the exact wave shape for longer and longer before diverging off to zero or $\infty$. Thus, by running the bisection process for more iterations than are needed to measure the front velocity, we can use the same calculation to give us an estimate of the front width, which we define as the distance between the points at which $n(z) = 0.05$ and 0.95. The values shown in Fig. 2b were determined by a calculation which terminated whenever the estimates of this quantity from the current upper and lower bracket velocities differed by less that 1 part in $10^3$. In practice this normally requires between 5 and 50 extra iterations, so the computational cost, although much increased over that of simple velocity determination, remains trivial.

We see from Fig. 2b that the scaled width of the invasion front changes quite markedly with its velocity. As the front slows down, it becomes steadily narrower—an effect we would expect if local growth to carrying capacity took a (scaled) time which was essentially constant.

### 3. TWO-COMPONENT SYSTEMS

We now turn our attention to systems with two players. Since these systems are not as amenable to analysis as their single species counterparts, we examine specific examples.

#### 3.1. The S–I Model

Our first example is a spatial extension of the well-known “susceptible–infective” (SI) epidemic model. We assume that susceptibles are immobile, and that in the absence of infection their population density, $S(X, T)$, would grow logistically. The conventional version of this model assumes that the per capita rate at which susceptibles become infected is proportional to the local density of infectives, $I(X, T)$, with the constant of proportionality ($B$) being known as the transmission coefficient. It then assumes that infected individuals suffer per capita mortality at rate $D_0$ and disperse diffusively with diffusion constant $\Psi$. Hence the dynamics of the system are

$$\frac{\delta S}{\delta T} = QS \left(1 - \frac{S}{K}\right) - BSI \quad \frac{\delta I}{\delta T} = (BS - D_0) I + \Psi \frac{\delta^2 I}{\delta X^2}. \quad (13)$$

The usual interpretation of deterministic models of this kind is that they describe the dynamics of average densities calculated over a (very) large ensemble of replicate systems. Practical biological experiments generally encompass only a modest degree of replication, so we shall take an alternative view in which Eq. (13) are seen as describing averages calculated over a finite-sized ensemble containing $N_B$ replicates.

At time $T$, each replicate will contain a number of infective and susceptible individuals distributed over the available space. To estimate the ensemble average infective density at $X$, $I(X, T)$, we define a quadrat of area $A_0$.
centred at \( X \) in each system, count the number of infective individuals contained in this set of quadrats, and divide the result by the product of the quadrat area and the number of replicates. The minimum non-zero value of \( I(x) \) occurs when quadrat \( X \) contains zero individuals in every replicate except one, where it contains a single individual—a configuration which implies that \( I = (N_A Q_0)^{-1} \).

It is thus clear that all values of \( I \) in the range \([0, (N_A Q_0)^{-1}]\) should be interpreted as representing a condition in which the infective population of quadrat \( X \) is extinct. In discrete space/time models which use continuous state variables it is common practice to recognise this by setting local population densities below an appropriate threshold value to zero. However, in a formally continuous space/time model this option is profoundly unappealing since it induces discontinuities in the spatial distribution.

We choose instead to eliminate unbiological behaviour which might result from effectively extinct local populations of infectives by preventing such local populations from producing infections. The most obvious way of doing this would be to set the transmission coefficient \( (B) \) to zero for infective populations below a threshold \( I_T \approx (N_A Q_0)^{-1} \).

We performed a number of our early experiments using this formulation. Although it turns out to be quite straightforward to generalise the bisection method for numerical velocity determination to tolerate a discontinuity in \( B \) for zero infective populations below a threshold \( I_T \approx (N_A Q_0)^{-1} \).

Here, \( I_T \) represents the infective density below which the infective population cannot increase, even when the susceptible population is at carrying capacity.

Defining scaled variables \( s \equiv S/K, \quad i \equiv I/K, \quad t \equiv QT, \quad x \equiv X/\sqrt{Q}\), and parameter groups \( b_0 \equiv B_0 K/Q, \quad d_0 \equiv D_0/Q, \quad i_T \equiv I_T/K \) allows us, without losses of generality, to re-express this model as

\[
\frac{\partial s}{\partial t} = s(1-s) - bsi, \quad \frac{\partial i}{\partial t} = (bs - d_0)i + \frac{\partial^2 i}{\partial x^2},
\]

where

\[
b = \begin{cases} \frac{b_0}{d_0} & i > b_0 i_T/d_0 \\ d_0/i_T & \text{otherwise.} \end{cases}
\]

This model has three, non-trivial, spatially homogeneous stationary states: one with no infectives and susceptibles at carrying capacity, one with high \( s^* \) and low \( i^* \), and one with low \( s^* \) and high \( i^* \). Provided \( i_T < (d_0/b_0^2)(1 - d_0/b_0) \), the high \( i^* \) steady state is identical to that of the equivalent un-thresholded model, namely

\[
s^* = \frac{d_0}{b_0}, \quad i^* = \frac{b_0 - d_0}{b_0^2}.
\]

So long as this steady state is positive \( (b_0 > d_0) \) it is locally stable. Spatially uniform perturbations return to it monotonically if

\[
b_0 - d_0 < \frac{d_0}{b_0^2}
\]

and return via damped oscillations otherwise.

The properties of the non-threshold version of this model have been investigated by Dunbar (1983, 1984), whose work is summarised in a more approachable form by Murray (1989). For all biologically sensible parameters, introduction of a propagule of infected individuals into a universe previously occupied by susceptibles eventually produces a stable-shaped epidemic front moving at a (scaled) velocity \( v_0 \) given by

\[
v_0 = 2\sqrt{b_0 - d_0}.
\]

When Inequality (18) is fulfilled, the front has at most one overshoot before settling to the steady state given by Eq. (17). Otherwise, the front trails a “wake” of damped oscillations.

When the local dynamics are non-oscillatory or the scaled threshold \( i_T \) is small, the behaviour of our modified model is unchanged except for a reduction in front velocity (Fig. 3a). However, the combination of oscillatory dynamics and a finite threshold can produce a radical change in behaviour. In Fig. 3c, the threshold is above the minimum infective density which would occur in the trough behind the initial epidemic if there were no threshold. Now, the infective population does not recover from this trough, but instead declines asymptotically to zero, allowing the susceptible population to recover to the carrying capacity. Thus, in addition to changing the velocity of the front, in this case the
threshold has changed the post-epidemic state of the system from a persistent admixture of susceptibles and infectives to one containing only susceptibles.

Guided by the results of the last section we study the behaviour of this system in a frame of reference moving at velocity $v_R$. Since we are looking for a constant-shape solution, we assume that $i$ and $s$ depend only on $z \equiv x + v_R t$, define $\phi(z) \equiv d i(z)/d z$, and hence re-express Eq. (15) as

$$\frac{d\phi}{dz} = v_R \phi - (bs - d_0) i, \quad \frac{di}{dz} = \phi, \quad (20)$$

$$\frac{ds}{dz} = v_R^{-1} [s(1-s) - bs i], \quad (21)$$

where $b$ is defined by Eq. (16). If we place the zero of $z$ at a point to the left of the front, where $i = i_0 \ll i_T$, then we can be sure that for all $z < 0, s(z) \approx 1$ and hence $i(z) = i_0 e^{\lambda z}$, where

$$\lambda = \frac{1}{2} [v_R + \sqrt{v_R^2 + 4d_0}]. \quad (22)$$

Hence, for $z > 0$ we must solve Eq. (20) and (21) subject to the initial conditions $s(0) = 1$, $i(0) = i_0$, and $\phi(0) = 2i_0$.

The solutions thus produced, which we illustrate in Figs. 3b and 3d, are strongly reminiscent of those we obtained for the Fisher equation in the last section. In particular, they follow the true shape of the wavefront for a distance which depends on how close the frame of reference moving at velocity $-v_R$ is to a single value, which we hypothesize must be the front velocity in a static frame of reference. If the frame of reference velocity is less than the front velocity then the density of infectives eventually becomes negative, while if $v_R > v$ the infective density diverges to $+\infty$. Although in this case we cannot prove this behaviour to be generic, we have performed a wide-ranging numerical investigation without uncovering a single counterexample.

This regularity of behaviour again presents us with an opportunity to make an efficient numerical determination of the velocity of the epidemic front using a bisection search. In Fig. 4a we show curves of front velocity against threshold infective density calculated in this way. Since we cannot prove that this procedure converges to the correct front velocity, we have also performed direct simulations of the model defined by Eqs. (15) and (16) and measured the asymptotic properties of the epidemic front. We show that the velocities thus observed are in excellent agreement with those predicted by the bisection method. We also notice that as the threshold goes to zero, the bisection method velocity predictions converge

FIG. 3. Epidemic waves in the S-I model with $b_0 = 12, d_0 = 4$. Frames (a) and (c) show snapshots of the wavefront at $t = 25$ for $i_T = 10^{-5}$ and $5 \times 10^{-3}$, respectively. Frames (b) and (d) show equivalent solutions in frames of reference moving at different (marked) velocities. To obtain (b) and (d) we solved Eqs. (20) and (21) with $s(0) = 1$, $i_T = i_0$, using RK4 with an allowed error of $1 \times 10^{-6}$. To obtain (a) and (c) p.d.e.’s (15) were discretised by the method of lines onto a grid with $\Delta x = 0.25$ and the resulting system of o.d.e.’s integrated using RK4 with an allowed error of $1 \times 10^{-4}$.

FIG. 4. Properties of epidemic fronts in the S-I model with $d_0 = 4$ and $b_0$ as marked, plotted against normalised threshold $i_T$. Smooth curves show results from Eqs. (20) and (21) using the bisection method. Points show results from numerical solution of Eq. (15); $b_0 = 4.5$ (circles), $b_0 = 5$ (squares), $b_0 = 12$ (diamonds), $b_0 = 40$ (triangles). (a) Front velocity, shown as $v_F$ where the zero threshold value $v_0 = 2 \sqrt{b_0 - d_0}$. (b) $i_{\text{max}} = \text{maximum abundance of infectives}$. (c) $w_F = \text{distance between the point where } i \text{ first exceeds } 0.95 i_{\text{max}}$ and the point where it first exceeds $0.05 i_{\text{max}}$. 

Epidemics and Invasions with Thresholds

285
Our second two-component example is a spatial extension of the Rosenzweig–McArthur predator–prey model (Rosenzweig, 1971). Here we assume that the prey \( F(X, T) \) is immobile and grows logistically with intrinsic growth rate \( R \) to a carrying capacity \( K \) in the absence of predation. The predators \( (C(X, T)) \) disperse diffusively with diffusion constant \( \Psi \) and suffer density-independent mortality at per capita rate \( D_0 \). They kill prey at a rate \( U \) given by a Holling type II functional response with maximum uptake rate \( U_m \) and half-saturation prey abundance \( H \),

\[
U(F) = \frac{U_m F}{F + H}. \tag{23}
\]

Ingested prey are turned into new predator individuals with efficiency \( E \). Hence, the dynamics of the system are described by

\[
\begin{align*}
\frac{\partial F}{\partial T} &= QF \left(1 - \frac{F}{K}\right) - UC \\
\frac{\partial C}{\partial T} &= (EU - D_0) C + \Psi \frac{\partial^2 C}{\partial X^2}.
\end{align*} \tag{24}
\]

Provided that predators never become locally scarce this model can safely be assumed to have a constant value of prey–predator conversion efficiency, \( E_0 \). However, if reproduction is sexual, then predators present in very small abundances will be unable to mate and hence turn ingested prey into new predators. To model this effect, we write

\[
E = \begin{cases} 
E_0 & C \geq C_T E_0 U_m / D_0 \\
D_0 C(U_m C_T) & \text{otherwise.}
\end{cases} \tag{25}
\]

We identify \( Q^{-1}, \sqrt{\Psi/Q}, \) and \( E_0 H \) as the natural scales of time, distance, food, and consumer population, respectively, and re-express the model in terms of dimensionless variables \( t = QT,\; x = X / \sqrt{\Psi} Q,\; f = F / H,\; c = C / (E_0 H) \), and parameter groups \( k = K / H,\; u_m = U_m / Q,\; c_T = C_T / (E_0 H) \); thus

\[
\begin{align*}
\frac{df}{dt} &= f \left(1 - \frac{f}{k}\right) - uc \\
\frac{dc}{dt} &= (eu - d_0) c + \frac{\partial^2 c}{\partial x^2}.
\end{align*} \tag{26}
\]

where

\[
u = \frac{u_m f}{1 + f} \tag{27}
\]

and

\[
e = \begin{cases} 
1 & c \geq c_T u_m / d_0 \\
d_0 c / u_m c_T & \text{otherwise.}
\end{cases} \tag{28}
\]

This model has many similarities to the S–I epidemic model discussed above. It has spatially homogeneous exterior steady states at \((c^*, f^*) = (0, 0)\) and \((0, k)\). Provided

\[
c_T < \frac{d_0}{u_m - d_0} \left(1 - \frac{d_0}{k(u_m - d_0)}\right), \tag{29}
\]

it has a spatially homogeneous interior steady state

\[
f^* = \frac{d_0}{u_m - d_0},\;\; c^* = (1 + f^*) \left(1 - \frac{f^*}{k}\right), \tag{30}
\]

which is identical to that of the equivalent non-thresholded model. However, in contrast to the S–I model case, the interior steady state is locally unstable against spatially uniform perturbations if

\[
k > \frac{u_m + d_0}{u_m - d_0}. \tag{31}
\]

Invasions in this system behave in a very similar way to the epidemics shown in Fig 3. When the local dynamics are stable and either strongly damped or non-oscillatory, the invasion front trails a short wake, leading
eventually to spatially uniform co-existence (cf. Fig. 3a).
When the local dynamics show oscillatory instability or weakly damped oscillations, the solution takes the form of a soliton (cf. Fig. 3c).
As before, we study the behaviour of such solutions by transforming the system into a frame of reference moving at velocity $-v_R$. Since we are looking for a constant-shape solution, we assume that $c$ and $f$ depend only on $z \equiv x + v_R t$, define $\phi(z) \equiv dc(z)/dz$, and re-express Eq. (26) as

$$\frac{df}{dz} = v_R \phi - (eu - d_0) c, \quad \frac{dc}{dz} = \phi,$$

(32)

$$\frac{df}{dz} = v_R [f(1-f/k) - uc],$$

(33)

where $u$ is defined by Eq. (27) and $e$ by Eq. (28). If we place the zero of $z$ at a point to the left of the front, where $c = c_0 \ll c_T$, then we can be sure that for all $z < 0$, $f(z) \approx k$ and hence $c(z) = c_0 e^{\lambda z}$ where

$$\lambda = \frac{1}{2} [v_R + \sqrt{v_R^2 + 4d_0}].$$

(34)

Hence, for $z > 0$ we must solve Eq. (32) and (33) subject to the initial conditions $f(0) = k$, $c(0) = c_0$, and $\phi(0) = \lambda c_0$.

The behaviour of this transformed system is essentially identical to that of the S-I system illustrated in Figs. 3b and 3d. If the frame of reference velocity $v_R$ is greater than the front velocity ($v$) then the normalized consumer population ($c(z)$) tends to $\infty$ as $z$ becomes large. If $v_R < v$ then the normalized consumer population eventually becomes negative. As the frame of reference velocity approaches the true front velocity the solution of the transformed system follows the true solution to progressively larger values of $z$. Although we cannot prove that this behaviour is generic, we have conducted a wide-ranging numerical investigation without finding any counter-examples.

As in the case of the S-I model, this behaviour provides us with an efficient numerical route to determining the velocity of the front by a bisection search. Extending the bisection process to sufficiently high accuracy allows us to determine the characteristics of the early part of the front—namely the peak consumer density and the spatial width of the initial invasion ahead of the peak consumer density.

In Fig. 5 we plot the results of four such investigations, together with a small number of results measured (at many orders of magnitude more labour) from direct solution of the scaled dynamic equations (26). As for the S-I model we see excellent agreement between our bisection method estimates and the confirmatory simulations. In further confirmation, we note that as $c_0 \rightarrow 0$ the velocity of the invasion front tends (albeit slowly) to the analytic value derived for the threshold-free system.

**4. THE S-I MODEL AND EUROPEAN FOX RABBIES**

Our final example is a spatial variant of the well-known S–E–I epidemic model developed to represent a disease, such as rabies, which has a long incubation period. We consider a one-dimensional universe containing individuals who can be susceptible, exposed (infected) but not infectious, or infectious (rabid). The local susceptible population grows logistically to carrying capacity $K$. Susceptible individuals are infected by contact with rabid individuals, whereupon they enter the exposed state, from which they have a constant probability per unit time of progressing to the infective stage. This model expresses the dynamics of the normalised populations of susceptibles, exposed and infectives ($S(X, T)$, $E(X, T)$, and $I(X, T)$) as
TABLE I

Parameters for the SIR Fox–Rabies Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carrying capacity</td>
<td>$K$</td>
<td>0.2 $\to$ 4</td>
<td>km$^{-2}$</td>
</tr>
<tr>
<td>Intrinsic growth rate</td>
<td>$Q$</td>
<td>0.5</td>
<td>yr$^{-1}$</td>
</tr>
<tr>
<td>Exposed mortality</td>
<td>$D_e$</td>
<td>0.5</td>
<td>yr$^{-1}$</td>
</tr>
<tr>
<td>Infective (rabid) mortality</td>
<td>$D_i$</td>
<td>73</td>
<td>yr$^{-1}$</td>
</tr>
<tr>
<td>Transmission coefficient</td>
<td>$B$</td>
<td>80</td>
<td>km$^2$/yr</td>
</tr>
<tr>
<td>Transfer rate</td>
<td>$\theta$</td>
<td>13</td>
<td>yr$^{-1}$</td>
</tr>
<tr>
<td>Diffusion coefficient</td>
<td>$\Psi$</td>
<td>200</td>
<td>km$^2$/yr</td>
</tr>
</tbody>
</table>

$B = \begin{cases} 
B_0 & \text{if } I \geq KB_0I_T \theta/(D_i(\theta + D_e)) \\
[\{\theta + D_e\}D_i/(\theta K)](I/I_T) & \text{otherwise,} 
\end{cases}$

so that the parameter $I_T$ represents the minimum local density of rabid individuals which can grow in the absence of immigration, provided the susceptibles are at carrying capacity.

The relationship between the behaviour of the model defined by Eqs. (35), (36), (37) and its non-thresholded cousin is very similar to that described by Gurney and Nisbet (1998) and discussed earlier in this paper in the context of the SI model. However, this version of the thresholded SEI model turns out to be susceptible to the treatment described earlier for the SI epidemic model and the Rosenzweig–McArthur predator–prey model. If we transform the system into a frame of reference moving with an arbitrary velocity $V_R$, then the properties of the resulting solutions depend critically on the relation between $V_R$ and the epidemic front velocity $V$. If $V_R > V$ then the solution of the transformed system has $I \to \infty$ as $z$ becomes large, whereas $V < V_R$ implies that $I$ eventually passes below zero. Since the interior steady state is always unstable in the transformed system, all solutions eventually go to $\infty$ or become negative except in the special case $V_R = V$. As $V_R$ approaches $V$ the solution in the transformed system follows the correct wave shape for an increasing distance in the transformed co-ordinate.

This behaviour implies that the numerical technique explored earlier for finding the properties of epidemic fronts in simpler systems, can also be expected to work here. In Fig. 6 we show the relationship between threshold rabid density ($I_T$) and front velocity, height, and width for a series of values of carrying capacity spanning the plausible range for the European fox. On the same diagram we show a number of values of the same quantities calculated from direct simulations of Eqs. (35), (36), and (37). Since we see excellent agreement between bisection method results and direct simulations at a widely distributed set of points, we can have confidence that the broad picture shown in this diagram is accurate.

Gurney and Nisbet (1998) showed that with the parameters given in Table I, a threshold as low as $10^{-6}$ km$^{-2}$ gives rise to a soliton solution of the type shown in Fig. 3c. Their simulation results suggest that such a low threshold value should affect the epidemic velocity by only a few percent, and this is confirmed by the more accurate calculation leading to Fig. 6. However, this figure also shows that higher values of $I_T$...
FIG. 6. Properties of epidemic fronts as a function of threshold ($I_T$) in the thresholded S-I model with parameters from Table I. Curves show values derived from the bisection search method for a given value of $K$. Points show direct simulation results. (a) Front velocity; (b) peak infective (rabid) density; (c) front width defined as in Fig. 5.

can produce very significant changes in epidemic properties.

To make an estimate of an appropriate threshold in this case, we note that the average lifetime of an individual showing clinical symptoms (and hence acting as a source of infection) is 5 days (0.014 yr). It seems highly unlikely that any infected individual can live for more than twice this period (0.028 yr). Murray (1989) argues that although the diffusion coefficient for rabid foxes is hard to estimate accurately, its value must lie in the range from 70 to 330 km$^2$ per year. The root mean square displacement of a rabid individual between becoming infective and dying must therefore be between 2.8 and 6.1 km. Since the probability distribution of this displacement is Gaussian, it seems reasonable to assume that any susceptible which does not have at least one rabid individual within three times the root mean square displacement is safe from infection. This implies that $I_T$ lies between $1 \times 10^{-3}$ km$^{-2}$ and $4.5 \times 10^{-3}$ km$^{-2}$.

In terms of our earlier discussion of the interpretation of thresholds, we note that since the peak infective density is on the order of 0.3 km$^{-2}$ (Fig. 6) we should think of the densities in our rabies model as being defined on quadrats with $A_Q \approx 3$ km$^2$. This implies that with thresholds in the range given above, we should interpret our model as describing the dynamics of averages taken over an ensemble containing between 75 and 330 members.

It is clear from Fig. 6 that a threshold of the order of $2 \times 10^{-3}$ km$^{-2}$ will decrease the velocity and increase the width of the epidemic front by around 30% compared to the values predicted by a model with zero threshold and otherwise identical parameters. However, we recall that the value given in Table I for the transmission coefficient ($B_0$) was chosen to match the observation that rabies is not self-sustaining when the fox carrying capacity is below a critical value, $K^c$. Figure 6 implies that at a given value of $B_0$, the minimum value of $K$ for a self-sustaining epidemic ($K^c$) depends on $I_T$. Thus, once a value for $I_T$ has been selected, $B_0$ must be re-adjusted to return $K$ to its observed value, $K^c$.

Observed values of $K^c$ cover a range from 0.25 to 1.0 km$^{-2}$ (Lloyd 1976, Andral and Toma 1977). We arbitrarily choose the middle of this range and set $K^c = 0.63$ km$^{-2}$. Since the ratio of the smallest to the largest possible value of diffusion coefficient is nearly 5, we choose as our best estimate of this quantity a value, $\Psi = 150$ km$^2$ yr$^{-1}$, which differs from both ends of the range by roughly the same factor (±22). In Figs. 7a and 7b we show the best estimate of front velocity and peak rabid abundance calculated using these parameters. Although introducing a threshold with all other parameters held constant always reduces front velocity, the adjustment to $B_0$ required to return $K$ to its observed value $K^c$ implies that the results differ little from those derived by Murray (1989) from the zero-threshold version of this model.

FIG. 7. Predicted characteristics of rabies epidemics in the European fox population. Solid curves show best estimates, dotted curves show values calculated from extremal values of diffusion coefficient. Upper frames show velocity (a) and peak incidence (b) with constant diffusion coefficient. $\Psi = 150$ km$^2$ yr$^{-1}$, $I_T = 2.1 \times 10^{-3}$ km$^{-2}$, $B_0 = 221$ km$^2$ yr$^{-1}$. Lower frames show velocity (c) and peak incidence (d) with $\Psi \propto 1/K$. $\Psi = 300/K$ km$^2$ yr$^{-1}$, $I_T = 1.05K \times 10^{-3}$ km$^{-2}$, $B_0 = 164$ km$^2$ yr$^{-1}$.
A characteristic feature of the predictions of both thresholded and non-thresholded SEI models is that the predicted front velocity is a strongly increasing function of carrying capacity. This conflicts with field observations suggesting that the epidemic velocity is always in the range 30–60 km yr\(^{-1}\). In their (linear) treatment, van den Bosch et al. (1990) overcome this problem by postulating that the size of fox territories, and hence the diffusion coefficient, is inversely proportional to fox density. To investigate the effect of this assumption in our model, we assume that territory size cannot adjust rapidly to changes in instantaneous density, so that the diffusion constant may be regarded as inversely proportional to carrying capacity \(K\). As a direct corollary of this assumption we also take \(I_0\) to be proportional to \(K\). We assume, rather arbitrarily, that our previous evaluation of diffusion constant produced the appropriate result for a fox density of 2 km\(^{-2}\), so we set \(\Psi = 300/K\) km\(^{-2}\) yr\(^{-1}\) and \(I_0 = (1.05 \times 10^{-3}) K\) km\(^{-2}\).

In Figs. 7c and 7d we show the best estimates of front velocity and peak rabid abundance calculated using this prescription and adjusting \(B_0\) so that \(K_c = K_e = 0.63\) km\(^{-2}\). Again, the absolute size of the predicted velocity is surprisingly close to the equivalent zero-threshold results (van den Bosch et al. 1990). As we might expect by analogy with that calculation, the key feature of the results from this variant of the model is that over most of the plausible range of carrying capacity, the epidemic velocity is almost independent of \(K\).

The most striking conclusion from both calculations shown in Fig. 7 is that the difference between the velocity predicted by our modified S–E–I model and that predicted by its less biological cousin is dwarfed by the uncertainty in the absolute size of the diffusion coefficient and the dependence (if any) of that diffusion coefficient on carrying capacity. This suggests that those features of the model predictions which are robust against the elimination of unbiological infectivity—mainly the existence and properties of the initial epidemic wave—can safely be modelled by deterministic reaction–diffusion methods with or without thresholds. Those features which are critically dependent on the occurrence of unbiological infection—specifically the passage to a stable endemic—can safely be eliminated from the repertoire of a reaction–diffusion formulation by the addition of a threshold or Allee effect.

5. DISCUSSION

In this paper, we describe an efficient numerical method for determining the characteristics of an epidemic or invasion when the invading organism cannot grow from arbitrarily low densities. This method depends on observing that when we transform the reaction–diffusion description of the system into a moving frame of reference, its properties change dramatically. All the interior steady states become violently unstable and generating a solution which ends on any of them poses a problem akin to firing a ball bearing up a smooth mountainside so that it finishes up balanced on the summit.

The key to exploiting this situation turns out to be the fact that although hitting the steady state requires exact determination of the correct frame of reference velocity and is thus functionally impossible, determining whether the ball passes over the ridge or falls back is simple. Moreover, the distinction between overshooting and falling back turns out to correlate exactly with the relation between the frame of reference velocity and the front velocity—providing a natural route to determining the front velocity to an arbitrary degree of accuracy by a bisection search.

Although we have only proved the correctness of this methodology for the class of single component models formed by generalizing the Fisher model, we have observed experimentally that it works excellently for a wide class of multi-component models including the S–I and S–E–I epidemic models and the Rosenzweig–McArthur predator–prey model, with a variety of alternative representations of low density behaviour.

Although the solution of the transformed system always diverges from the “correct” solution eventually, we observe experimentally that as the frame of reference and front velocities converge, the transformed solution follows the shape of the untransformed solution for progressively longer and longer. This implies that if the bisection process designed to yield velocity estimates is continued to rather higher accuracy than would normally be warranted by epidemic velocity data, the transformed solution follows the untransformed solution far enough for us to be able to reliably determine the characteristics of the leading part of the epidemic wave—specifically the spatial extent of the epidemic onset and the peak incidence of the disease.

The work reported here has shown that the characteristics predicted for the initial epidemic or invasion wave by reaction–diffusion formalisms are qualitatively robust against the elimination of the atto-fox effect (Mollison, 1991). The effect of introducing a preventative threshold or Allee effect is generally to slow up the epidemic front and reduce the peak disease incidence. However, as our application to European fox rabies demonstrated, the particular route by which parameters are estimated can reduce these changes to the point
where they are insignificant compared to the inaccuracies inherent in parameter or structural uncertainty.

Because of the inherent difficulty of estimating diffusion coefficients, explicit inclusion of thresholds or Allee effects seems unlikely to change estimated velocities by observable amounts. However, models incorporating these effects yield robust estimates of the characteristics of the initial epidemic, and are very amenable to efficient numerical estimation of these quantities. In common with their non-thresholded cousins, these models currently have nothing useful to say about the long term persistence of the disease—a problem which clearly requires new observational as well as theoretical insights. However, there are indications that they form a more promising basis for theoretical developments in this area than models which permit unbiological reinfection.

APPENDIX

We seek solutions of

\[ \frac{dn}{dz} = \phi, \quad n(0) = n_0, \quad (38) \]

\[ \frac{d\phi}{dz} = v_R \phi - g(n), \quad \phi(0) = \phi_0. \quad (39) \]

This is equivalent to

\[ \frac{d\phi}{dn} = \gamma(n, v_R, \phi), \quad \phi(n_0) = \phi_0, \quad (40) \]

where

\[ \gamma(n, v_R, \phi) = v_R - \frac{g(n)}{\phi}. \quad (41) \]

We represent the solution of this system by \( \phi(n, v_R, \phi_0) \) and define \( \phi_1(v) \) such that \( \phi(n, v, \phi_1(v)) \rightarrow 0 \) as \( n \rightarrow 1^- \).

We first note that Eq. (41) implies that

\[ v_1 > v_2 \Rightarrow \gamma(n, v_1, \phi) > \gamma(n, v_2, \phi), \quad (42) \]

which in turn implies that, so long as \( \phi(n, v_1, \phi_0) > 0, \)

\[ v_1 > v_2 \Rightarrow \phi(n, v_1, \phi_0) > \phi(n, v_2, \phi_0). \quad (43) \]

Now consider two trajectories, the first calculated with \( \phi(n_0) = \phi_1(v_1) \) and \( v_R = v_1 \), and the second calculated from the same initial condition but with \( v_R = v_2 < v_1 \). The first trajectory hits \((1, 0)\). Inequality (43) tells us that while the first trajectory is above the \( n \)-axis, the second trajectory cannot be above it and will generally be below it. The second trajectory thus crosses the \( n \) axis \((\phi = 0)\) at or to the left of \((1, 0)\). If it hits \((1, 0)\) then we know that \( \phi(v_1) = \phi(v_2) \). If it crosses to the left of \((1, 0)\) then the phase plane analysis shown in Fig. 1c implies that to cause it to hit \((1, 0)\) we must use an increased value of \( \phi(n_0) \), so \( \phi_1(v_1) > \phi_1(v_2) \). Hence, we know that

\[ v_1 > v_2 \Rightarrow \phi_1(v_1) > \phi_1(v_2). \quad (44) \]

ACKNOWLEDGMENTS

The authors acknowledge grant support from the Natural Environment Research Council (Grant GST 02/0998). IC gratefully acknowledges the receipt of a research studentship from EPSRC. WSCG is grateful for the support and hospitality of the University of Calgary during the initial phase of the work.

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