Modelling *Aedes aegypti* mosquito control via transgenic and sterile insect techniques: Endemics and emerging outbreaks

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**AUTHOR HIGHLIGHTS**

- Spatial model considering control strategies for two variants of sterile male releases (SIT and RIDL).
- Discussing the minimal release resources and optimal strategy costs for complete eradication of mosquitoes.
- The control effectiveness and optimal control strategy vary depending on scenarios and associated strategies.

**ABSTRACT**

The invasion of pest insects often changes or destroys a native ecosystem, and can result in food shortages and disease endemics. Issues such as the environmental effects of chemical control methods, the economic burden of maintaining control strategies and the risk of pest resistance still remain, and mosquito-borne diseases such as malaria and dengue fever prevail in many countries, infecting over 100 million worldwide in 2010. One environmentally friendly method for mosquito control is the Sterile Insect Technique (SIT). This species-specific method of insect control relies on the mass rearing, sterilization and release of large numbers of sterile insects. An alternative transgenic method is the Release of Insects carrying a Dominant Lethal (RIDL). Our objective is to consider contrasting control strategies for two invasive scenarios via SIT and RIDL: an endemic case and an emerging outbreak. We investigate how the release rate and size of release region influence both the potential for control success and the resources needed to achieve it, under a range of conditions and control strategies, and we discuss advantageous strategies with respect to reducing the release resources and strategy costs (in terms of control mosquito numbers) required to achieve complete eradication of wild-type mosquitoes.

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**1. Introduction**

The history of pest control is as old as human agriculture or disease. The invasion of pest insects often changes or destroys a native ecosystem, and can result in food shortages and disease endemics. As a result, the development of biological control methods has received widespread attention and, in some cases, they have been successful (Benedict and Robinson, 2003; Dyck et al., 2005; Vreysen et al., 2007). However, issues such as the environmental effects of chemical control methods, the economic burden of maintaining control strategies and the risk of pest resistance still remain, and mosquito-borne diseases such as Malaria and Dengue fever prevail in many countries in East Asia, South America and Africa, infecting over 100 million and killing at least half a million in 2010 (WHO, 2012a, 2012b, 2012c). Furthermore, repeated invasions are observed in regions where the vector mosquito species have been eradicated completely in the past. For example, *Aedes aegypti* and *Aedes albopictus* are observed in Northern European countries as well as Asia (Hulden and Hulden, 2008; Paupy et al., 2012). Global warming and the human transportation system also promote such situations (Ensink, 2010). As such, continued research into the development of better pest control methods remains vital (Dyck et al., 2005; Pimentel, 2011).

One environmentally friendly alternative for mosquito control is the sterile insect technique, SIT (Knipling, 1955). This species-specific method of insect control relies on the mass rearing, sterilization and release of large numbers of sterile insects, preferably males (Dyck et al., 2005), which, it is hoped, mate with wild-type insects, thereby reducing their reproductive output and,
potentially, the pest population abundance (see Black et al., 2011; Wilke et al., 2012 for recent reviews). Mixed-sex sterile releases are avoided where practical as they are generally less efficient and, for species such as mosquitoes, it is only the females that bite. This means that their release could potentially aid disease spread in the short-term (see Alphey et al., 2010 for a recent review).

Other transgenic technologies have recently been developed to improve SIT control (Benedict and Robinson, 2003; Wimmer, 2003; Alphey et al., 2010); these include genetic sexing (Robinson et al., 1999), genetic marking (Pelouquin et al., 2000) and genetic female-specific lethality (Seawright et al., 1978). One such transgenic strategy is RIDL, i.e. "Release of Insects carrying a Dominant Lethal" (Thomas et al., 2000; Phuc et al., 2007). Here the released transgenic males are homozygous for a dominant lethal gene that is expressed in both male and female (bisex) progeny that result from mating with wild-type insects. Female-specific RIDL strategies have also been developed (Fu et al., 2010), but here we focus on bisex RIDL control strategies. Hereafter, we use the terms SIT and sterile to refer to early-acting lethality of the progeny of released insects, for example classical SIT using radiation-induced sterility, and the terms RIDL and transgenic to refer to late-acting lethality in both sexes.

We note also that the developmental stage at which the dominant lethal gene is expressed, for instance the embryonic or the larval stages, can have a substantial effect on the control strategy. In particular, late acting genes, which induce death after the density-dependent larval stage, have a significant advantage over SIT strategies because of an additional reduction in pest abundance that arises as a result of larval competition (Atkinson et al., 2000; Phuc et al., 2007; White et al., 2010).

The details of mosquito dispersal behaviour are not completely understood (Reiter et al., 1995; Harrington et al., 2005), though there have been mathematical modelling studies highlighting that Ae. aegypti invasion rates have a critical influence on the success of the control strategy (Lewis and Driessche, 1993; Takahashi et al., 2004; Yakob et al., 2008; Magori et al., 2009; Seirin-Lee et al., 2013). Nonetheless, studies that explore the effects of Ae. aegypti invasive dynamics upon the efficacy of SIT and RIDL control strategies in eliminating mosquitoes are limited to those by Yakob et al. (2008) and Yakob and Bonsall (2009), which consider the interplay of stage structuring and dispersion on a lattice with a small control region that is embedded within an established pest population. These investigations reveal complex dynamics and focus on the differences between SIT and RIDL control strategies for a very limited variation in spatial parameters, other than dispersal rates. However, firstly, it is not clear whether a strategy aimed at eliminating an established pest is appropriate for eradicating an emergent, invading, outbreak. In addition, the influence of systematically varying the size of the region in which control insects are released is an aspect of spatially heterogeneous models that is essentially unexplored and merits detailed study, given the concern that spatial dynamics such as mosquito invasion is becoming a critical issue on global scale (Benedict et al., 2007; Jansen and Beebe, 2010).

Furthermore, such detailed investigations are facilitated in the continuum modelling approach considered here, which allows the ready prediction of scaling laws, as illustrated below for the influence of dispersal rates. More generally the continuum approach is typically an appropriate and efficient framework, and thus often advantageous, when the lengthscale and timescale under consideration are large compared to those describing the population’s individuals.

Our objective is thus to consider control strategies for two control scenarios via SIT and RIDL: an endemic case and an emerging outbreak for a mosquito vector. In the former case, a mosquito vector is endemic. In contrast, in the latter case invading mosquitoes establish and cause a local outbreak in a previously mosquito-free region; see Fig. 1. An important question is how such differences in the initial scenario induce different responses to variations in control strategies with SIT and RIDL. In particular, we are concerned with how these responses are influenced by spatial parameters such as dispersal rates and especially the lengthscales of the regions in which control insects are released. Thus for the two contrasting scenarios, we investigate how varying the release rate in conjunction with the size of release region influences both the potential for control success and the resources needed to achieve it, in terms of control mosquito numbers, under a range of conditions. We thus discuss the relationships between the size of the control zone, the mosquito dispersal rate and advantageous strategies with respect to reducing control insect numbers and thus improving the strategy costs required to achieve eradication of mosquitoes. Finally, we briefly note that in the emerging outbreak case, we explore release efforts and strategy-costs with a control strategy that can eradicate the wild-type females. This is in distinct contrast to halting the spread of an outbreak using a barrier zone method of our previous study (Seirin-Lee et al., 2013).

2. Materials and methods

2.1. Mathematical models

We build upon the temporal model of mosquito population dynamics developed by Dye (1984), which was validated on data for the larval and adult ecology of Ae. aegypti in Wat Samphaya,
Bangkok, Thailand, published in Sheppard et al. (1969) and from unpublished reports of the World Health Organization’s Aedes Research Unit (ARU) in Bangkok [ibid].

The densities of wild-type female mosquitoes and sterile/transgenic male mosquitoes at time $t$ are respectively denoted by $N(t)$ and $S(t)$. Following Dye (1984) we firstly assume that mosquito proliferation proceeds via a stage-structured process approximated by a delayed density-dependent mortality acting on a pre-adult developmental stage, reflecting larval competition. In addition, equal numbers of male and female wild-type mosquitoes are assumed, and it is taken that wild-type females mate in proportion to their relative abundance (Knipping, 1955; Phuc et al., 2007), at a rate given by $N(t)/N(t) + cS(t))$ where $0 < c \leq 1$ represents the reduced mating competitive ability of sterile male or transgenic male mosquitoes. We also impose the same per capita death rate, denoted $\mu$, below, for the female wild-type and male sterile/transgenic mosquitoes. In addition, the control framework is modelled by the release of sterile or transgenic male mosquitoes at a constant rate, denoted $\kappa = \theta N^*$, where $N^*$ is the control-free equilibrium density of wild-type mosquitoes and $\theta$ is defined as the release rate ratio.

By balancing mosquito numbers, these assumptions yield the following equations:

\[
\frac{dN(t)}{dt} = rN(t-T)\left(\frac{N(t-T)}{N(t-T) + cS(t-T)}\right)\phi(t) - \mu N(t),
\]

\[
\frac{dS(t)}{dt} = \kappa - \mu S(t).
\]

Here $\phi(t)$ captures density-dependent competition in the larval stage, the delay time, $T$ represents the mosquito developmental time in the stage-structuring and, finally, the egg production rate per adult female is denoted by $r$ and is multiplied by a corrective factor to account for fertile matings with steriles and imperfect survival while reaching the adult stage.

The late-acting lethal induced by RIDL is anticipated to participate in larval competition and thus $\phi$ is unaffected by the perturbations induced by such control strategies and hence is independent of transgenic mosquitoes. Following the classical insect population dynamics of Gurney et al. (1980), we therefore have

\[
\phi(t) = \exp[-\alpha E N(t-T)],
\]

with RIDL control. Here $\alpha$ is a parameter representing the strength of density-dependent competition that facilitates fitting with field data, as detailed by Dye (1984). Note that $\alpha, E$ occur only in the parameter grouping $\alpha E$ and thus one cannot separate the interpretation of these two parameters. They are distinct here to maintain notational similarity with Dye’s (1984) model formulation, where $1/\alpha$ is interpreted as the size at which the wild-type female mosquito population reproduces at maximum rate and $E$ is the egg production rate of adult mosquitoes. Nonetheless, below we treat $\alpha E$ as a single parameter grouping.

For SIT, the matings with control mosquitoes do not give rise to any offspring, and thus larval competition is reduced in proportion to the number of fertile matings. Hence, for SIT, we have (Phuc et al., 2007; White et al., 2010; Seirin-Lee et al., 2013)

\[
\phi(t) = \exp\left[-\alpha E \left(\frac{N(t-T)}{N(t-T) + cS(t-T)}\right)\right].
\]

thus accounting for how the SIT interventions interfere with larval competition. The general extent to which such models concur with alternative representations of stage structure in mosquito dynamics, for instance the models based on the framework of Focks et al. (1993a, 1993b) such as Erickson et al. (2010), is an open question that we do not address here.

We proceed to generalise the temporal model (1)–(3) to consider spatial dynamics in a one-dimensional homogeneous domain (see Fig. 1 for a schematic). The larvae are not motile and hence there is no dispersive kernel linking the stages of mosquito maturation, though the adults are taken to diffuse at constant rate. Hence, for $t > 0$ we have

\[
\frac{\partial N(x, t)}{\partial t} = D \frac{\partial^2 N(x, t)}{\partial x^2} + rN(x, t-T)
\]

\[
\left(\frac{N(x, t-T)}{N(x, t-T) + cS(x, t-T)}\right)\phi(x) - \mu N(x, t),
\]

\[
\frac{\partial S(x, t)}{\partial t} = D \frac{\partial^2 S(x, t)}{\partial x^2} + \kappa(x) - \mu S(x, t),
\]

where $x \in \Omega$, the spatial domain, with $D$ denoting the diffusion rate of both wild-type females and sterile/transgenic males. The competition term, $\phi(x, t)$, is given by (2) or (3) by simply exchanging $S(t)$ and $N(t)$ for $S(x, t)$ and $N(x, t)$ respectively. We also assume that the boundary of region $\Omega$ does not permit mosquito transport and thus we have zero flux boundary conditions

\[
\frac{\partial N}{\partial x} = \frac{\partial S}{\partial x} = 0, \quad x \in \partial \Omega.
\]

To model control strategies we consider the continuous release of sterile/transgenic males within the delivery region at a constant rate per unit length, $\theta N^*$, which defines $\theta$ given $N^*$ denotes the control free equilibrium pest insect density. This is described in detail via the release function

\[
k(x) = \theta N^* \chi(x),
\]

\[
\chi(x) = \begin{cases} 1 & x \in A \\ 0 & x \in \partial A \end{cases}
\]

where $A$ is the region of $\Omega$ in which sterile/transgenic males are released at rate $\theta N^*$. In Fig. 1, $A$ becomes the interval $[x, x + \gamma s]$. We use this general functional form to explore two different scenarios and their respective control strategies.

2.2. Scenarios and control strategies

We consider two scenarios. The first is an endemic case in which female mosquitoes are widespread over an isolated region $\Omega$, so that the width of the wild-type female habitat, $\gamma_N$, is equal to $\Omega$. The control is applied by releasing sterile/transgenic males locally within the region (Fig. 1(a)). The second scenario is an emerging outbreak case, in which female mosquitoes are invading a new environment. In this case, $\Omega$ is large enough so that $\gamma_N < \Omega$ (Fig. 1(b)). For both cases, control success will mean a complete eradication of wild-type female mosquitoes rather than just an invasion arrest or a decrease in pest population density.

2.2.1. Endemic outbreaks and the local release strategy

This scenario is described in Fig. 1(a) in detail and we call it the local release strategy. We assume that the female wild-type mosquito population has already approached carrying capacity in an isolated homogeneous region. The simplest control strategy for complete eradication in this scenario is the release of a sufficiently large number of sterile/transgenic males over the whole region, $\gamma_S = \gamma_N$, where $\gamma_S$ is the width of the release region. The success of this control method can be explored in a straightforward manner via the temporal model (1), because success depends only on the release rate of sterile/transgenic males per unit time. We obtain a minimal release ratio for complete eradication, as in Phuc et al. (2007) and Seirin-Lee et al. (2013). However, it is not clear how the minimal release ratios change when release is over only a portion of the region, $\gamma_S < \gamma_N$, nor how critically this ratio depends on the mosquito dispersal rate. Hence, we explore a measure of the resource cost required for the successful eradication of female mosquitoes, namely the product of the release region size and the release ratio, which below we refer to as the release effort, $[EF]_{\text{loc}}$. This measure therefore is the total number of released sterile/transgenic males per unit time,
Table 1
The values of $(r, \beta)$ associated with parameter sets A and B have been chosen from the parameter ranges estimated by Dye (1984), as also used in other modelling investigations (Phuc et al., 2007; White et al., 2010). The parameter grouping $ae^\theta$ for parameter set B has been fixed to ensure the same control-free equilibrium of approximately six million mosquitoes per kilometre.

<table>
<thead>
<tr>
<th>Parameter/variable</th>
<th>Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N$</td>
<td>Density/number of female wild-type mosquitoes</td>
<td>500 km</td>
</tr>
<tr>
<td>$S$</td>
<td>Density/number of male sterile or transgenic mosquitoes</td>
<td>500 km</td>
</tr>
<tr>
<td>$\Omega$</td>
<td>Whole spatial region</td>
<td>$0, 500$ km</td>
</tr>
<tr>
<td>$\gamma_x$</td>
<td>Width of wild-type females habitat $^*$</td>
<td>$0, 500$ km</td>
</tr>
<tr>
<td>$\gamma_S$</td>
<td>Width of sterile/transgenic male release region</td>
<td>$0, 500$ km</td>
</tr>
<tr>
<td>$D$</td>
<td>Diffusion coefficient for mosquitoes</td>
<td>$0.01, 25$ [km$^2$/day]</td>
</tr>
<tr>
<td>$T$</td>
<td>Mosquito development time</td>
<td>18.84 days</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Coefficient of reduced mating competitive ability of sterile/transgenic male mosquitoes</td>
<td>0.95</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Death rate of wild-type adult females</td>
<td>$0.12$ days$^{-1}$</td>
</tr>
<tr>
<td>$k$</td>
<td>Release rate of control strategy males</td>
<td>$ae^\theta$ days$^{-1}$</td>
</tr>
<tr>
<td>$\theta$</td>
<td>Release rate ratio of control strategy males</td>
<td>$0, 20$</td>
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</tbody>
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<table>
<thead>
<tr>
<th>Parameter set A</th>
<th></th>
<th></th>
</tr>
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<tbody>
<tr>
<td>$r$</td>
<td>Birth rate of adults corrected for egg to adult survival</td>
<td>0.367 days$^{-1}$</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Density-dependent coefficient</td>
<td>3.02</td>
</tr>
<tr>
<td>$ae^\theta$</td>
<td>Density-dependent coefficient</td>
<td>0.01$^b$</td>
</tr>
<tr>
<td>$N^*$</td>
<td>Control-free female mosquito equilibrium</td>
<td>$6.064 \times 10^{10}$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter set B</th>
<th>Parameter set B</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>$r$</td>
<td>Birth rate of adults corrected for egg to adult survival</td>
<td>1.31 days$^{-1}$</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Density-dependent coefficient</td>
<td>1.0</td>
</tr>
<tr>
<td>$ae^\theta$</td>
<td>Density-dependent coefficient</td>
<td>$3.94 \times 10^{10}$</td>
</tr>
<tr>
<td>$N^*$</td>
<td>Control-free female mosquito equilibrium</td>
<td>$6.064 \times 10^{10}$</td>
</tr>
</tbody>
</table>

$^*$ For the model, this value is given with appropriate length units, i.e. per (km)$^2$ for $ae^\theta$ and per km for $N^*$.

and is given mathematically by

$$[EF]_{loc} = rfiN^*.$$

As mentioned in the Introduction, the indefinite release of sterile/transgenic mosquitoes imposes a heavy economic burden, and hence we estimate the time to complete eradication, in particular because many of the insects involved are likely to be influenced either socially or by climate change (Purse et al., 2005; White et al., 2010). The time required for complete control will also be a very important issue in determining improved strategies. Thus we also define the strategy-cost as the product (release effort $\times$ time to eradication). Mathematically, this is given by

$$[SC]_{loc} = [EF]_{loc} \times T_{ex}.$$

where $T_{ex}$ is the extinction time of the wild-type female mosquito population, which includes definition in terms of a tolerance (or detection threshold), characterised by $\epsilon$ below. In particular, $T_{ex}$ is the smallest time such that whenever $t > T_{ex}$ we have

$$\frac{1}{|\Omega|} \int_\Omega \int_0^{N(x, t)} \frac{dx}{N^*} < \epsilon < 1.$$

Typically in our simulations we take $\epsilon = 10^{-2}$. The strategy-cost is therefore the total number of sterile/transgenic males released up until effective eradication of the wild-type female mosquito population.

2.2.2. Emerging outbreaks and the wavefront cover strategy

In the modern era of developed human transport systems, the transmission of disease over several thousands of kilometres by vector insects is common (Shigesada and Kawasaki, 1997; Enserink, 2010). We expect, with a uniform environment, mosquitoes will disperse in a wave-like manner away from their initial site of invasion, with the population approaching its carrying capacity behind the wave. We suppose that sterile/transgenic males are released over a single region of length $\gamma_S$, as depicted in Fig. 1(b), which covers the invasive wavefront.

As a measure of cost resource, we define the release effort by

$$[EF]_{cov} = \frac{rfiN^*}{\gamma^*}.$$

where $\gamma^*$ denotes the above-threshold region which wild-type female mosquitoes have invaded when control is initiated. Noting that the invasive profile is unimodal, as depicted in Fig. 1(b), we have $\gamma^*$, which satisfies the constraint $N(x, 0)/N^* = N(x + \gamma^* x, 0)/N^* = \tau$ where $\tau$ is the threshold and thus an extremely small density (which the results are insensitive to).

However, note that $\gamma^*$ is defined differently for parameter sets A and B in the numerical simulation, as these induce invasive waves with different spatial profiles. Thus the release effort function (9) has been defined per unit length and the release effort for an emerging outbreak (9) constitutes the average number of sterile/transgenic males released per unit time and per unit length of the initial above-threshold outbreak domain. With the extinction time given by (8) the strategy-cost is

$$[SC]_{cov} = [EF]_{cov} \times T_{ex} \times \gamma_N = \frac{rfiN^*}{\gamma^*} \times T_{ex}.$$

which is the total number of sterile/transgenic males released during the control period.

2.3. Parameter values

As with many other studies (e.g. Phuc et al., 2007; Yakob et al., 2008; White et al., 2010) we use Dye's (1984) estimates for the life-history parameter values for $Ae. aegypti$, which incorporate a range of values for the intrinsic birth rate, $r$, and the density-dependent coefficient, $\beta$. As presented in Table 1, we focus on two sets of parameters which represent the extremes of $r$ and $\beta$ (White et al., 2010; Seirin-Lee et al., 2013), with the grouping $ae^\theta$ chosen so that the
equilibrium density, \( N^* \), is the same for each parameter set and of the order of one million mosquitoes per kilometre for the spatial models. The first parameter set, denoted A, has a lower intrinsic birth rate, \( r \), in combination with weaker density-dependent competition, \( \beta \), and gives rise to a stable equilibrium which is approached monotonically in the absence of control strategies. In contrast, parameter set B has substantially larger birth rate, \( r \), and higher density-dependent competition, \( \beta \), which induces overcompensating density-dependent competition, giving rise to oscillatory dynamics in an uncontrolled population for the spatially homogeneous model. This dynamics arise as a peak in the adult population results in an increase in reproduction, leading to competition and a subsequent drop in the following generation. Population recovery then follows as a result of a drop in competition.

These two parameter sets also result in very different predictions concerning the control of \( A e. \ aegypti \) mosquitoes (see, for example, Phuc et al., 2007). While SIT and RIDL control strategies give rise to similar results in decreasing the population wild-type female mosquitoes in the case of parameter set A, for parameter set B, a moderate release rate of sterile mosquitoes may undesirably increase the wild-type mosquito population due to a reduction in competition offsetting the reduced birth rate.

It should be noted that we take the density-dependence parameters from Dye (1984), following many previous studies. However, Legros et al. (2009) have called these values into question by using an alternative technique, and finding different values. The qualitative results that follow do not change for these alternative values and we detail this further in the Discussion and both parameter sets are considered given the uncertainty in their estimates. Also, in the absence of explicit empirical estimates for the diffusion rates of sterile or transgenic \( A e. \ aegypti \) mosquitoes (Reiter et al., 1995; Harrington et al., 2005), we assume that the sterile/transgenic mosquitoes have the same diffusion rate as wild-type mosquitoes, and this is varied across a broad range, from hundreds of square meters per day to several square kilometres per day.

Recent studies in radiation dose optimisation have led to marked improvements in SIT in general, with some studies showing little competitive reduction from radiation (Mastrangelo et al., 2012; Oliva et al., 2012; Sow et al., 2012). Similarly, the mating competitiveness of genetically sterile RIDL male mosquitoes has been shown to be comparable to that of their wild-type counterparts in semi-field conditions (Lee et al., 2013). Therefore we assume that, for both control strategies, the mating competition coefficient \( c \) is close to unity, reflecting a small fitness cost. An extensive investigation into this parameter can be found in White et al. (2010).

For instance the effect of variations in the parameter grouping \( aE^\beta \) can be inferred from the fact that the model equations are invariant under the mapping

\[
(aE^\beta) \rightarrow (\zeta aE^\beta), \quad N \rightarrow \zeta N, \quad N^* \rightarrow \zeta N^*; \quad S \rightarrow \zeta S.
\]

Finally, a detailed numerical scheme for the model given by Eqs. (4) and (5) is described in Appendix A.

3. Results

3.1. Endemic outbreaks and the local release strategy

We consider the local release strategy, asking two main questions: (i) To what extent does the dispersal rate affect the potential for eradicating female mosquitoes? (ii) If the local release strategy is effective, what is the minimal release region and how does it relate to the release rate ratio and dispersal rate? Our simulation results show that for some release regions and rates the local release strategy is not always successful in eradicating female wild-type mosquitoes (see Fig. 2). In particular, with parameter set B, application of a local release strategy using SITs in fact induces an increase in the total female population if the release rate is not large enough, as observed in spatially homogeneous modelling (Phuc et al., 2007). Below, we explore the relationship between duration for complete eradication, the release rate and the release region size, plus their influence in reducing resources, as measured via controlled mosquito numbers.

3.1.1. Minimal release region size for complete eradication

We denote the minimal release region size by \( r_0^{\text{min}} \) and define it as the release region size at which we are able to achieve complete eradication for a given release rate ratio, \( \theta \). In order to find the minimal release region size required for complete eradication of female wild-type mosquitoes we plot, in Fig. 3(a), the threshold values of \( (\gamma_5, \theta) \) at which female mosquitoes become extinct throughout the entire habitat. Note that we have assumed in our calculations that complete eradication is achieved when the constraint (8) is satisfied.

Regardless of the control strategy and parameter choice, when the release rate ratio is small, the size of release region required for successful eradication of female mosquitoes depends sensitively on the release rate ratio. However, for large release rate ratios the minimum size of release region becomes insensitive to changes in the release rate, as shown in Fig. 3(a), although the size of release rate at which this insensitivity arises, and the size of release region there, are dependent on the control strategy and parameters chosen.
We explore the dependence of the minimal release region size upon $\gamma_N$ for a fixed release rate ratio in Fig. 3(b). The results highlight that the minimal release region size increases with female habitat size but, surprisingly, $\gamma_N^{-s_{\text{min}}}$ (denoting $s_{\text{opt}}$) is constant (approximately 30 km for parameter set A) when $\gamma_N$ is sufficiently large; however, $\gamma_N^{-s_{\text{min}}}$ decreases and tends to zero as $\gamma_N$ is reduced to zero. This enables us to suggest an intuitive result that the local release strategy is more effective for a small habitat than a large one. For example, when the female habitat is very large, we need to release sterile/transgenic males over a very wide region to achieve eradication. In contrast, when the habitat is very close in size to $s_{\text{opt}}$ or less than it, release in a very small region compared to $s_{\text{opt}}$ will be sufficient to eradicate the female population over the whole habitat. Furthermore, we note that this result is not highly sensitive to the choice of parameter set or control strategy.

The sensitivity of $s_{\text{opt}}$ to the diffusion rate is shown in Fig. 3(c) where we see the, again, intuitive result that $s_{\text{opt}}$ decreases as the diffusion rate increases. Further, as detailed in Appendix B, a scaling relation exists for the variation of the minimal release region size with the diffusion rate

$$s_{\text{min}}(D) = \frac{C}{\sqrt{D}} s_{\text{opt}},$$

where $C$ is a constant given by $C = \sqrt{D_0/D}$ with $D_0$ and $D$ denoting a fixed diffusion rate and habitat size of wild-type females, respectively. Since the choice of optimal release region is highly sensitive to the value of the diffusion rate, one would require careful experimental measurement of mosquito diffusion rates in order to be able to minimise the release effort. Nonetheless, the local release strategy is potentially applicable to small endemic regions, regardless of the parameter values and control method used.

### 3.1.2. Release effort and strategy-cost

In Fig. 3(d), we plot the release effort for each strategy and parameter set on restriction to the threshold curve, Fig. 3(a). Note that the minimal release effort is given at the minimal release rate ratio and the release effort increases monotonically as the release rate ratio increases, regardless of the choice of SIT and RIDL, or parameter set.

Further, the extinction time of the wild-type female mosquitoes at points $(\theta, s_{\text{opt}})$ taken from Fig. 3(a) is fairly constant except for small release rate ratios or sufficiently large release rate ratios, as shown in Fig. 4(a), (b). The reason that the extinction time is almost constant for intermediate release rate ratios is that it is governed by the invasion timescale of the control mosquito for the domain, given $s_{\text{opt}}$ is approximately constant. Once the minimal release region size, $s_{\text{opt}}$, becomes insensitive to increases in the release rate ratio (for example, around $\theta = 8.0$ in Fig. 4(a)), the extinction time decreases as the release rate ratio increases. This is because the extinction time for wild-type females on $\Omega s_{\text{opt}}^{-1}$ decreases as the number of sterile/transgenic males migrating into $\Omega s_{\text{opt}}^{-1}$ increases, which is promoted when the sterile/transgenic males are released as quickly as possible. In contrast, with very small numbers of released males, a long time is required for the sterile males to reach each boundary of the female habitat, so that the eradication time of the females increases.
For smaller release rates, we obtain a monotonically increasing strategy-cost, $[SC]_{loc}$, as a function of the release rate ratio, as illustrated in Fig. 4(c). This is because the value of the release effort at small release rates is small enough to counteract the influence of any increase in extinction time in the strategy-cost, Eq. (7), so that the strategy-cost monotonically increases as a function of the release rate ratio, as shown in Fig. 4(c), (d). However, the strategy cost, $[SC]_{loc}$, slightly decreases around $\theta = 8.0$ because the extinction time decreases with a large release rate ratio, as shown in Fig. 4(a).

Although the strategy cost, $[SC]_{loc}$, decreases with reductions in the release rate ratio, $\theta$, the eradication time is, in fact, sensitive to the fact we are working with the minimum release range, $\chi^{min}$ of Eq. 3(a). The white points (○, □) in Fig. 4(d) illustrate this: the strategy cost, $[SC]_{loc}$, of the white points has been calculated with the extinction time for the same release range but a very slightly elevated release rate ratio compared to the black points (●, ●). Obviously, the strategy-costs for the white points are smaller than the respective black points, a result of the decrease in $T_{ex}$. We thus can reduce strategy-costs by selecting higher release rate ratios than the threshold value, at which elimination just occurs.

### 3.1.3. Sensitivity of the surviving population of wild-type females to the diffusion rate and release rate ratio

In the endemic scenario, $\gamma_N$ is always greater than $\gamma_S$ so that if one does not choose the release region size greater than $\chi^{min}$, complete eradication of wild-type female mosquitoes will not be achieved. However, one can still achieve local eradication, as shown in Fig. 2. In what follows, we explore how mosquito dispersal rates and release rates affect the decrease in the wild-type female population. The numerical results are shown in Fig. 5 where the average number of surviving wild-type female mosquitoes is plotted as a function of both the mosquito dispersal rate and release rate ratio. The former has a negligible effect when the release rate is small. In contrast, for a large enough release rate and parameter set A, we see that an increase in the dispersal rate causes a decrease in the new population equilibrium. This is because the dispersion of sterile/transgenic mosquitoes to outlying regions increases, though such an effect is negligible for diffusion rates on the order of hundreds m$^2$/day. For parameter set B, an insufficient number of sterile males using SITs can lead to an increase in the female population as diffusion rates are increased (see Fig. 5(c)). This is consistent with the results of the discrete model formulated by Yakob et al. (2008). In Fig. 5(d), we find that the RIDL method has a clear switch around $\theta = 1.0$ but the average fraction of surviving wild-type females is not sensitive to the release rate ratio for a given diffusion rate.

#### 3.2. Emerging outbreaks and the wavefront cover strategy

One finds four kinds of representative dynamics, determined by the release rate ratio, $\theta$. Fig. 6 illustrates results for SIT controls, whilst RIDL controls exhibit similar dynamics, except for the absence of an increase in the wild-type female population observed in Fig. 6(b), (c).

In Fig. 6(a) a sufficiently large release rate ratio drives the wave of wild-type female mosquitoes extinct before it can extensively disperse outside of the sterile male release region, and successful control is established. In (b), with a decrease in the release rate ratio, the invading wave of female mosquitoes reverses its direction of travel (i.e. the infested region contracts) and eventually the population becomes, again, extinct, though the wild-type female population size increases on reversal using SIT with parameter set B. In (c), in contrast, the wave of female mosquitoes ceases contraction and, in the SIT case, the female population increases.
Eradication is not achieved. Finally, in (d), with a further decrease in the release rate ratio we see that the female population wave is able to invade through the boundaries of the control region and eventually occupy the entire habitat. In cases (a) and (b), control is successful but in the cases (c) and (d), control fails. In what follows, we explore optimal strategies for control success.

3.2.1. Minimal release region size needed for complete eradication

Fig. 7 shows that the values of \( \theta, \frac{S_{\text{min}}}{S_N} \) are not sensitive to the choice of SIT or RIDL methods. For both parameter sets A and B, the variation in \( \frac{S_{\text{min}}}{S_N} \) is very small for \( \theta \in [0, 10] \). This implies that the minimal release region size, \( S_{\text{min}} \), varies only within several kilometres on a dimensional scale. Hence, the sensitivity of the minimal release region size to the release rate ratio is much less than in the case of the local release strategy. Such insensitivity is observed regardless of diffusion rates (results not shown). Since \( S_{\text{min}} \) converges to a constant value as the release rate ratio increases, once \( S_{\text{min}} \) is less than the threshold of \( S_{\text{min}}^\text{crit} \), the sterile/transgenic males always fail to impede the female wild-type wave, even for large release rate ratios. However, if \( S_{\text{min}} > S_{\text{min}}^\text{crit} \), the release rate ratio critically influences the dynamics of the wild-type females, as shown in Fig. 6 and the parameter region sketches of Fig. 7, and it determines the extent of control success.

Fig. 5. Average fraction of surviving wild-type females for different diffusion rates and release rate ratio with \( S_N = 500 \) km and \( S_S = 250 \) km, whereby eradication is not feasible. The plots give the normalised equilibrium female wild-type population in terms of \( \theta \), the control release rate ratio. The dotted line indicates 1.0 which is the normalised equilibrium population of female mosquitoes before control.

Fig. 6. Control success/fail scenarios for the emerging outbreak scenario. (a)-(d) plot representative cases for the SIT method with parameter set B. Similar dynamics are observed for the other parameter set or RIDL except that local increases in female density are not observed in cases (b, c). A and B, the variation in \( \frac{S_{\text{min}}}{S_N} \) is very small for \( \theta \in [0, 10] \). This implies that the minimal release region size, \( S_{\text{min}}^\text{crit} \), varies only within several kilometres on a dimensional scale. Hence, the sensitivity of the minimal release region size to the release rate ratio is much less than in the case of the local release strategy. Such insensitivity is observed regardless of diffusion rates (results not shown). Since \( S_{\text{min}}^\text{crit} \) converges to a constant value as the release rate ratio increases, once \( S_S \) is less than the threshold of \( S_{\text{min}}^\text{crit} \), the sterile/transgenic males always fail to impede the female wild-type wave, even for large release rate ratios. However, if \( S_S > S_{\text{min}}^\text{crit} \), the release rate ratio critically influences the dynamics of the wild-type females, as shown in Fig. 6 and the parameter region sketches of Fig. 7, and it determines the extent of control success.
In contrast to the results for the local release strategy, shown in Fig. 3(a), the threshold requirement of complete eradication for either the SIT or RIDL strategy induces relatively small changes in the minimal release region size even for parameter set B (Fig. 7(b)). In particular, the female wild-type population in the local release strategy for the endemic scenario remains at high levels and the density-dependency impacts strongly at the edges of the released sterile/transgenic male zone inducing different minimal release region sizes not only between SIT and RIDL but also between parameter sets. However, for wavefront covering strategies both edges of the female wild-type wave have low population density so that the effect of the density-dependence is slight, explaining the similarity of the behaviour of the SIT and RIDL strategies here.

3.2.2. Time to extinction and release rate ratio

In Fig. 8, we show the dependence of extinction time upon release rate ratio, given a sufficiently large and fixed release region size, $\gamma_S$. Since the extinction time is not measured precisely in the deterministic model, we define the extinction time for the female mosquitoes to be the minimal time satisfying Eq. (8). As expected, and also observed in a spatially homogeneous study by Atkinson et al. (2007), this eradication time increases drastically as the release rate ratio reduces towards the threshold. Indeed, for a release rate ratio on the order of the threshold value, and an eradication time of several years is predicted (Fig. 8(a), (c)). In contrast, release rate ratios significantly higher than threshold can reduce the time to extinction to the order of months (Fig. 8(b), (d)). Such predictions of the temporal dynamics can be made regardless of the choice of parameters or SIT/RIDL strategies. Nevertheless, the threshold release rate ratio for the RIDL technique is less than that for SIT and RIDL which always offers faster eradication, especially near threshold.

3.2.3. Release effort and strategy-cost

Before discussing results, we note that $[EF]_{\text{low}}$ and $[SC]_{\text{low}}$ given by Eqs. (9) and (10), respectively, depend on the initial size of the female

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**Fig. 7.** Wavefront cover strategy in the emerging outbreak scenario. Release rate ratio and relative release region size for complete eradication of the wild-type female mosquitoes. The threshold curve indicates successful control strategies. Above the curve control is successful and below the curve control fails. In (a), the parameter regions for the four representative dynamics of Fig. 6 are sketched. Similar parameter regions are also obtained in the case of parameter set B (details not shown). $D=1$ km$^2$/day, $\gamma_N = 373.5$ km in (a) and $\gamma_N = 325.5$ km in (b). For numerical simulations, we calculate $\gamma_N$ using Eq. (11). As we take very small values for $\epsilon$ in (11), the value of $\gamma_N$ used in our result is usually larger than $\gamma_{\text{min}}^S$ so that $\gamma_{\text{min}}^S/\gamma_N$ is less than 1.

**Fig. 8.** The dependence of extinction time on release rate ratio using the wavefront cover strategy in the emerging outbreak scenario given a fixed release region, $\gamma_S$, above $\gamma_{\text{min}}^S$ for all $\theta$. The diffusion constant is $D=1$ km$^2$/day. When the release rate ratio is small, the extinction time shows extreme sensitivity to the choice of control method. (a, b): Parameter set A. (c, d): Parameter set B.
mosquito wave, $\gamma N$, which is determined slightly differently depending on parameter sets A and B because the initial size of the wild-type female wave is given by simulation data for an invasive wave, using Eq. (4) with $S(x,t) = 0$. This differs between parameter sets A and B. Thus, strictly, we cannot use these two strategy measures directly for comparing the influence of the choice of parameter set. However, these two measurements are effective for exploring the effectiveness of SIT or RIDL using the same parameter set.

In Fig. 9 we present the results of a more detailed exploration of the release effort and strategy-cost for a fixed domain size $\gamma s$, in excess of $\gamma s_{\text{min}}$ for all release rates considered. The minimal effort values are subsequently given by the minimal release rate ratio regardless of the choice of SIT/RIDL strategies or parameter sets. Furthermore, as expected from Fig. 7, the release efforts using SIT and RIDL are identical. Nonetheless, we see non-trivial results for the strategy-cost, $[SC]_{\text{cov}}$, as shown in Fig. 9(c) and (d). Note that the eradication time decreases very rapidly once the release rate ratio is increased above the minimal release rate ratio required for complete eradication for the fixed value of $\gamma s$ used; furthermore, it becomes constant as the release rate ratio increases, as shown in Fig. 8. Therefore, the minimal value of strategy-cost exists not at the minimal release rate ratio but at a slightly larger value than the minimum, and it increases monotonically as the release rate ratio is further increased.

In general, the extinction time will decrease if we take a small initial size, $\gamma s$. This means that $[SC]_{\text{cov}}$ is dependent on $\gamma s$ and will decrease for smaller initial values of $\gamma s$. Obviously, an earlier initiation of a control strategy will be economically beneficial in the emerging outbreak scenario.

4. Discussion

When the release region of sterile/transgenic insects is sufficiently large, a temporal model for sterile/transgenic technologies may be enough to understand the potential for controlling pest insect populations. However, in practical situations this requires the release of sterile or transgenic insects over a long lengthscale, and therefore results in a heavy economic burden (Vreysen et al., 2007). Thus we are interested in finding the minimal value of the release region size, the release rate ratio (i.e. the number of sterile/transgenic males released per unit time) and time required for complete eradication. In particular, the minimal release region size is likely to be affected by the dispersal rate of the mosquitoes (Seirin-Lee et al., 2013). Thus a temporal model is insufficient and spatial models must be investigated carefully for a given invasion scenario. In addition, though an immediate difficulty in modelling studies is determining the levels of insect dispersal, with very limited empirical data and, potentially, a very wide range of estimates (Reiter et al., 1995; Harrington et al., 2005), a simple rescaling analysis can be used to account for the influence of dispersion in our modelling study, as illustrated in Appendix B.

In the first scenario where the wild-type female mosquitoes are endemic, our study demonstrates that sterile/transgenic males released locally in the habitat of the wild-type female mosquitoes can eradicate the vector insects completely with a larger size of release region. Nonetheless such a local release strategy easily fails if the diffusion rate of sterile/transgenic males is not high enough to ensure dispersal over the entire habitat. This result is consistent with those of a previous discrete model (Yakob et al., 2008).

Furthermore, our theoretical observations suggest that the local strategy is likely to be more applicable in a small region rather than a wide region because $\theta_{\text{opt}} = \gamma s_{\text{min}} \frac{\delta}{\gamma s}$ is determined independently of $\gamma s$ but depends on the diffusion rate. Furthermore, this difference in the size of the minimal release region relative to the region containing the established pest is predicted to be substantially larger than one might expect from the diffusive scale and the

Fig. 9. Release effort and strategy-cost values. The diffusion rate is $D = 1 \text{km}^2/\text{day}$ and the release region is the same as Fig 8, and thus fixed above $\gamma s_{\text{min}}$ for all $\theta$. (a, b) are the release effort values as a function of release rate ratio, and (c, d) are the strategy cost values, as given by (10), for varying release rate ratio, $\theta$. 

[Diagram of release effort and strategy-cost values]
timescale of either mosquito reproduction or death. Hence a local release strategy is predicted to be more readily applicable than one might initially anticipate from the scales of mosquito population dynamics. Nonetheless, in the local release strategy, the mosquito diffusion rate is a critical parameter in determining the optimal release region size, though the relation is a simple scaling law that can be readily predicted (see Appendix B). In turn, this means that one must carefully estimate mosquito dispersal rates in order to reduce control costs. Finally, we note that minimal overall strategy costs, in terms of total released mosquito numbers, are not minimised at the threshold of mosquito extinction, as shown in Fig. 4(d). Hence, increases in the release efforts, i.e. the unit time rate of release of control insects, can reduce the overall strategy cost regardless of the influence of spatial heterogeneity.

In the emerging outbreak scenario, our modelling study shows that several possible types of dynamics, depending on the release rate of sterile/transgenic males. However, the population dynamics are relatively insensitive to the release region size once the latter is larger than $r_{\text{min}}^S$ for all release rates. Furthermore, control interventions with a smaller strategy-cost do not always correspond to values of $(\gamma_S, \theta)$ that induce smaller release efforts. This demonstrates that a longer term picture, also considering eradication times, is required for efficient interventions aimed at eradicating an emerging outbreak.

The detailed requirements for inducing cost effective controls are predicted to differ with these two scenarios of a stable endemic and an emerging outbreak. For the endemic, the mosquito diffusion rate critically influences the minimal release region size for complete eradication. In contrast, control success is not highly sensitive to the diffusion coefficient for an emerging outbreak; instead the release rate ratio is an important and relatively sensitive parameter in determining the dynamics of the wild-type female wave.

Observations of the improved outcomes associated with RIDL strategies are inherited from the temporal model dynamics. In particular, once the suppression of larval competition by SIT interventions induces dynamically significant effects, as with parameter set B, RIDL strategies are substantially more effective in almost all aspects of control. Consequently, the typical conclusions that RIDL interventions are superior to SIT as a result of previous modelling (Atkinson et al., 2007; Phuc et al., 2007; White et al., 2010) do transfer in the context of local release and wavefront cover strategies. Similarly, local increases in pest populations can be associated with a SIT local release strategy or wavefront cover strategy, as observed in other contexts with overcompensating density-dependent competition (as in parameter set B) (Yakob et al., 2008; Yakob and Bonsall, 2009). These conclusions hinge on the fact that SITs reduce larval populations, enhancing the survival of insects resulting from wild-type matings and thus offsetting the reductions in proliferation. Thus RIDL strategies are never inferior in either control scenario considered. Nonetheless, once the release rate is chosen sufficiently large, both SIT and RIDL perform similarly for wavefront cover strategies with either parameter set, indicating that the governing dynamics of the model are then driven by the wild-type wavefront, where larval competition is minimal. This is in distinct contrast to predictions for control strategies designed to act as barriers to prevent the spread of mosquitoes into a pest-free region from an endemic area; here RIDL is predicted to be significantly superior (Seirin-Lee et al., 2013), highlighting that the control strategies are highly context dependent.

The timescale for a vector insect to become extinct is critical in terms of preventing a pandemic disease in a human society (Atkinson et al., 2007) and its increases are likely to induce serious fluctuations in insect populations by combining with external effects such as seasonality (Purse et al., 2005; Altizer et al., 2006; Yang et al., 2009; White et al., 2010). Large timescales are observed, in a spatially homogeneous modelling study on approaching the extinction threshold, by Atkinson et al. (2007) and we have analogous observations in our spatially heterogeneous setting. Thus, although a low release rate reduces the production costs of sterile/transgenic mosquitoes, it is also likely to be difficult to estimate or confirm control success in a situation where several years are required for eradication. Such long extinction times also drive our observation that the strategic cost illustrated in Fig. 9(d) for the emerging case has a local minimum, further reflecting the need to consider the longer term picture when designing interventions.

Throughout this manuscript, we have used fecundity and density-dependence parameter values based upon Dye (1984), concentrating on the extreme best and worst case scenarios, following previous approaches (Phuc et al., 2007; Yakob et al., 2008; White et al., 2010; Seirin-Lee et al., 2013). These parameters are derived from field data to which a simple regression is used to obtain the values. Legros et al. (2009) questioned this method and used a two-stage fitting method. They concluded that for their method (a) when density-independent processes are taken into consideration they account for a large part of the mortality of immature stages and density-dependence is much weaker than the Dye approach, (b) the functional responses of the two approaches are significantly different for the range of densities in the study, and (c) whilst both methods give reasonable accounts of the “characteristics of density-dependence”, they deviate when low densities are concerned, primarily due to the lack of data. Hence, it is critical that full life-table analyses are conducted in order to ensure that suitable estimates of these, and other (e.g. development periods, dispersal distances, differential density-dependent coefficients throughout the larval stages), life-history parameters be calculated, and at a local scale. For example, it has recently been shown that the dispersal ability of two lines of RIDL Ae. aegypti mosquitoes may be reduced compared to their wild-type counterparts in laboratory conditions (Bargielowski et al., 2012). This is likely to have an impact of the effectiveness of barrier zone techniques for population control. However, the difference in diffusion rates of the transgenic and wild-type mosquitoes is likely to add greater model complexity (Billingham and King, 2001). Furthermore, since it is likely that many additional biotic and abiotic factors may dynamically influence the life-histories of Ae. aegypti populations, both spatially and temporally (e.g. seasonality), further fine-tuning of control strategies will require these factors to be explicitly modelled. Extensions to our modelling approach could be adopted to incorporate these processes, but alternative approaches may also yield informative results, such as simulation models (e.g. Focks et al., 1993a, 1993b), additionally motivating a comparative study of differing modelling formulations.

In summary, the dispersion of mosquitoes appears in various invasive scenarios and our modelling study suggests successful control strategies for each scenario. Our results show that the requirements for understanding control effectiveness and efficient control strategy vary depending on the invasive and endemic scenario. Furthermore, SIT control is never more effective though the difference between RIDL and SIT strategies can be weak in the emerging outbreak strategy as the dynamics are dictated by the wavefront where competition is weak. Finally, we note that the long term picture is important in considering controls, due to the sensitivity of the extinction time for instance.

Finally, although the focus of our models is the mosquito, Ae. aegypti, which can spread yellow fever, dengue fever and Chikungunya disease, our modelling approach and results can be applied more broadly to other species. A further generalisation would be the consideration of more realistic measures of economic cost
rather than ones based on simply mosquito numbers. In addition, a pulse releasing schedule for sterile/transgenic mosquitoes may be more pragmatic and thus merits study, generalising the spatially homogeneous study of White et al. (2010). This is in progress, along with comparing whether and when modelling predictions are sensitive to the detailed representation of stage structure, for example contrasting models built on Dye’s (1984) delay formulation on one hand and ordinary differential equation representations of stage structure on the other (Focks et al., 1993a, 1993b; Erickson et al., 2010). Questions concerning higher dimensional geometries are also relevant, including smaller scale, three-dimensional models in high-rise buildings’ water tanks. In general the eikonal approximation indicates that the local frontwave behaviour possesses a curvature correction, which is sufficient to stabilise perturbations of a planar wave as well offering the prospect of complex global spatial dynamics such as spiral and scroll waves (Grindrod, 1991); whether such behaviours exist in mosquito models is a further open question.

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Appendix A. Numerical method

The reaction–diffusion systems formulated in this paper were solved numerically via standard techniques, which can readily accommodate the time delay; the kinetics are considered explicitly within a standard, fully implicit, finite difference treatment of the parabolic transport term (Morton and Mayers, 1994). In particular, storing the history of the system for the duration of the time delay allows the generation of the kinetic terms within the numerical algorithm. A fully implicit treatment of the diffusive terms then generates a set of linear algebraic equations for the mosquito populations at each new timepoint, which may be solved using a choice of numerical techniques; we use an LU-decomposition. This numerical algorithm has been validated against independent code simulations, used in Seirin-Lee et al. (2010), and we have checked timestep and grid spacing refinements do not influence the results presented.

Appendix B. Minimal release region size and diffusion rates

To explore the effects of diffusion rate in the model we use a scaling argument. Let \( D_{\text{ndim}} \) be a non-dimensionalised diffusion coefficient and define an arbitrary diffusion rate

\[
D = kD_0,
\]

for a given diffusion rate \( D_0 \) and arbitrary positive constant \( k \). Then for a time scale \( T \) and a given spatial length \( l \), we have

\[
D_{\text{ndim}} = \frac{DT}{l^2} = \frac{kD_0 T}{l^2} = \frac{D_0 T}{(l/k)^2}.
\]

From the above equation, we set a female habitat size, \( \gamma_N \), to be an arbitrary value by taking

\[
\gamma_N = l^2 / k,
\]

instead of choosing the diffusion rates arbitrarily.

On one hand, from Fig. 3(b) we know the optimal release region size, \( \delta_{\text{opt}} \), is independent of the spatial length scale so that it is also independent of the diffusion rate. That is, we have

\[
\delta_{\text{opt}} = \gamma_N (D) - \gamma''_N (D).
\]

Hence, we obtain the relationship between the diffusion rate and the optimal release region size directly from Eqs. (B.1)–(B.3), as

\[
y_N^{\text{opt}} (D) = y_N (D) - \delta_{\text{opt}} = \frac{\rho}{K} - \sqrt{D} \frac{\rho}{K} - \delta_{\text{opt}}.
\]

In Fig. 3(c), \( D_0 = 1 \text{ km}^2/\text{day} \), \( \gamma_0 = 500 \text{ km} \) and \( \delta_{\text{opt}} = 30 \text{ km} \) have been chosen.

References


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