Mathematical Biology



Pulled, pushed or failed: the demographic impact of a gene drive can change the nature of its spatial spread

Léna Kläy¹ · Léo Girardin² · Vincent Calvez² · Florence Débarre¹

Received: 26 October 2022 / Revised: 10 March 2023 / Accepted: 18 April 2023 /

Published online: 16 July 2023

© The Author(s), under exclusive licence to Springer-Verlag GmbH Germany, part of Springer Nature 2023

Abstract

Understanding the temporal spread of gene drive alleles—alleles that bias their own transmission—through modeling is essential before any field experiments. In this paper, we present a deterministic reaction-diffusion model describing the interplay between demographic and allelic dynamics, in a one-dimensional spatial context. We focused on the traveling wave solutions, and more specifically, on the speed of gene drive invasion (if successful). We considered various timings of gene conversion (in the zygote or in the germline) and different probabilities of gene conversion (instead of assuming 100% conversion as done in a previous work). We compared the types of propagation when the intrinsic growth rate of the population takes extreme values, either very large or very low. When it is infinitely large, the wave can be either successful or not, and, if successful, it can be either pulled or pushed, in agreement with previous studies (extended here to the case of partial conversion). In contrast, it cannot be pushed when the intrinsic growth rate is vanishing. In this case, analytical results are obtained through an insightful connection with an epidemiological SI model. We conducted extensive numerical simulations to bridge the gap between the two regimes of large and low growth rate. We conjecture that, if it is pulled in the two extreme regimes, then the wave is always pulled, and the wave speed is independent of the growth rate. This occurs for instance when the fitness cost is small enough, or when there is stable coexistence of the drive and the wild-type in the population after successful drive invasion. Our model helps delineate the conditions under which demographic dynamics can affect the spread of a gene drive.

Mathematics Subject Classification 92-10 · 92D25

Institut Camille Jordan, UMR 5208 CNRS and Universite Claude Bernard Lyon 1, Villeurbanne, France



 [∠] Léna Kläy lena.klay@sorbonne-universite.fr

Institute of Ecology and Environmental Sciences Paris (IEES Paris), Sorbonne Université, CNRS, IRD, INRAE, Université Paris Est Creteil, Université de Paris, Paris Cedex 5, Paris, France

30 Page 2 of 54 L. Kläy et al.

Contents

1	Introduction	3
2	Methodology	
	2.1 Models	5
	2.2 Setting of the problem	7
	Traveling waves	7
	Numerical simulations	7
	2.3 Glossary	8
	Allelic densities and frequencies	8
	Classification of the dynamics	8
	Monostable/Bistable systems	9
	Pulled and pushed waves	9
2		10
3	Results	
	3.1 Model with perfect conversion in the zygote	10
	3.1.1 Preliminary statements on the model	10
	$3.1.2 r = +\infty \dots $	11
	$3.1.3 r = 0 \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	12
	3.1.4 Comparison between the outcomes when $r = +\infty$ and $r = 0$	13
	3.2 Models with partial conversion	14
	3.2.1 Conversion occurring in the zygote	14
	3.2.2 Conversion occurring in the germline	21
	3.2.3 Conclusion	23
4	Discussion	24
	On the final state of the population	24
	On the transient regime (propagation of waves)	24
	Perspectives	25
A	ppendix	26
	Model with partial conversion: growth term details	26
^	A.1 Conversion occurring in the zygote	27
	A.2 Conversion occurring in the germline	28
Б	A.2 Conversion occurring in the germline	28 29
В	System rewritten with variables (n, p_D)	
	B.1 Model with perfect conversion	29
	B.2 Model with partial conversion	30
	B.2.1 Conversion in the zygote	30
	B.2.2 Conversion in the germline	31
C	Proofs for model (12) with perfect conversion in the zygote	33
	C.1 Numerical evidence for the continuity when $r \to 0$	33
	C.2 Proof of the statements in Tables 3 and 4 when perfect conversion occurs in the zygote	
	C.2.1 Gene drive clearance for $s \in (0.5, 1)$ when $r = 0$	35
	C.2.2 Numerical approximation of s threshold value for the pulled/pushed wave	
	when $r = +\infty$	36
D	Critical travaling wave for an SI similar model	36
	D.1 Existence of critical traveling wave solutions	37
	D.2 Proof of the theorem	37
	D.2.1 Construction of sub- and super-solutions	38
	D.2.2 Existence and positivity of a critical traveling wave solution	42
E	Study of the reaction term when $r = +\infty$ in section 3.2	42
_	E.1 Conversion occurring in the zygote	42
	E.1.1 Monostable drive invasion	44
	E.1.2 Monostable wild-type invasion	45
		43
	E.1.3 Monostable coexistence state	
	E.2 Conversion occurring in the germline	46
	E.2.1 Monostable drive invasion	47
	E.2.2 Monostable wild-type invasion	49
	E.2.3 Monostable coexistence state	50
F.	Heatmap supplementary materials	51



Effect of fitness disadvantage (s) and dominance coefficient (h) on drive		
dynamics, for $r = +\infty$	51	
F.2 Heatmap lines	52	
F.2.1 Pure drive line	52	
F.2.2 Composite persistence line	52	
afarances	53	

1 Introduction

A highly accurate, cost-effective and easy-to-use technology, the CRISPR-Cas genome editing system has been favoring the development of promising innovations (Jinek et al. 2012). Among them, CRISPR-Cas9 gene drive (Alphey 2020), which aims to spread a trait of interest in a wild type population in a relatively short number of generations (Kyrou et al. 2018). Application fields are numerous, and include (i) the eradication of insect-borne diseases (Buchman 2020; Gantz et al. 2015; Kyrou et al. 2018); (ii) the elimination of herbicide and pesticide resistance in pest populations (Neve 2018); (iii) the control of destructive invasive species (Gantz and Bier 2015; Grunwald et al. 2019); (iv) the conservation of biodiversity by spreading beneficial traits in endangered species (Esvelt et al. 2014; Rode et al. 2019).

Targeting sexually reproducing species, CRISPR-Cas9 gene drive biases the transmission of an allele from a parent to its offspring. This biased inheritance occurs through gene conversion (also called "homing" (Deredec et al. 2008)): in a heterozygous cell, the gene drive cassette present on one chromosome induces a double-strand break at a specific target site on the homologous chromosome, and the repair process duplicates the cassette. Overall, this process increases the chances of transmitting the gene drive cassette compared to its wild-part counterpart, and the mechanism repeats through the generations. Gene conversion can potentially take place at different timings of the life cycle: from very early on, in the zygote, meaning that potentially every single cell of the individual could become homozygous for the gene drive, to, in the germline, where only the gametes are converted.

Gene drives can be classified into two main categories depending on the purpose of their use (Dhole et al. 2020; Girardin and Débarre 2021). A "replacement drive" is aimed at spreading a genetic modification in order to introduce an important and durable feature in the natural population. Population size is then not significantly affected and the drive construct may in principle persist indefinitely in the environment. A "suppression drive" on the other hand is meant to reduce population size by spreading a detrimental trait, such as a sex ratio distorter (Male Bias and Female Fertility 2022) or by altering fertility (Kyrou et al. 2018), for example. The term "eradication drive" can be used for the extreme case where population extinction is the aim.

As with any new tool, it is essential to balance risks (safety) and benefits (efficacy) of the technique before running any field trials. Experiments currently conducted in laboratories provide small- to medium-scale information; mathematical models can help to extend these empirical results and identify the features that are the most important in determining the dynamics at larger scales (Committee on Gene Drive Research in Non-Human Organisms 2016).



30 Page 4 of 54 L. Kläy et al.

Early gene drive models (Burt 2003; Deredec et al. 2008; Unckless et al. 2015) used classical population genetics frameworks, and considered discrete non-overlapping generations in a well-mixed population. These simplifications helped to draw general conclusions, but it is important to challenge them. First of all, most of the species targeted in the context of gene drive do not have synchronous generations (for instance mosquitoes (Gantz et al. 2015; Hammond et al. 2015; Buchman 2020; Kyrou et al. 2018), flies (Gantz and Bier 2015), mice (Grunwald et al. 2019)). Secondly, the assumption of a single well-mixed collection of individuals living across a uniform space is usually not realistic. In fact, most of the natural landscapes are heterogeneous. Individuals are also more likely to interact with others that are in closer proximity, which might result in local genetic variations. Finally, releases of transgenic individuals are limited in range, which is another factor of spatial heterogeneity.

Taking into account spatio-temporal dynamics of the population size is another key step towards more realistic models. For the sake of simplicity, most early models focused on allele frequencies and considered a constant population density. However in the context of gene drive, the introduction of maladapted transgenic individuals can lead to the reduction (or even extinction) of the population (Dhole et al. 2020). When considering a spatially structured population, variations in population density naturally generate a demographic flux from denser to less dense areas. This demographic flux is directed in opposition to the spread of the drive allele. It was previously shown (Girardin and Débarre 2021) that the advantage conferred by gene conversion may nevertheless counteract the demographic effect linked to the fitness cost.

The main goal of this paper is to clarify the impact of variations in population density over the course of drive propagation over space.

We study partial differential equations which follow the propagation of the drive in space and time. We explore numerically and analytically two models: a first model based on perfect conversion in the zygote, already introduced in Girardin and Débarre (2021) in a spatially structured population, corresponding to an idealized case where gene conversion always succeeds; second, a more realistic model with partial conversion and presence of heterozygous individuals, already studied in Rode et al. (2019) in a well-mixed, non spatial population. In order to investigate the possible spreading of gene drives through space after local introduction, we focus on the description of traveling waves solutions, that is, particular solutions which are stationary in a frame moving at constant speed. Our analysis goes beyond (Girardin and Débarre 2021) by several means: we extend it to the case of partial conversion, and we systematically analyze the case where the demographic effects are the strongest, in the regime of vanishing growth rate. The latter is possible through an insightful connection with an epidemiological SI model.



2 Methodology

2.1 Models

We present our model step-by-step. For a genetically and spatially homogeneous population, we consider the following (non-dimensionalized) equation:

$$\partial_t n(t) = \left(1 + r\left(1 - n(t)\right)\right) f(t) - n(t) \quad (\forall t > 0). \tag{1}$$

where the unit of time is generations. Fecundity is density-dependent, and parametrized by the fitness f and the rate r at which the (f-dependent) carrying capacity is restored. When f = 1, the carrying capacity is 1, and we recover the logistic equation $\partial_t n(t) = r (1 - n(t)) n(t)$. Other modeling options are discussed in Girardin and Débarre (2021).

Then, we add genetic diversity in the population. We still denote by n the total density, and by n_i the density of individuals with genotype i. The population we consider is diploid, sexually reproducing, and the fitness f_i depends on the genotype. The dynamics are given by the following equations:

$$\partial_t n_i(t) = \left(1 + r(1 - n(t))\right) f_i \ n(t) \underbrace{\sum_{l,k} \pi_{l,k}^i \frac{n_l(t)}{n(t)} \frac{n_k(t)}{n(t)}}_{\text{Mating term}} - n_i(t) \qquad (\forall t > 0) \ (\forall i).$$

The mating term takes into account the probability for each couple of parents l,k to have offspring of type i $(\pi_{l,k}^i)$, multiplied by the probability of a mating event l,k $(\frac{n_l(t)n_k(t)}{n(t)^2})$, assuming random mating.

Last but not least, we consider a spatially structured population. We assume that the movement of individuals is described by a diffusion term with equal diffusion coefficients, normalized to 1. Since we focus on traveling wave solutions, we restrict our analysis to a one-dimensional space. We obtain the following equations:

$$\partial_{t} n_{i}(t, x) - \partial_{xx}^{2} n_{i}(t, x) = \left(1 + r(1 - n(t, x))\right) f_{i} n(t) \sum_{l, k} \pi_{l, k}^{i} \frac{n_{l}(t)}{n(t)} \frac{n_{k}(t)}{n(t)} - n_{i}(t, x)$$

$$(\forall t > 0) \ (\forall x \in \mathbb{R}) \ (\forall i). \tag{3}$$

There are two possible alleles at the locus that we consider: the wild-type allele (W) and the drive allele (D). We have three genotypes: wild-type homozygotes (i = WW), drive homozygotes (i = DD) and heterozygotes (i = DW). Wild-type homozygotes have fitness $f_{WW} = 1$, drive homozygotes have fitness $f_{DD} = 1 - s$, where s is the fitness cost of the drive, and drive heterozygotes have fitness $f_{DW} = 1 - sh$, where h is the dominance parameter (see Table 1).

All along the paper, we assume $s \in (0, 1)$, corresponding to a fitness cost carried by the drive alleles. Furthermore, we assume that the fitness of heterozygotes cannot be greater than the fitness of either homozygote $(h \in [0, 1])$.



30 Page 6 of 54 L. Kläy et al.

Gene conversion turns a heterozygous cell into a drive homozygous cell. To determine the probability $\pi^i_{l,k}$ (probability for a couple l,k to have offspring of type i), we need to take into account both the probability $c \in [0, 1]$ with which gene conversion occurs in heterozygotes, and the stage of the life cycle at which it occurs: either in the zygote, or in the germline. This last feature modifies significantly the probabilities: for example, a couple W, D of gametes has a probability 1-c to lead to heterozygous offspring if conversion occurs in the zygote, whereas this probability becomes one if conversion occurs in the germline. We detail all $\pi^i_{l,k}$ values in Appendix A. For the sake of clarity, we now omit variables in the notation $(n_i = n_i(t, x))$.

The parameters are summarized in Table 2.

In this article, we will analyse the three following versions/variations of model (3):

Partial conversion occurring in the zygote:

$$\begin{cases} \partial_{t} n_{\text{DD}} - \partial_{xx}^{2} n_{\text{DD}} = (1 - s)(r (1 - n) + 1) & \frac{c \ n_{\text{WW}} n_{\text{DW}} + 2 \ c \ n_{\text{WW}} n_{\text{DD}} + (\frac{1}{2} \ c + \frac{1}{4}) \ n_{\text{DW}}^{2} + (c + 1) \ n_{\text{DW}} n_{\text{DD}} + n_{\text{DD}}^{2}}{n} - n_{\text{DD}}, \\ \partial_{t} n_{\text{DW}} - \partial_{xx}^{2} n_{\text{DW}} = (1 - sh)(r (1 - n) + 1) \ (1 - c) & \frac{n_{\text{WW}} n_{\text{DW}} + 2 \ n_{\text{WW}} n_{\text{DD}} + \frac{1}{2} \ n_{\text{DW}}^{2} + n_{\text{DW}} n_{\text{DD}}}{n} - n_{\text{DW}}, \\ \partial_{t} n_{\text{WW}} - \partial_{xx}^{2} n_{\text{WW}} = (r (1 - n) + 1) & \frac{n_{\text{WW}}^{2} + n_{\text{WW}} n_{\text{DW}} + \frac{1}{4} \ n_{\text{DW}}^{2}}{n} - n_{\text{WW}}. \end{cases}$$

$$(44)$$

Partial conversion occurring in the germline:

$$\begin{cases} \partial_{t} n_{\text{DD}} - \partial_{xx}^{2} n_{\text{DD}} = (1 - s)(r (1 - n) + 1) & \frac{\frac{1}{4} (1 + c)^{2} n_{\text{DW}}^{2} + (1 + c) n_{\text{DW}} n_{\text{DD}} + n_{\text{DD}}^{2}}{n} - n_{\text{DD}}, \\ \partial_{t} n_{\text{DW}} - \partial_{xx}^{2} n_{\text{DW}} = (1 - sh)(r (1 - n) + 1) & \frac{(1 + c) n_{\text{WW}} n_{\text{DW}} + 2 n_{\text{WW}} n_{\text{DD}} + \frac{1}{2} (1 - c^{2}) n_{\text{DW}}^{2} + (1 - c) n_{\text{DW}} n_{\text{DD}}}{n} - n_{\text{DW}}, \\ \partial_{t} n_{\text{WW}} - \partial_{xx}^{2} n_{\text{WW}} = (r (1 - n) + 1) & \frac{n_{\text{WW}}^{2} + (1 - c) n_{\text{WW}} n_{\text{DW}} + \frac{1}{4} (1 - c)^{2} n_{\text{DW}}^{2}}{n} - n_{\text{WW}}. \end{cases}$$

$$(5)$$

Table 1 Population characteristics (D: Drive allele, W: Wild-type allele)

Adult genotype	Density	Fitness
Drive Homozygote (DD)	n_{DD}	1-s
Heterozygote (DW)	n_{DW}	1 - sh
Wild-type Homozygote (WW)	n_{WW}	1

Table 2 Model parameters

Parameters	Range values	Description
r	$(0, +\infty)$	Intrinsic growth rate
c	[0, 1]	Conversion rate
S	(0, 1)	Fitness cost of drive homozygotes
h	[0, 1]	Drive dominance



Perfect conversion occurring in the zygote (no heterozygotes):

For a perfect conversion occurring in the zygote (c = 1), model (4) reduces to the following set of two equations, which was introduced in Girardin and Débarre (2021):

$$\begin{cases} \partial_{t} n_{\text{DD}} - \partial_{xx}^{2} n_{\text{DD}} &= (1 - s) \left(r \left(1 - n_{\text{DD}} - n_{\text{WW}} \right) + 1 \right) \frac{n_{\text{DD}}^{2} + 2 n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}} - n_{\text{DD}} = F_{D}(n_{\text{DD}}, n_{\text{WW}}) \\ \partial_{t} n_{\text{WW}} - \partial_{xx}^{2} n_{\text{WW}} &= \left(r \left(1 - n_{\text{DD}} - n_{\text{WW}} \right) + 1 \right) \frac{n_{\text{WW}}^{2}}{n_{\text{WW}} + n_{\text{DD}}} - n_{\text{WW}} = F_{W}(n_{\text{DD}}, n_{\text{WW}}). \end{cases}$$
(6)

This last model only follows the two homozygous genotypes, drive and wild-type. Due to perfect gene conversion (c=1), no heterozygous individuals are ever produced: heterozygous eggs are all transformed into homozygotes. Further assuming that there are no heterozygotes initially, we only need to follow the densities of homozygotes.

Note that system (6) can also be obtained from model (5) by assuming perfect conversion in the germline (c=1) and drive dominance (h=1). In this case, heterozygotes and drive homozygotes have the same fitnesses, and both only produce gametes with the drive allele. We can then group them together and follow their density $n_{\rm DW} + n_{\rm DD}$, whose dynamics are given by the first line of (6).

2.2 Setting of the problem

Traveling waves

We seek stationary solutions in a reference frame moving at speed v, where v is some unknown:

$$\begin{cases} n_{\mathrm{DD}}(t,x) = n_{\mathrm{DD}}(x-vt) & (\forall t > 0) \ (\forall x \in \mathbb{R}), \\ n_{\mathrm{DW}}(t,x) = n_{\mathrm{DW}}(x-vt) & (\forall t > 0) \ (\forall x \in \mathbb{R}), \\ n_{\mathrm{WW}}(t,x) = n_{\mathrm{WW}}(x-vt) & (\forall t > 0) \ (\forall x \in \mathbb{R}). \end{cases}$$
(7)

Traveling wave solutions contain important information for the biological interpretation of the results, such as the speed of invasion v, the genetic composition of the expanding population, or the final equilibrium. In this paper, we focus our study on this mathematical object and detail below the vocabulary we use. Key to our analysis are the notions of monostable or bistable systems, and whether the traveling wave is pulled or pushed. There may be confusion around these concepts in the literature, so we clarify their definitions below.

Numerical simulations

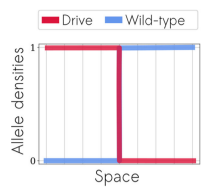
We complement our mathematical analysis with numerical simulations of the Cauchy problem, with initial conditions for each genotype specified as in Fig. 1. The outcomes of the simulations are heatmaps of the expansion speed over a wide range of parameters.

Initial conditions for numerical simulations are as follows: the left half of the domain is full of drive ($n_{DD} = 1$), and the right half is full of wild-type ($n_{WW} = 1$) (see Fig. 1).



30 Page 8 of 54 L. Kläy et al.

Fig. 1 Initial conditions for numerical simulations. The left half of the domain is full of drive $(n_{\rm DD}=n_D=1)$, and the right half is full of wild-type $(n_{\rm WW}=n_W=1)$



The code is available on GitHub (https://github.com/LenaKlay/gd_project_1). We ran our simulations in Python 3.6, with the Spyder environment. Heavy heatmaps 4, 5, 6A, 6B; 7 have been computed thanks to the INRAE Migale bioinformatics facility (doi: 10.15454/1.5572390655343293E12). We are grateful to them for providing these computing resources.

2.3 Glossary

Allelic densities and frequencies

For our analysis, it is convenient to introduce the allelic (half-) densities $(n_{\rm D}, n_{\rm W})$. The precise definition depends on the model, and more specifically on the timing of conversion. In fact, we have $n_{\rm D}=n_{\rm DD}+\alpha$ $n_{\rm DW}$ and $n_{\rm W}=n_{\rm WW}+(1-\alpha)$ $n_{\rm DW}$, with $\alpha=\frac{1}{2}$ when conversion occurs in the zygote, and $\alpha=\frac{1+c}{2}$ when conversion occurs in the germline (see Sect. 3.2). Depending on the regime of parameters, it may be more appropriate to study the allelic frequencies $p_{\rm D}=\frac{n_{\rm D}}{n_{\rm D}+n_{\rm W}}$, $p_{\rm W}=\frac{n_{\rm W}}{n_{\rm D}+n_{\rm W}}$.

Classification of the dynamics

It can happen that the dynamics lead to the decay of the drive allele uniformly in space. In this case, there cannot exist a traveling wave for the drive population: we use the term *gene drive clearance* to describe this situation. Then, the problem boils down to the standard Fisher-KPP traveling wave problem for the expansion of the wild-type in the absence of a drive (see (Girardin and Débarre 2021)).

When traveling waves do exist, we distinguish between two cases depending on the sign of the speed. When v > 0, the wave moves to the right: it is a *drive invasion*. When v < 0, the wave moves to the left: it is a *wild-type invasion*. In some specific cases, drive and wild-type invasions can happen simultaneously: the waves decompose into two sub-traveling wave solutions over half of the domain. They move in opposite directions and lead to the coexistence of both alleles in-between.

In case of drive invasion, we distinguish several cases depending on the state of the population in the wake of the front(s): i) *eradication* drives are those for which the population vanishes in the wake of the front(s); ii) *suppression* drives are those for



which population persists in the wake of the front(s). In the latter case, two scenarios are possible: persistence of drive homozygotes only; persistence of all genotypes.

Monostable/Bistable systems

To illustrate useful concepts in the theory of reaction-diffusion equations, we consider the following standard equation of population genetics (Otto and Day 2011) describing the dynamics of the frequency p of an allele of interest:

$$\partial_t p - \partial_{xx}^2 p = p (1 - p) \sigma(p) \quad \text{with } p \in [0, 1], \tag{8}$$

where $\sigma(p)$ is the selection term, which we consider frequency-dependent (i.e., function of p).

If σ is of constant sign, say $\sigma > 0$, this equation is referred to as a monostable case. Then, the solution converges locally to the unique stable equilibrium p=1 (or p=0 if $\sigma < 0$). If σ is changing signs once in (0,1), being negative below some threshold, and positive above, it is referred to as a bistable case. In the latter case, the solution converges locally to one of the two stable equilibria p=0 or p=1, depending on the initial condition. Moreover, each equilibria has a basin of attraction and there is a threshold effect – hence the name "threshold-dependent drives" in the gene drive literature to describe this kind of case (for example in reference (Tanaka et al. 2017)).

In both cases, there exist traveling waves connecting the two equilibria p=0 and p=1. A straightforward integration by parts shows that, whatever the stability, the sign of the wave speed satisfies

$$\operatorname{sign}(v) = \operatorname{sign}\left(\int_0^1 p(1-p)\sigma(p)dp\right). \tag{9}$$

In monostable cases with $\sigma > 0$, this sign is positive; in bistable cases, however, it depends on the details of the frequency-dependence σ . Moreover, under some circumstances (bistable case, or degenerate monostable case), the invasion outcome for the Cauchy problem can be changed by modifying the inoculum size. Even if traveling waves exist such that p=1 is invading p=0, small initial conditions may not succeed in propagating in space, see the discussion in Tanaka et al. (2017); Turelli and Barton (2017); Nadin et al. (2018).

By analogy with the scalar case, we consider that a system is monostable if it has exactly one stationary stable state, and bistable if it has exactly two stationary stable states.

Pulled and pushed waves

Usually, a wave is said to be pulled if the wave speed coincides with the minimal speed of the linearized problem at low density (resp. low frequency). This occurs when the population at low density (resp. low frequency) has sufficient reproductive success to determine the dynamics of the full invasion.



30 Page 10 of 54 L. Kläy et al.

Conversely, a wave is said to be pushed if the wave speed is strictly larger than the minimal speed of the linearized problem. In contrast with pulled waves, the whole population contributes to the dynamics of invasion.

A bistable wave is clearly pushed (Hadeler and Rothe 1975). However, a monostable wave can be either pulled or pushed, see (An et al. 2022; Holzer 2022; Birzu et al. 2018) and discussion therein. Nonetheless, a monostable wave is necessarily pulled if the per-capita growth rate is maximal at low density (resp. low frequency). In the particular case of the scalar problem (8), this criterion simply writes:

$$\sigma(0) > (1-p) \, \sigma(p) \quad (\forall p \in [0,1]).$$
 (10)

3 Results

In part 3.1, we study the model with perfect conversion in the zygote (6) and compare the qualitative behavior of the solution when r=0 and $r=+\infty$. In part 3.2, we proceed the same way on models with partial conversion (4) and (5), obtaining more general results.

3.1 Model with perfect conversion in the zygote

3.1.1 Preliminary statements on the model

We introduce a few general results on model (6) when r > 0, which will be useful in the study.

When $s \le \frac{1}{2}$, system (12) is monostable: the only stable state is $(n_{\text{DD}} = n_{\text{DD}}^*, n_{\text{WW}} = 0)$ with $n_{\text{DD}}^* = \min(0, 1 - \frac{s}{r(1-s)})$ (Girardin and Débarre 2021), leading to a drive invasion if any. We introduce the minimal speed of problem (6) linearized at low drive density, i.e. the speed of any pulled wave in case of a drive invasion:

$$2\sqrt{\partial_{n_{\rm DD}}F_D(0,1)} = 2\sqrt{1-2s}. (11)$$

When $s > \frac{1}{2}$, system (12) is bistable. Consequently traveling waves are either semi-trivial ($n_{\rm DD} = 0$ identically, standard Fisher-KPP problem for $n_{\rm WW}$) or pushed.

For our analysis, it will be convenient to rewrite model (6) so that it follows the frequency of the drive $p_{\rm D} = \frac{n_{\rm D}}{n_{\rm D} + n_{\rm W}} = \frac{n_{\rm DD}}{n_{\rm WW} + n_{\rm DD}}$ (because $n_{\rm DW} = 0$) and total population density $n = n_{\rm WW} + n_{\rm DD}$ (details in B.1):

$$\begin{cases} \partial_{t} p_{\mathrm{D}} - \partial_{xx}^{2} p_{\mathrm{D}} = 2 \, \partial_{x} (\log n) \, \partial_{x} p_{\mathrm{D}} + (r \, (1-n)+1) \, s \, p_{\mathrm{D}} \, (1-p_{\mathrm{D}}) \, \left(p_{\mathrm{D}} - \frac{2s-1}{s} \right), \\ \partial_{t} n - \partial_{xx}^{2} n &= (r \, (1-n)+1) \left(1-s + s (1-p_{\mathrm{D}})^{2} \right) \, n - n. \end{cases}$$
(12)

System (12) differs from standard equations often used in populations genetics as it contains an advection term 2 $\partial_x(\log n)$ $\partial_x p_D$. This term appears when calculating $\partial_{xx}^2 p_D = \partial_{xx}^2 \frac{n_{DD}}{n}$ (details in B.1) and represents a demographic flux from denser to



s value	$0 < s \lesssim 0.35$	$0.35 \lesssim s < 1/2$	$1/2 < s \lesssim 0.70$	$0.70 \lesssim s < 1$
Stability	Monostal	ble	В	istable
Speed	$v = 2\sqrt{1 - 2s}$	$v > 2\sqrt{1-2s}$	v > 0	v < 0
Wave	Pulled wave	Push	ed wave	Pushed wave
Invasion	Drive frequency	Drive invasion		Wild-type invasion

Table 3 Traveling waves study for Model (13), limit of system (6) when $r = +\infty$. All statements in the table are proved in Appendix C.2

less dense areas, due to variations in population density. It is opposed to the spread of the (costly) drive allele (see Figure 2 (Girardin and Débarre 2021)). We observe a singularity for n=0 in both formulations of the system: in (6) due to $\frac{1}{n_{\rm DD}+n_{\rm WW}}$ and in (12) due to $\log(n)$. This should be handle with care.

$3.1.2 r = +\infty$

The limit of system (12) when $r \to +\infty$ has already been determined in Girardin and Débarre (2021). Using the Strugarek-Vauchelet rescaling (Strugarek and Vauchelet 2016), the following limit equation is obtained, which was also previously introduced in Tanaka et al. (2017):

$$\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D} = \frac{s \ p_{\rm D} (1 - p_{\rm D}) \left(p_{\rm D} - \frac{2s - 1}{s} \right)}{1 - s + s(1 - p_{\rm D})^2}.$$
 (13)

Interestingly, equation (13) is independent of the population density n and it does not contain the advection term $2 \partial_x (\log n) \partial_x p_D$. This is due to the fact that the population size n(t, x) remains spatially homogeneous after the introduction of drive individuals, when $r \to +\infty$. Intuitively, so many offspring are produced at each generation that the carrying capacity is instantaneously restored, and losing a fraction s of these offspring by selection has no consequence. Therefore the variations in population density (n), and consequently the demographic flux, are negligible.

Equation (13) has a single parameter, the fitness cost of the drive s. The numerical value of the threshold for the transition from positive to negative speed (≈ 0.70) was already known (Tanaka et al. 2017; Girardin and Débarre 2021), and can be computed to arbitrary precision by the formula (9). The numerical value of the threshold for the transition from pulled to pushed (≈ 0.35 up to two digits) was numerically computed by a continuation method following (Avery et al. 2022; Holzer 2022).

Note that equation (13) shows qualitative similarities with a common equation in the population genetics literature (Barton 1979) (and which is actually an approximation of (13) under weak selection, i.e. $s \to 0$):



30 Page 12 of 54 L. Kläy et al.

$$\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D} = s \ p_{\rm D} (1 - p_{\rm D}) \left(p_{\rm D} - \frac{2s - 1}{s} \right).$$
 (14)

Quantitatively, the thresholds are $\frac{2}{5}$ instead of 0.35, and $\frac{2}{3}$ instead of 0.70 (analytical values) (Hadeler and Rothe 1975).

3.1.3 r = 0

When the intrinsic growth rate r is finite, it is expected that the final population density after the invasion of the drive (if any) is strictly below 1, because of the fitness cost. The smaller r, the lower the final size. The spatial effect of demography on gene drive expansion is expected to be maximal as r vanishes, when the population can hardly restore its carrying capacity, leading to a high amplitude of the population size gradient $2 \partial_x (\log n)$. In this section we focus on the limit r = 0, which maximizes the demographic impact of the fitness cost on drive propagation.

In a purely wild-type population, the case r=0 corresponds to a number of births balancing exactly the number of deaths. As soon as the drive allele is introduced, this balance is locally broken, yielding a net decrease in the population size. Then, the drive can either propagate by leaving empty space behind, or disappear. The same conclusion holds as long as $r < \frac{s}{1-s}$, see Sect. 3.1.1.

We checked numerically that the wave speed is continuous in the limit $r \to 0$. Therefore, each conclusion on the case r = 0 sheds some light on the case of small r (see heatmap in Appendix C.1).

As discussed above, we cannot just consider a single equation on the drive frequency p_D when r is finite because of the demographic contribution $2\partial_x \log n$. Interestingly, in the case r = 0, the demographic system (6) reduces to the following pair of equations:

$$\begin{cases} \partial_t n_{\text{WW}} - \partial_{xx}^2 n_{\text{WW}} = \frac{-n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}}, \\ \partial_t n_{\text{DD}} - \partial_{xx}^2 n_{\text{DD}} = (1 - s) \frac{n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}} - s n_{\text{DD}}. \end{cases}$$
(15)

Noticeably, the previous system shares some features with density-dependent epidemiological SI models. In particular, the dynamics of $n_{\rm WW}$ is always decreasing. The dynamics of $n_{\rm DD}$ is the balance of creation and linear decay. By changing notations $n_{\rm WW} \leftrightarrow S$ (susceptible individuals), and $n_{\rm DD} \leftrightarrow I$ (infected individuals), (15) can be recast as follows:

$$\begin{cases} \partial_t S - \partial_{xx}^2 S = -\beta_1 \frac{SI}{S+I}, \\ \partial_t I - \partial_{xx}^2 I = \beta_2 \frac{SI}{S+I} - \gamma I. \end{cases}$$
 (16)

with $\beta_1 = 1$, $\beta_2 = (1 - s)$ (transmission parameters), and $\gamma = s$ (disease clearance). Usually, in SI models, individuals of type S are all transformed into individuals of type I at infection, hence $\beta_1 = \beta_2$. In our case, these two rates are distinct because



s value	0 < s < 1/2	1/2 < s < 1	
Stability	Monostable	Degenerate case	
Speed	$v = 2\sqrt{1 - 2s}$	No wave	
Wave	Pulled wave		
	Drive invasion	Gene drive clearance	
Invasion	Drive frequency Space	Drive frequency Space	

Table 4 Traveling waves study for Model (15), limit of system (6) when r = 0. All statements in the table are proved in Appendix C.2

of the fitness cost of the drive. The existence of traveling waves for model (15) with $\beta_1 = \beta_2$ has been studied recently in the literature (Zhou et al. 2019). Here, we extend the results of reference (Zhou et al. 2019) to a more general case $0 < \beta_1$ and $0 < \beta_2$. This leads to the characterization in Table 4 and Appendix D.

In contrast to the results obtained when $r=+\infty$, when r=0 there is only one threshold value of s determining the outcome of the model (Table 4). When 0 < s < 1/2, the system is monostable, the drive necessarily invades. Moreover, the wave is pulled and travels at speed $v=2\sqrt{1-2s}$ (11). When 1/2 < s < 1, the problem is degenerate: there exists a family of steady states, corresponding to homogeneous $n_{\rm WW} \in [0,1]$ and $n_{\rm DD}=0$. It is a case of gene drive clearance, as $n_{\rm DD}$ converges to zero uniformly in space (at rate at least 1-2s). However, the final density of wild-type is not clearly determined, as it boils down to diffusion only in the large time asymptotics (details in Appendix C.2.1). Note that this conclusion holds in a well-mixed population (without spatial consideration): the drive decays uniformly and the final density of wild type depends on the initial data.

3.1.4 Comparison between the outcomes when $r = +\infty$ and r = 0

The differences between the two regimes are strongest for intermediate values of s. When $1/2 < s \le 0.70$, the drive can spread when the demographic consequences are negligible $(r = +\infty)$. However, such a costly drive cannot invade when the intrinsic growth rate r is very low (r = 0). When $0.35 \le s < 1/2$, the drive wave advances for both $r = +\infty$ and r = 0. However, it is of different nature: the wave is pulled when r = 0, while it is pushed when $r = \infty$.

By providing analytical results for r=0, our study is complementary to (Girardin and Débarre 2021), where the invasion outcome was described numerically in (Girardin and Débarre 2021, Figure 3.A) together with a series of analytical estimates of the sign of the speed.



30 Page 14 of 54 L. Kläy et al.

3.2 Models with partial conversion

In this section, we extend the study to models (4) and (5) with partial conversion. The models are reformulated in terms of allelic densities ($n_{\rm D}$ and $n_{\rm W}$) rather than genotype densities ($n_{\rm DD}$, $n_{\rm DW}$ and $n_{\rm WW}$). This reformulation enables reducing the number of equations from three to two equations. Even if we are not able to determine the genotypic composition of the population (which individual genotype a gamete comes from, either homozygote or heterozygote), the spreading properties are equivalent. Interestingly, the same roadmap as in the full conversion model can be followed. Again, we focus on the two extreme regimes $r = +\infty$ and r = 0.

3.2.1 Conversion occurring in the zygote

When conversion occurs in the zygote, we can deduce the following system from model (4), with $n_D = n_{DD} + \frac{1}{2} n_{DW}$ and $n_W = n_{WW} + \frac{1}{2} n_{DW}$:

$$\begin{cases} \partial_{t} n_{D} - \partial_{xx}^{2} n_{D} = n_{D} \left[\frac{r(1-n)+1}{n} \left[(1-s)(n_{D}+2c n_{W}) + (1-sh)(1-c) n_{W} \right] - 1 \right] = F_{D}^{z}(n_{D}, n_{W}), \\ \partial_{t} n_{W} - \partial_{xx}^{2} n_{W} = n_{W} \left[\frac{r(1-n)+1}{n} \left[n_{W} + (1-sh)(1-c) n_{D} \right] - 1 \right] = F_{W}^{z}(n_{D}, n_{W}). \end{cases}$$

$$(17)$$

The density $n_{\rm W}$ (resp. $n_{\rm D}$) corresponds to one half of the wild-type (resp. drive) allele density at the time of zygote formation. When conversion happens in the zygote, heterozygous individuals are the result of conversion failures and produce one half of each type of gamete, drive or wild-type.

3.2.1.1 Preliminary statements on the model

This model brings more variety in terms of traveling waves than the previous one (6). Cases of monostable wild-type invasion can occur, as well as cases of monostable coexistence. We introduce first all the possible minimal speeds of the problem linearized at low densities and detail later under which parameters they arise.

The minimal speed of the problem linearized at low drive density, i.e. the speed of any pulled monostable wave with a positive speed is given by:

$$2\sqrt{\partial_{n_{\rm D}}F_D^z(0,1)} = 2\sqrt{2c(1-s) + (1-sh)(1-c) - 1} = 2\sqrt{c(1-2s) - sh(1-c)},\tag{18}$$

provided that the quantity is non-negative. The minimal speed of the problem linearized at low wild-type density depends on the stable steady state of a population only bearing drive alleles (i.e. a population of drive homozygotes): $n_{\rm D}^* = n_{\rm DD}^* = \min(0, 1 - \frac{s}{r(1-s)})$ (Girardin and Débarre 2021). If $n_{\rm D}^* = 1 - \frac{s}{r(1-s)}$, this minimal speed is given by:

$$-2\sqrt{\partial_{n_{\mathbf{W}}}F_{\mathbf{W}}^{z}(n_{\mathbf{D}}^{*},0)} = -2\sqrt{\frac{(1-sh)(1-c)}{1-s}} - 1.$$
 (19)



If $n_{\rm D}^* = 0$, the minimal speed of the problem linearized at low wild-type density is given by the classical Fisher-KPP formula:

$$-2\sqrt{\partial_{n_{\mathbf{W}}}F_{W}^{z}(0,0)} = -2\sqrt{r}.$$
 (20)

Note that (20) is the only minimal speed depending on parameter r: it corresponds to the case of gene drive clearance, the only configuration where the drive allele disappears uniformly in space.

For our analysis, it will be convenient to rewrite model (17) so that it follows the frequency of the drive $p_D = \frac{n_D}{n_W + n_D}$ and the total population density $n = n_{WW} + n_{DW} + n_{DD} = n_W + n_D$ (details in B.2.1):

$$\begin{cases} \partial_{t}n - \partial_{xx}^{2}n = \left(r(1-n)+1\right)\left((1-s)p_{D}^{2}+2p_{D}(1-p_{D})\left[c(1-s)+(1-c)(1-sh)\right]+(1-p_{D})^{2}\right)n-n, \\ \partial_{t}p_{D} - \partial_{xx}^{2}p_{D} = 2\partial_{x}\log(n)\partial_{x}p_{D} \\ +\left(r(1-n)+1\right)\left(\left[1-2(1-c)(1-h)\right]sp_{D}-s\left[1-(1-c)(1-h)\right]+c(1-s)\right)(1-p_{D})p_{D}. \end{cases}$$

$$(21)$$

3.2.1.2 $r = +\infty$

Similarly as in Sect. 3.1.2, we compute formally the limiting equation on (21) when $r \to +\infty$:

$$\frac{\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D}}{\left[1 - 2(1 - c)(1 - h)\right] s \ p_{\rm D} - s[1 - (1 - c)(1 - h)] + c(1 - s)\right) (1 - p_{\rm D}) \ p_{\rm D}}{(1 - s) \ p_{\rm D}^2 + 2 \ p_{\rm D} \ (1 - p_{\rm D}) \ [c \ (1 - s) + (1 - c) \ (1 - sh)] + (1 - p_{\rm D})^2}.$$
(22)

Note that as in Sect. 3.1, this equation does not depend on n. We introduce:

$$\mathscr{A}_z := s \left[2(1-c)(1-h) - 1 \right], \quad s_1 := \frac{c}{1 - h(1-c)}, \quad s_{2,z} := \frac{c}{2c + h(1-c)}.$$
(23)

where z stands for zygote. Note that $\mathcal{A}_z > 0 \iff s_1 < s_{2,z}$.

We distinguish between two cases, depending on the sign of \mathcal{A}_z . If $\mathcal{A}_z > 0$, which can only happen if we have both $c < \frac{1}{2}$ and $h < \frac{1}{2}$, the system is always monostable. We observe a drive invasion for $s < s_1$, a coexistence state for $s_1 < s < s_{2,z}$ and a wild-type invasion for $s_{2,z} < s$. Criterion (10) is always verified (see Appendix E.1), consequently every traveling wave (or sub-traveling wave in case of coexistence) is pulled, moving at speed (18) or (19). These statements are summarized in Table 5.

Now we turn to the case $\mathcal{A}_z < 0$. The system is monostable for $s < s_{2,z}$ (drive invasion), bistable for $s_{2,z} < s < s_1$ and monostable for $s_1 < s$ (wild-type invasion). In case of monostable drive invasion, we define a set \mathcal{S}_z of s values:

$$\mathcal{S}_z := \Big\{ s \in (0,1) | \Big(1 - 2s[1 - (1-h)(1-c)] \Big) \Big((-2c - h + ch)s + c \Big) + s \left[2(1-c)(1-h) - 1 \right] > 0 \Big\}. \tag{24}$$



30 Page 16 of 54 L. Kläy et al.

Table 5 Traveling waves study for Model (22) (limit of system (21) when $r = +\infty$) when $\mathcal{A}_z > 0$, with $v_{lin+} = 2\sqrt{c(1-2s)-sh(1-c)}$ (18) and $v_{lin-} = -2\sqrt{\frac{(1-sh)(1-c)}{1-s}-1}$ (19). For all values of s, there exists only one stable state (monostability). In particular in the case of coexistence, the stable state (in the center) invades two unstable states (on the right and left)

s value $s < s_1$		$s_1 < s < s_{2,z}$	$s_{2,z} < s$	
Stability	Monostable			
Speed	$v = v_{lin+}$	$v = v_{lin+}$ and $v = v_{lin-}$	$v = v_{lin-}$	
Wave		Pulled Wave		
	Drive invasion	Coexistence	Wild-type invasion	
Invasion	Appace Space	Acuanda yaya Space	Duve (redreuch	

Table 6 Traveling waves study for Model (22) (limit of system (21) when $r = +\infty$) when $\mathcal{A}_z < 0$, with $v_{lin+} = 2\sqrt{c(1-2s)-sh(1-c)}$ (18)

s value	$s\in \mathcal{S}_z$	$s \in (0, s_{2,z}) \backslash \mathscr{S}_z$	$s_{2,z} < s < s_1$	$s_1 < s < 1$
Stability	Monostable		Bistable	Monostable
Speed	$v = v_{lin+}$	$v \ge v_{lin+}$		v < 0
Wave	Pulled wave		Pushed wave	
Invasion	Drive invasion		Drive or Wild-type invasion	Wild-type invasion

For all s in \mathcal{S}_z and $\mathcal{A}_z < 0$, criterion (10) is verified and consequently, there exists a pulled monostable traveling wave with positive speed (see Appendix E.1). Note that such s values are necessarily strictly below $s_{2,z}$, condition for a monostable drive invasion. In case of wild-type invasion, criterion (10) is never verified (see Appendix E.1). These statements are summarized in Table 6.

3.2.1.3 r = 0

Using the relation $n = n_{WW} + n_{DW} + n_{DD} = n_W + n_D$, system (17) can be rewritten as follows when r = 0:



s value	s value $0 < s < s_{2,z}$ $s_{2,z}$		
Stability	Monostable	Degenerate case	
Speed	$v = v_{lin+}$	No wave	
Wave	Pulled wave		
	Drive invasion	Gene drive clearance	
Invasion	Space Space	Space	

Table 7 Traveling waves study for Model (25) (limit of system (21) when r = 0), with $v_{lin+} = 2\sqrt{c(1-2s)-sh(1-c)}$ (18)

$$\begin{cases} \partial_{t} n_{D} - \partial_{xx}^{2} n_{D} = \left(c (1 - s) + s (1 - c) (1 - h) \right) \frac{n_{D} n_{W}}{n_{D} + n_{W}} - s n_{D}, \\ \partial_{t} n_{W} - \partial_{xx}^{2} n_{W} = -\left(1 - (1 - sh) (1 - c) \right) \frac{n_{D} n_{W}}{n_{D} + n_{W}}. \end{cases}$$
(25)

We apply the results of Appendix D with $\beta_1 = 1 - (1 - sh)(1 - c)$ and $\beta_2 = c(1 - s) + s(1 - c)(1 - h)$. There exists a monostable and pulled drive invasion wave if:

$$\beta_2 > \gamma \iff s < s_{2,z} = \frac{c}{2c + h(1 - c)}.$$
 (26)

On the other hand when $\beta_2 < \gamma$, the reaction term of n_D in (25) is strictly negative. As before, the density n_D converges to zero uniformly in space at rate at least $\beta_2 - \gamma$ (gene drive clearance) and the final density of wild-type is not clearly determined: the problem boils down to diffusion only in the large time asymptotics (details in Appendix C.2.1). These statements are summarized in Table 7.

Note that, when $\mathcal{A}_z > 0$, a condition for having a pulled wave with positive speed for both r = 0 and $r = +\infty$ is $s < s_{2,z}$. When $\mathcal{A}_z < 0$, a condition for having a pulled wave with positive speed for both r = 0 and $r = +\infty$ is $s \in \mathcal{S}_z \subseteq [0, s_{2,z}]$. This suggests that, under those conditions, whatever the value of the demographic parameter r is, the drive invasion wave is always pulled and consequently, travels at a speed which does not depend on r either (speed given by (18)). We verify this intuition numerically (vertical level lines) in the following section.

3.2.1.4 Numerical illustrations

$$\mathcal{A}_z > 0$$

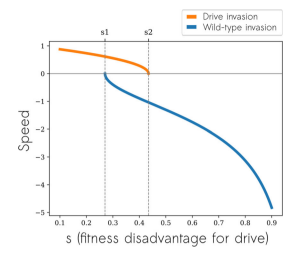
In a first example we choose c=0.25 and h=0.1 such that $\mathscr{A}_z>0$. The s threshold values are $s_1\approx 0.27$ and $s_{2,z}\approx 0.43$. As discussed in the previous sections, when $s< s_{2,z}$, all waves are pulled (sub-)traveling waves for $r=+\infty$ and r=0. Note that $s>s_1$ is the condition for the existence of pulled (sub-)traveling waves with negative speed only when $r=+\infty$.

We show the value of the speed (18) and (19) of the pulled waves as a function of s when $r = +\infty$, with c = 0.25 and h = 0.1 (in Fig. 2).



30 Page 18 of 54 L. Kläy et al.

Fig. 2 Speed of the drive invasion (18) in orange and speed of the wild-type invasion (19) in blue, as a function of s when $r = +\infty$, with c = 0.25 and h = 0.1 (color figure online)



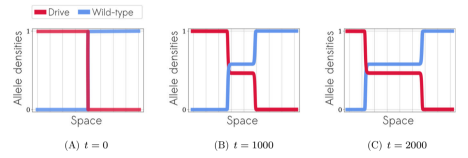


Fig. 3 Allele densities as a function of space, at different times, for s = 0.35, c = 0.25, h = 0.1 and r = 3

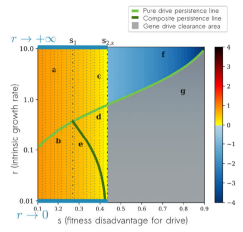
Note that for $0.27 \approx s_1 < s < s_{2,z} \approx 0.43$, we have both a drive and a wild-type invasion, leading to a stable coexistence state. This case is illustrated in Fig. 3 with s = 0.35.

We now compute numerically the values of the wave speed for intermediate values of r (in Fig. 4). In case of coexistence, for $s_1 < s < s_{2,z}$, we choose to show only the positive speed value.

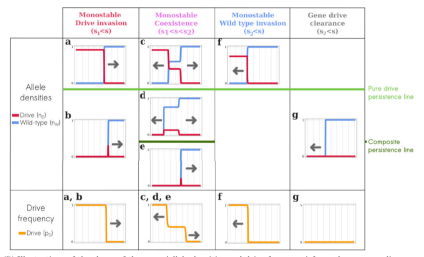
For a better understanding of Fig. 4, we detail the effect of fitness disadvantage s and dominance coefficient h on drive dynamics for $r = +\infty$ and c = 0.25, without spatial structure, in Appendix F.1 (in Fig. 13A).

In Fig. 4, the speed value for $s < s_{2,z}$ seems not to depend on the demographic parameter r: whatever the final equilibrium is, going from population extinction to full replacement of the wild-type genotypes by drive genotypes, the invasion occurs at the same speed. This is in agreement with the fact that the values of the speed coincide when $r = +\infty$ and r = 0 for $s < s_{2,z}$. If correct, the value of the speed can be found in Fig. 2.





(A) Heatmap representing the speed of the wave for c = 0.25, h = 0.1 when conversion occurs in the zygote.

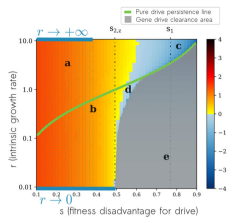


(B) Illustrations of the shape of the wave (allele densities and drive frequency) for each corresponding case in the heatmap.

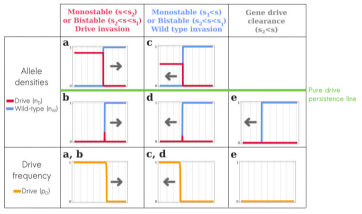
Fig. 4 A Heatmap representing the speed of the waves for c = 0.25, h = 0.1 when conversion occurs in the zygote When the drive invades the population, the speed is positive (in yellow-orange). On the contrary, when the wild-type invades the population, the speed is negative (in blue). When both drive and wild types invade (coexistence), only the speed of the drive is shown in the heatmap, resulting in an apparent discontinuity at $s = s_{2,z}$. As $\mathcal{A}_z > 0$, the system is always monostable for $r = +\infty$: when $s < s_1$ the drive always invades; when $s_1 < s < s_{2,z}$ the final state is a coexistence state; when $s > s_{2,z}$ the wild-type invades or there is gene drive clearance. The turquoise horizontal lines at the bottom and at the top of the heatmap indicate the theoretical values of s such that there exists a pulled wave with positive speed, respectively for $r = +\infty$ and r = 0. Below the pure drive persistence line (light green), a well-mixed population containing only drive homozygous individuals will necessarily go extinct. Below the composite persistence line (dark green), it is the whole population that gets extinct (calculations for both lines available in Appendix F.2). The gray zone corresponds to the gene drive clearance area. Outside the gray zone, the level lines are apparently vertical, meaning that the wave speed would be independent of r. This is in agreement with the fact that the values of the speed coincide when $r = +\infty$ and r = 0 for $s < s_{2,z}$. If correct, the value of the speed can be found in Fig. 2. B Shape of the wave for each case indicated by a letter in the heatmap above. The position of the graphs in the table reflects the position in the heatmap with respect to the persistence lines (color figure online)



30 Page 20 of 54 L. Kläy et al.



(A) Heatmap representing the speed of the wave for $c=0.75,\,h=0.1$ when conversion occurs in the zygote.



(B) Illustrations of the shape of the wave (allele densities and drive frequency) for each corresponding case in the heatmap.

Fig. 5 A Heatmap representing the speed of the wave for c=0.75, h=0.1 when conversion occurs in the zygote When the drive invades the population, the speed is positive (in yellow-orange-red). On the contrary, when the wild-type invades the population, the speed is negative (in blue). We have $\mathscr{A}_z < 0$, therefore when $r=+\infty$: when $s< s_{2,z}$ the system is monostable and the drive always invades; when $s_{2,z} < s < s_1$ the system is bistable and the final state depends on the initial condition; when $s>s_1$ the system is monostable and the wild-type invades or there is gene drive clearance. The turquoise horizontal lines at the bottom and at the top of the heatmap indicate the theoretical values of s such that there exists a pulled wave with positive speed, respectively for $r=+\infty$ and r=0. Below the pure drive persistence line (light green), a well-mixed population containing only drive homozygous individuals will necessarily go extinct (calculations for this line available in Appendix F.2). For $s\in \mathscr{S}_z$, i.e. s<0.38, the level lines are apparently vertical: this is in agreement with the fact that the values of the speed coincide when $r=+\infty$ and r=0 in this area. B Shape of the wave for each case indicated by a letter in the heatmap above. The position of the graphs in the table reflects the position in the heatmap with respect to the pure drive persistence line (color figure online)



$$\mathcal{A}_z < 0$$

In a second example, we choose c=0.75 and h=0.1 such that $\mathcal{A}_z < 0$. The s threshold values are $s_1 \approx 0.77$ and $s_{2,z} \approx 0.49$. As discussed in the previous section, when $s \in \mathcal{S}_z$ (in our case $s \lesssim 0.38$), all waves are pulled traveling waves. However, the latter criterion is not a sufficient condition. It is expected that waves are indeed pulled beyond this approximate value of 0.38. However, this would require to use numerical continuation methods as in Sect. 3.1.2. We computed numerically the speed values for intermediate values of r, as shown in Fig. 5. We believe that the wave speed is independent of the demographic parameter r when the wave is pulled (visual observation for $s \lesssim 0.38$).

For a better understanding of Fig. 5, we detail the effect of fitness disadvantage s and dominance coefficient h on drive dynamics for $r = +\infty$ and c = 0.75, without spatial structure, in Appendix F.1 (in Fig. 13B).

3.2.2 Conversion occurring in the germline

When conversion occurs in the germline, we can deduce the following system from model (5), with $n_D = n_{DD} + (1+c) \frac{1}{2} n_{DW}$ and $n_W = n_{WW} + (1-c) \frac{1}{2} n_{DW}$:

$$\begin{cases} \partial_{t} n_{D} - \partial_{xx}^{2} n_{D} = n_{D} \left[\frac{r(1-n)+1}{n} \left[(1-s)n_{D} + (1-sh)(1+c)n_{W} \right] - 1 \right] = F_{D}^{g}(n_{D}, n_{W}), \\ \partial_{t} n_{W} - \partial_{xx}^{2} n_{W} = n_{W} \left[\frac{r(1-n)+1}{n} \left[n_{W} + (1-sh)(1-c)n_{D} \right] - 1 \right] = F_{W}^{g}(n_{D}, n_{W}). \end{cases}$$
(27)

The density $n_{\rm W}$ (resp. $n_{\rm D}$) corresponds to one half of the wild-type (resp. drive) allele density at the time of zygote formation. When conversion happens in the germline, heterozygous individuals undergo a conversion of their wild-type alleles with probability c, and produce a fraction (1+c)/2 of drive-carrying gametes.

3.2.2.1 Preliminary statements on the model

As before, we detail the minimal speed of the problem linearized at low densities, for both drive and wild-type alleles.

In case of drive invasion, the minimal speed of the problem linearized at low drive density, i.e. the speed of any pulled monostable wave with positive speed is given by:

$$2\sqrt{\partial_{n_D} F_D^g(0,1)} = 2\sqrt{(1-sh)(1+c)-1}. (28)$$

Note that $F_W^g(n_D, n_W) = F_W^z(n_D, n_W)$: in case of a wild-type invasion, the minimal speeds are already given by (19) and (20) (Sect. 3.2.1).

For our analysis, it will be convenient to rewrite model (27) so that it follows the frequency of the drive $p_D = \frac{n_D}{n_W + n_D}$ and the total population density $n = n_{WW} + n_{DW} + n_{DD} = n_W + n_D$ (details in Appendix B.2.2):

$$\begin{cases} \partial_{t} n - \partial_{xx}^{2} n &= \left(r (1 - n) + 1 \right) \left((1 - s) \ p_{D}^{2} + 2 (1 - sh) \ p_{D} (1 - p_{D}) + (1 - p_{D})^{2} \right) n - n, \\ \partial_{t} p_{D} - \partial_{xx}^{2} p_{D} &= 2 \ \partial_{x} \log(n) \ \partial_{x} p_{D} + \left(r (1 - n) + 1 \right) \left((2h - 1) \ s \ p_{D} + (1 - sh)(1 + c) - 1 \right) \ p_{D} (1 - p_{D}). \end{cases}$$
(29)



30 Page 22 of 54 L. Kläy et al.

$3.2.2.2 r = +\infty$

3.1.2, we can compute formally the limiting equation on p_D when $r = +\infty$:

$$\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D} = \frac{\left(-(1-2h) s p_{\rm D} + [(1-sh)(1+c)-1]\right) p_{\rm D} (1-p_{\rm D})}{(1-s) p_{\rm D}^2 + 2 (1-sh) p_{\rm D} (1-p_{\rm D}) + (1-p_{\rm D})^2}. (30)$$

Note that as in Sect. 3.1, this equation does not depend on n. We introduce:

$$\mathcal{A}_g := s \ (1-2h), \qquad s_1 := \frac{c}{1-h(1-c)}, \qquad s_{2,g} := \frac{c}{2ch+h(1-c)} = \frac{c}{h(1+c)}.$$
(31)

where g stands for germline. Note that $\mathcal{A}_g > 0 \iff s_1 < s_{2,g}$. We define a set \mathcal{S}_g of s values:

$$\mathcal{S}_g := \left\{ s \in (0,1) | (1 - 2sh) \left(c - sh(c+1) \right) + s(1 - 2h) > 0 \right\}. \tag{32}$$

Results are exactly the same as in Sect. 3.2.1.2, substituting \mathscr{A}_z by \mathscr{A}_g , $s_{2,z}$ by $s_{2,g}$, \mathscr{S}_z by \mathscr{S}_g , and the minimal speed of the problem linearized at low drive density (18) by (28) (see Appendix E.2).

3.2.2.3 r = 0

Using the relation $n = n_W + n_D$, system (27) can be rewritten as follows when r = 0:

$$\begin{cases} \partial_{t} n_{D} - \partial_{xx}^{2} n_{D} = \left(c (1 - sh) + s (1 - h) \right) \frac{n_{D} n_{W}}{n_{D} + n_{W}} - s n_{D}, \\ \partial_{t} n_{W} - \partial_{xx}^{2} n_{W} = -\left(1 - (1 - sh) (1 - c) \right) \frac{n_{D} n_{W}}{n_{D} + n_{W}}. \end{cases}$$
(33)

We apply the results of Appendix D with $\beta_1 = 1 - (1 - sh)(1 - c)$ and $\beta_2 = c(1 - sh) + s(1 - h)$. There exists a monostable and pulled drive invasion wave if:

$$\beta_2 > \gamma \iff s < s_{2,g} = \frac{c}{h(1+c)} = \frac{c}{2ch + h(1-c)}.$$
 (34)

On the other hand when $\beta_2 < \gamma$, the reaction term of n_D in (33) is strictly negative. As before, the density n_D converges to zero uniformly in space at rate at least $\beta_2 - \gamma$ (gene drive clearance) and the final density of wild-type is not clearly determined: the problem boils down to diffusion only in the large time asymptotics (details in Appendix C.2.1).

Note that the same intuitions as in Sect. 3.2.1.3 hold: when $\mathcal{A}_g > 0$, a condition for having a pulled wave with positive speed for both r = 0 and $r = +\infty$ is $s < s_{2,g}$; when $\mathcal{A}_g < 0$, a condition for having a pulled wave with positive speed for both r = 0 and $r = +\infty$ is $s \in \mathcal{S}_g \subseteq [0, s_{2,g}]$. This suggests that, under those conditions, whatever the value of the demographic parameter r is, the drive invasion wave is always pulled and consequently, travels at a speed which does not depend on r either (speed given



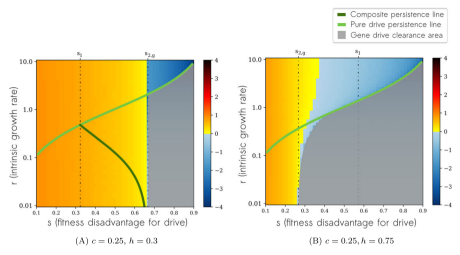


Fig. 6 Heatmap representing the speed of the wave when conversion occurs in the germline. When the drive invades the population, the speed is positive (in yellow-orange-red). On the contrary, when the wild-type invades the population, the speed is negative (in blue). Below the pure drive persistence line (light green), a well-mixed population containing only drive homozygous individuals will necessarily go extinct. Below the composite persistence line (dark green), it is the whole population that gets extinct (calculations for both lines available in Appendix F.2). The gray zone corresponds to the gene drive clearance area. In (a) we have $\mathcal{A}_g > 0$, therefore the system is always monostable for $r = +\infty$: when $s < s_1$ the drive always invades; when $s_1 < s < s_{2,g}$ the final state is a coexistence state; when $s > s_{2,g}$ the wild-type invades or there is gene drive clearance. When both drive and wild types invade (coexistence), only the speed of the drive is shown in the heatmap, resulting in an apparent discontinuity at $s = s_{2,z}$. In (b) we have $\mathcal{A}_z < 0$, therefore when $r = +\infty$: when $s < s_{2,g}$ the system is monostable and the drive always invades; when $s_2 < s_3$ the system is bistable and the final state depends on the initial condition; when $s > s_1$ the system is monostable and the wild-type invades or there is gene drive clearance (color figure online)

by (28)). As before, we verify this intuition numerically (vertical level lines) in the following section.

3.2.2.4 Numerical illustrations

For a better understanding of Fig. 6A and B, we detail the effect of fitness disadvantage s and dominance coefficient h on drive dynamics for $r = +\infty$ and c = 0.25, without spatial structure, in Appendix F.1 (in Fig. 14A).

3.2.3 Conclusion

When conversion occurs in the zygote (resp. in the germline) for $\mathcal{A}_z < 0$ (resp. $\mathcal{A}_g < 0$), demographics influence the speed of the drive propagation at least for $s \in (s_{2,z},s_1)$ (resp. $s \in (s_{2,g},s_1)$). More precisely, the sign of the speed can switch, changing the type of the invasion (drive or wild-type). When $\mathcal{A}_z > 0$ (resp. $\mathcal{A}_g > 0$) however, a model following only frequencies will always predict the correct speed of expansion. However, a model following only frequencies will not provide information on population size, and in particular whether the population is suppressed or eradicated, while this point is of great biological relevance.



30 Page 24 of 54 L. Kläy et al.

For both zygote and germline conversion timings, the critical values of \mathscr{A}_z and \mathscr{A}_g can be interpreted as the values at which the fitness of adults who were born heterozygous (f'_H) is the arithmetic mean of the fitness of adults born homozygote $((f_D + f_W)/2)$. The fitness of adults who were born heterozygous depends on the timing of gene conversion. For germline conversion, $f'_H = f_H$, and $\mathscr{A}_g = 0$ when h = 1/2, i.e. when there is co-dominance between the drive and wild-type alleles, i.e. when $f'_H = f_H = (f_D + f_W)/2$. For zygote conversion, the fitness of adults born heterozygous depends on whether gene conversion has taken place or not $(f'_H = (1-c)(1-hs)+c(1-s))$. The condition $\mathscr{A}_z = 0$ is equivalent to (1-c)(1-h) = 1/2, which happens for $f'_H = (f_D + f_W)/2$.

4 Discussion

Following (Girardin and Débarre 2021), we quantified the impact of demography in the case of the propagation of a super-Mendelian drive. We extended the analysis of reference (Girardin and Débarre 2021) to the case of partial conversion (0 < c < 1), implying the presence of heterozygotes.

On the final state of the population

The final size of the population naturally varies. In case where no wild type can survive, the final size is the same regardless of the details of gene conversion(timing nor probability): $n_{\rm D}^* = \min(0, 1 - \frac{s}{r(1-s)})$. In case of coexistence between wild-type and drive alleles, the final size depends on all parameters (see Appendix F.2.2). Interestingly, in the case of coexistence, the drive allele can persist in the population even if a pure drive population would not $(n_{\rm D}^* = 0)$, see Fig. 4 (note the area between the *composite persistence line* and the *pure drive persistence line*). In contrast with standard Mendelian genetics (corresponding to c = 0), coexistence can occur even if the dominance parameter is such that $h \in (0, 1)$ (Deredec et al. 2008; Rode et al. 2019). More precisely, when conversion is partial and, either $h < 1 - \frac{1}{2(1-c)}$ (zygote conversion), or $h < \frac{1}{2}$ (germline conversion), there exists a stable coexistence state if s takes intermediate values $s \in (s_1, s_{2,z})$ (zygote conversion), or $s \in (s_1, s_{2,g})$ (germline conversion), where $s_1, s_{2,z}, s_{2,g}$ depend on (c, h) but do not depend on the demographic parameter r (see details in Sect. 3.2.1 and 3.2.2). While the final size of the population naturally depends on r.

On the transient regime (propagation of waves)

In order to evaluate the impact of demography on the dynamics of drive expansion, we compared the extreme cases $r \to \infty$ and $r \to 0$ (resp. low demographic variations versus large demographic variations).

For r = 0, we found that, when the drive propagates, it does so through a monostable and pulled wave. This happens when the drive is not too costly. In contrary, the drive gets uniformly extinct if it is too costly. The threshold on the fitness cost $s_{2,z}$



(zygote conversion), or $s_{2,g}$ (germline conversion), depends on (c, h). The situation is analogous to the spatial spreading of an epidemic following a SI type model.

The case r=0 gives the possibility to measure the importance of the demographic advection term $2\partial_x \log(n)\partial_x p$ when the problem is formulated in frequency, see equations (12), (21), (29). In fact, we show that ignoring this term can lead to an overestimation of the wave speed. This happens, for instance, in case of perfect conversion in the zygote, when $s \in (\frac{2}{5}, \frac{1}{2})$, then the equation (12) without $2\partial_x \log(n)\partial_x p$ would lead to a pushed front with velocity $\frac{2-3s}{\sqrt{2s}}$ (Hadeler and Rothe 1975). However, we proved that the front is actually pulled with velocity $2\sqrt{1-2s} < \frac{2-3s}{\sqrt{2s}}$. Intuitively, advection due to demographic variations slows down the expansion of the bulk. Noticeably, the effect is so strong that it prevents the front from being pushed.

In contrast, for $r = \infty$, the analysis boils down to a single equation on the drive allele frequency (Strugarek and Vauchelet 2016; Girardin and Débarre 2021). According to (Rode et al. 2019), where the case of germline conversion was investigated, there is a large panel of frequency-dependence relationships, including monostable fixation of one allele, bi-stability, and stable coexistence between the two alleles, even if $h \in (0, 1)$. This leads to a variety of propagation phenomena, either pulled or pushed, as described in Sect. 3.2.2. The same panel of relationships arises in the case of zygote conversion, with qualitative similarities but quantitative differences in the thresholds and in the wave speeds, compare Sect. 3.2.2 with Sect. 3.2.1.

To connect r=0 and $r=\infty$, we conjecture that, if the wave of the drive is pulled at $r=\infty$, then it is pulled for any value of r>0, and the wave speed is independent of r. In particular, this occurs when the frequency-dependence term induces monostable dynamics and s is small enough, or when there is stable co-existence. This conjecture is supported by numerical investigations (Figs. 4, 5 and 6A, B). Still, the final size of population naturally depends on r.

Perspectives

We have focused on the classical dichotomy between pulled and pushed waves, even if the transition between the two is subject to current research both in theoretical studies (An et al. 2021; Avery et al. 2022; Birzu et al. 2018), and in experimental works (Dahirel et al. 2021).

Pulled and pushed waves are associated with different outcomes on the maintenance of neutral diversity (which was not considered in our study). The genetic diversity of a population expanding by a pulled wave is very limited (with possible accumulation of deleterious mutations (Peischl et al. 2013)), while more diversity is maintained under a pushed wave (Roques et al. 2012). It could be interesting to investigate how gene conversion influences the maintenance of diversity along an expanding wave. More generally, the bottleneck following spread of a suppression drive will affect diversity, which may have long-lasting consequences even if wild-type individuals later recolonize the area.

It would be highly relevant to explore stochastic dynamics beyond our deterministic approach. When population sizes get to be small, as in the drive eradication case, large fluctuations and even chasing events are expected, as described in Champer (2021).



30 Page 26 of 54 L. Kläy et al.

It would also be extremely interesting to extend the scope of the model, including by distinguishing between males and females which may have different fitnesses (especially in transgenic mosquitoes (Beaghton et al. 2016; North et al. 2020; Kyrou et al. 2018; Hammond et al. 2015)). Plural life stages or haploid phases might also influence modeling conclusions (Liu and Champer 2022; Li et al. 2020).

Acknowledgements This work is funded by ANR-19-CE45-0009-01 "TheoGeneDrive". This project has received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programm (grant agreement No 865711). We are grateful to the INRAE MIGALE bioinformatics facility (MIGALE, INRAE, 2020. Migale bioinformatics Facility, DOI: 10.15454/1.5572390655343293E12) for providing computing resources.

Declarations

Conflict of interest The authors declare that they have no conflict of interest.

Appendix

A Model with partial conversion: growth term details

To obtain the global growth term for each genotypes in models (4) and (5), we calculate type proportions among the offspring for each possible couple, and then we sum the corresponding terms. The calculations follow standard lines of population genetics, differing only by the timing of gene conversion. When conversion occurs in the zygote, the parameter c appears in between the gametes and the offspring production, whereas when conversion occurs in the germline, it appears before gametes production. These equations in densities are consistent with the one obtained in frequency in the literature (when conversion occurs in the zygote (Deredec et al. 2008; Unckless et al. 2015), or in the germline (Deredec et al. 2008; Rode et al. 2019)).



A.1 Conversion occurring in the zygote

 Table 8 Growth term details when conversion occurs in the zygote

Parents	Egg	Adult	Fitness	Growth term
WW + WW	— 1 → WW —— 1 -	→ WW	1	$\frac{n_{\text{ww}}n_{\text{ww}}}{n}$
	, , , , , , , , , , , , , , , , , , ,	→ DD	1-s	$\frac{1}{2} c (1-s) \frac{2n_{\text{WW}} n_{\text{DW}}}{n}$
	$\begin{array}{c} \text{WD} \stackrel{c}{\underset{1-c}{\overset{c}{\longrightarrow}}} \\ \stackrel{1}{\underset{2}{\overset{c}{\longrightarrow}}} \\ \text{WW} \longrightarrow 1-c \end{array}$	\rightarrow WD	1-sh	$\frac{1}{2} (1-c) (1-sh) \frac{2n_{\text{ww}} n_{\text{DW}}}{n}$
WW + WD	1/2 → WW — 1 -	→ WW	1	$\frac{1}{2} \frac{2n_{\text{WW}}n_{\text{DW}}}{n}$
	1 > WD — c-	→ DD	1-s	$c (1-s) \frac{2n_{\text{WW}} n_{\text{DD}}}{n}$
WW + DD	$-1 \to WD \xrightarrow{c} DD$ $1 - c \to WD$	1-sh	$(1-c) (1-sh) \frac{2n_{\text{WW}}n_{\text{DD}}}{n}$	
	WW 1 -	→ WW	1	$\frac{1}{4} \frac{n_{\mathrm{DW}} n_{\mathrm{DW}}}{n}$
	$ \begin{array}{c} \frac{1}{4} \\ \frac{1}{2} \rightarrow \text{WD} \\ \frac{1}{1} - c \end{array} $	→ DD	1-s	$\frac{1}{2} c \left(1 - s\right) \frac{n_{\text{DW}} n_{\text{DW}}}{n}$
WD + WD	$\frac{1}{2} \longrightarrow WD \longrightarrow 1 - c$	\rightarrow WD	1-sh	$\frac{1}{2} (1-c) (1-sh) \frac{n_{\rm DW} n_{\rm DW}}{n}$
	DD 1 -	→ DD	1-s	$\frac{1}{4} (1-s) \frac{n_{\rm DW} n_{\rm DW}}{n}$
	WD c -	→ DD	1-s	$\frac{1}{2} c (1-s) \frac{2n_{\rm DW} n_{\rm DD}}{n}$
	WD $\stackrel{c}{\underset{1-c}{\overset{1}{\sim}}}$ $\stackrel{\text{WD}}{\underset{1-c}{\overset{c}{\sim}}}$ $\stackrel{1}{\underset{2}{\sim}}$ DD $\stackrel{1-c}{\underset{1-c}{\sim}}$	\rightarrow WD	1-sh	$\frac{1}{2} (1-c) (1-sh) \frac{2n_{\rm DW}n_{\rm DD}}{n}$
WD + DD	$\frac{1}{2}$ DD — 1 -	→ DD	1-s	$\frac{1}{2} \left(1 - s\right) \frac{2n_{\rm DW} n_{\rm DD}}{n}$
$\mathrm{DD} + \mathrm{DD}$	— 1 → DD —— 1 -	→ DD	1-s	$(1-s) \; \frac{n_{\rm DD} n_{\rm DD}}{n}$



30 Page 28 of 54 L. Kläy et al.

A.2 Conversion occurring in the germline

Table 9 Growth term details when conversion occurs in the germline

Parents	Gametes Adult	Fitness	Growth term
WW + WW	$1 \longrightarrow W,W+W,W-1 \longrightarrow WW$	1	$\frac{n_{\text{ww}}n_{\text{ww}}}{n}$
WW + WD	$(V,W + D,D - 1 \rightarrow WD)$ $(1-c)$ $W,W + W,D - \frac{1}{2} \rightarrow WW$ $\frac{1}{2} \rightarrow WD$	1 - sh 1 $1 - sh$	$\begin{array}{c} c \ (1-sh) \ \frac{2n_{_{\rm WW}}n_{_{\rm DW}}}{n} \\ \\ (1-c) \ \frac{1}{2} \ \ \frac{2n_{_{\rm WW}}n_{_{\rm DW}}}{n} \\ \\ (1-c) \ \frac{1}{2} \ \ (1-sh) \ \frac{2n_{_{\rm WW}}n_{_{\rm DW}}}{n} \end{array}$
WW + DD	$\longrightarrow 1 \longrightarrow W,W+D,D\longrightarrow 1 \longrightarrow WD$	1 - sh	$(1-sh) \; \frac{2n_{\scriptscriptstyle \rm WW} n_{\scriptscriptstyle \rm DD}}{n}$
WD + WD	$D,D + D,D \longrightarrow 1 \longrightarrow DD$ c^{2} $(1-c)^{2} \longrightarrow D,D + W,D \longrightarrow \frac{1}{2} \longrightarrow WD$ $W,D + W,D \longrightarrow \frac{1}{4} \longrightarrow WD$ WW		$c (1-c) (1-sh) \frac{n_{\text{DW}} n_{\text{DW}}}{n}$ $(1-c)^2 \frac{1}{4} (1-s) \frac{n_{\text{DW}} n_{\text{DW}}}{n}$
WD + DD	$ \begin{array}{c} c \\ 1-c \\ W,D+D,D \\ \hline \end{array} $ $ \begin{array}{c} 1 \\ 2 \\ DD \end{array} $ $ \begin{array}{c} WD \\ \frac{1}{2} \\ DD $	1 - s $1 - sh$ $1 - s$	$\begin{array}{c} c & (1-s) \; \frac{2n_{\rm DW}n_{\rm DD}}{n} \\ \\ (1-c) \; \frac{1}{2} \; (1-sh) \; \frac{2n_{\rm DW}n_{\rm DD}}{n} \\ \\ (1-c) \; \frac{1}{2} \; (1-s) \; \frac{2n_{\rm DW}n_{\rm DD}}{n} \end{array}$
DD + DD	$ \longrightarrow 1 \longrightarrow D,D + D,D \longrightarrow 1 \longrightarrow DD $	1-s	$(1-s) \; \frac{n_{\text{\tiny DD}} n_{\text{\tiny DD}}}{n}$



B System rewritten with variables (n, p_D)

Below, we present the details of the reformulation from models (6), (17) and (27) in terms of total population density n and drive allele frequency p_D .

B.1 Model with perfect conversion

We rewrite model (6) with variables:

$$n = n_{WW} + n_{DD},$$
 $p_{D} = \frac{n_{D}}{n_{W} + n_{D}} = \frac{n_{DD}}{n_{WW} + n_{DD}}.$ (35)

where n is the total population density and p_D is the drive allele frequency, or equivalently in this model, the frequency of drive homogeneous individuals.

Equation on n:

$$\partial_{t} n - \partial_{xx}^{2} n = \left(r (1 - n) + 1\right) \left((1 - s) \frac{n_{\text{DD}}^{2} + 2 n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}} + \frac{n_{\text{WW}}^{2}}{n_{\text{WW}} + n_{\text{DD}}}\right) - n,$$

$$= \left(r (1 - n) + 1\right) \left((1 - s) p_{\text{D}}^{2} + (1 - s) 2 p_{\text{D}} (1 - p_{\text{D}})\right)$$

$$+ (1 - s)(1 - p_{\text{D}})^{2} + s(1 - p_{\text{D}})^{2} n - n,$$

$$= \left(r (1 - n) + 1\right) \left((1 - s)(p_{\text{D}} + 1 - p_{\text{D}})^{2} + s(1 - p_{\text{D}})^{2}\right) n - n,$$

$$= \left(r (1 - n) + 1\right) \left((1 - s) + s(1 - p_{\text{D}})^{2}\right) n - n,$$

$$= \left(r (1 - n) + 1\right) \left((1 - s) p_{\text{D}} (2 - p_{\text{D}}) + (1 - p_{\text{D}})^{2}\right) n - n.$$
(36)

Equation on n_{DD}

$$\partial_{t} n_{\text{DD}} = \partial_{xx}^{2} n_{\text{DD}} + (1 - s) \left(r \left(1 - n_{\text{DD}} - n_{\text{WW}} \right) + 1 \right) \frac{n_{\text{DD}}^{2} + 2 n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}} - n_{\text{DD}},$$

$$\stackrel{1}{=} p_{\text{D}} \partial_{xx}^{2} n + 2 \partial_{x} n \partial_{x} p_{\text{D}} + n \partial_{xx}^{2} p_{\text{D}} + (1 - s) \left(r \left(1 - n \right) + 1 \right)$$

$$(2 - p_{\text{D}}) n p_{\text{D}} - n p_{\text{D}}.$$
(37)

Equation on $p_D = \frac{n_{DD}}{n}$:

$$\begin{split} \partial_{t} p_{\mathrm{D}} &= \frac{n \; (\partial_{t} n_{\mathrm{DD}}) - n_{\mathrm{DD}} \; (\partial_{t} n)}{n^{2}} = \frac{n \; (\partial_{t} n_{\mathrm{DD}}) - n \; p_{\mathrm{D}} \; (\partial_{t} n)}{n^{2}} = \frac{1}{n} \Big(\; \partial_{t} n_{\mathrm{DD}} - p_{\mathrm{D}} \; (\partial_{t} n) \Big), \\ &= \frac{1}{n} \Big[\; p_{\mathrm{D}} \; \partial_{xx}^{2} n + 2 \; \partial_{x} n \; \partial_{x} p_{\mathrm{D}} + n \; \partial_{xx}^{2} p_{\mathrm{D}} \; + \; \Big(r \; (1 - n) + 1 \Big) \; (1 - s) \\ &\qquad (2 - p_{\mathrm{D}}) \; n \; p_{\mathrm{D}} - n \; p_{\mathrm{D}} \; \Big] \end{split}$$

 $[\]overline{ \frac{1}{\partial_{xx} :^2 n_{\text{DD}} = \partial_{xx}^2 n p_{\text{D}} = \partial_x (p_{\text{D}} \partial_x n + n \partial_x p_{\text{D}}) } = p_{\text{D}} \partial_{xx}^2 n + 2 \partial_x p_{\text{D}} \partial_x n + n \partial_{xx}^2 p_{\text{D}}$



30 Page 30 of 54 L. Kläy et al.

$$-\frac{1}{n} \left[p_{D} \partial_{xx}^{2} n + \left(r (1-n) + 1 \right) \left((1-s) p_{D} (2-p_{D}) + (1-p_{D})^{2} \right) \right]$$

$$n p_{D} - n p_{D} ,$$

$$= \partial_{xx}^{2} p_{D} + 2 \partial_{x} \log(n) \partial_{x} p_{D} + \left(r (1-n) + 1 \right) p_{D} \left((1-s)(2-p_{D}) - (1-s)(2-p_{D})p_{D} - (1-p_{D})^{2} \right),$$

$$= \partial_{xx}^{2} p_{D} + 2 \partial_{x} \log(n) \partial_{x} p_{D} + \left(r (1-n) + 1 \right) s p_{D} (1-p_{D}) \left(p_{D} - \frac{2s-1}{s} \right).$$
(38)

Combining equations on n and p_D , we obtain model (12).

B.2 Model with partial conversion

B.2.1 Conversion in the zygote

We rewrite model (17) with variables:

$$n = n_{\rm W} + n_{\rm D}, \qquad p_{\rm D} = \frac{n_{\rm D}}{n_{\rm W} + n_{\rm D}}.$$
 (39)

where n is the total population density and p_D is the drive allele frequency.

Equation on n:

$$\begin{split} \partial_{t}n - \partial_{xx}^{2}n &= \frac{r(1-n)+1}{n} \left((1-s) n_{\mathrm{D}}^{2} + [2 c(1-s)+2 (1-sh) (1-c)] \right. \\ &\left. n_{\mathrm{D}}n_{\mathrm{W}} + n_{\mathrm{W}}^{2} \right) - n, \\ &= \frac{r(1-n)+1}{n} \left((1-s) (np_{\mathrm{D}})^{2} + [2 c(1-s)+2 (1-sh) (1-c)] \right. \\ &\left. p_{\mathrm{D}}(1-p_{\mathrm{D}})n^{2} + (1-p_{\mathrm{D}})^{2}n^{2} \right) - n, \\ &= \left(r(1-n)+1 \right) \left((1-s) p_{\mathrm{D}}^{2} + [2 c(1-s)+2 (1-sh) (1-c)] \right. \\ &\left. p_{\mathrm{D}}(1-p_{\mathrm{D}}) + (1-p_{\mathrm{D}})^{2} \right) n - n, \\ &= \left(r(1-n)+1 \right) \left((1-s) p_{\mathrm{D}}^{2} + 2 p_{\mathrm{D}} (1-p_{\mathrm{D}}) \left[c(1-s) + (1-c) (1-sh) \right] \right. \\ &\left. + (1-p_{\mathrm{D}})^{2} \right) n - n. \end{split}$$



Equation on n_D

$$\partial_{t} n_{D} = \partial_{xx}^{2} n_{D} + \frac{r(1-n)+1}{n} \left[(1-s)(n_{D}+2c n_{W}) + (1-sh) (1-c) n_{W} \right] n_{D} - n_{D},
= \partial_{xx}^{2} n_{D} + \frac{r(1-n)+1}{n} \left[(1-s)(n p_{D}+2c n (1-p_{D})) + (1-sh) (1-c) n_{D} \right] n_{D} - n_{D},
= n_{D} \partial_{xx}^{2} n_{D} + 2 \partial_{x} n_{D} \partial_{x} n_{D} + n_{D} \partial_{xx}^{2} n_{D}
+ (r(1-n)+1) \left[(1-s)(p_{D}+2c (1-p_{D})) + (1-sh) (1-c) (1-p_{D}) \right] n_{D} - n_{D}.$$
(41)

Equation on $\mathbf{p}_{\mathrm{D}} = \frac{\mathbf{n}_{\mathrm{D}}}{-}$:

$$\partial_{t} p_{D} = \frac{n (\partial_{t} n_{D}) - n_{D} (\partial_{t} n)}{n^{2}} = \frac{n (\partial_{t} n_{D}) - n p_{D} (\partial_{t} n)}{n^{2}} = \frac{1}{n} \Big(\partial_{t} n_{D} - p_{D} (\partial_{t} n) \Big), \\
= \frac{1}{n} \Big[2 \partial_{x} n \partial_{x} p_{D} + n \partial_{xx}^{2} p_{D} + (r (1 - n) + 1) \Big((1 - s)(p_{D} + 2 c (1 - p_{D})) \\
+ (1 - sh) (1 - c) (1 - p_{D}) \\
- (1 - s) p_{D}^{2} - 2 p_{D} (1 - p_{D}) [c (1 - s) + (1 - c) (1 - sh)] - (1 - p_{D})^{2} \Big) p_{D} n \Big], \\
= 2 \partial_{x} \log(n) \partial_{x} p_{D} + \partial_{xx}^{2} p_{D} + (r (1 - n) + 1) \Big(p_{D} (1 - s) (1 - p_{D}) \\
+ 2 (1 - s) c (1 - p_{D}) + (1 - sh) (1 - c) (1 - p_{D}) \\
- 2 p_{D} c (1 - s) (1 - p_{D}) - 2 p_{D} (1 - c) (1 - sh) (1 - p_{D}) - (1 - p_{D})^{2} \Big) p_{D} \Big], \\
= 2 \partial_{x} \log(n) \partial_{x} p_{D} + \partial_{xx}^{2} p_{D} + (r (1 - n) + 1) \Big(p_{D} \Big[(1 - s) (1 - 2c) \\
- 2 (1 - sh) (1 - c) + 1 \Big] + 2 (1 - s) c \\
+ (1 - sh) (1 - c) - 1 \Big) (1 - p_{D}) p_{D}, \\
= 2 \partial_{x} \log(n) \partial_{x} p_{D} + \partial_{xx}^{2} p_{D} \\
+ (r (1 - n) + 1) \Big([1 - 2(1 - c)(1 - h)] sp_{D} - s[1 - (1 - c)(1 - h)] \\
+ c(1 - s) \Big) (1 - p_{D}) p_{D}. \tag{42}$$

Combining equations on n and p_D , we obtain model (21).

B.2.2 Conversion in the germline

We rewrite model (27) with variables:

$$n = n_{\rm W} + n_{\rm D}, \qquad p_{\rm D} = \frac{n_{\rm D}}{n_{\rm W} + n_{\rm D}}.$$
 (43)

where n is the total population density and p_D is the drive allele frequency.

 $[\]overline{{}^2 \partial_{xx} : {}^2 n_{\mathrm{D}} = \partial_{xx}^2 n p_{\mathrm{D}} = \partial_x (p_{\mathrm{D}} \partial_x n + n \partial_x p_{\mathrm{D}}) = p_{\mathrm{D}} \partial_{xx}^2 n + 2 \partial_x p_{\mathrm{D}} \partial_x n + n \partial_{xx}^2 p_{\mathrm{D}}.$



30 Page 32 of 54 L. Kläy et al.

Equation on n:

$$\partial_{t}n - \partial_{xx}^{2}n = \frac{r(1-n)+1}{n} \left((1-s) n_{D}^{2} + 2(1-sh) n_{D}n_{W} + n_{W}^{2} \right) - n,$$

$$= \frac{r(1-n)+1}{n} \left((1-s) (np_{D})^{2} + 2(1-sh) p_{D}(1-p_{D})n^{2} + ((1-p_{D})n)^{2} \right) - n,$$

$$= \left(r(1-n)+1 \right) \left((1-s) p_{D}^{2} + 2(1-sh) p_{D}(1-p_{D}) + (1-p_{D})^{2} \right) n - n.$$
(44)

Equation on n_D

$$\partial_{t} n_{D} = \partial_{xx}^{2} n_{D} + \frac{r(1-n)+1}{n} \Big[(1-s)n_{D} + (1-sh)(1+c) n_{W} \Big] n_{D} - n_{D},
= p_{D} \partial_{xx}^{2} n + 2 \partial_{x} n \partial_{x} p_{D} + n \partial_{xx}^{2} p_{D} + \frac{r(1-n)+1}{n} \Big[(1-s)np_{D}
+ (1-sh)(1+c) n (1-p_{D}) \Big] n p_{D} - n p_{D},
= p_{D} \partial_{xx}^{2} n + 2 \partial_{x} n \partial_{x} p_{D} + n \partial_{xx}^{2} p_{D} + \Big(r(1-n)+1 \Big) \Big[(1-s)p_{D}
+ (1-sh)(1+c)(1-p_{D}) \Big] n p_{D} - n p_{D}.$$
Equation on $\mathbf{p}_{D} = \frac{\mathbf{n}_{D}}{\mathbf{n}}$:

$$\partial_{t} p_{D} = \frac{n (\partial_{t} n_{D}) - n_{D} (\partial_{t} n)}{n^{2}} = \frac{n (\partial_{t} n_{D}) - n p_{D} (\partial_{t} n)}{n^{2}} = \frac{1}{n} (\partial_{t} n_{D} - p_{D} (\partial_{t} n)),
= \frac{1}{n} \Big[2 \partial_{x} n \partial_{x} p_{D} + n \partial_{xx}^{2} p_{D} + (r (1 - n) + 1) ((1 - s) p_{D} + (1 - sh) (1 + c) (1 - p_{D}) - (1 - p_{D})^{2}) p_{D} n \Big],
+ (1 - s) p_{D}^{2} - 2 (1 - sh) p_{D} (1 - p_{D}) - (1 - p_{D})^{2}) p_{D} n \Big],
= 2 \partial_{x} \log(n) \partial_{x} p_{D} + \partial_{xx}^{2} p_{D} + (r (1 - n) + 1) ((1 - s) p_{D} + (1 - sh)(1 + c) - 2 (1 - sh) p_{D} - (1 - p_{D})) p_{D} (1 - p_{D}),
= 2 \partial_{x} \log(n) \partial_{x} p_{D} + \partial_{xx}^{2} p_{D} + (r (1 - n) + 1) ((2h - 1) s p_{D} + (1 - sh)(1 + c) - 1) p_{D} (1 - p_{D}).$$
(46)

Combining equations on n and p_D , we obtain model (29).

 $[\]overline{^3 \partial_{xx} :^2 n_{\mathbf{D}} = \partial_{xx}^2 n p_{\mathbf{D}} = \partial_x (p_{\mathbf{D}} \partial_x n + n \partial_x p_{\mathbf{D}}) = p_{\mathbf{D}} \partial_{xx}^2 n + 2 \partial_x p_{\mathbf{D}} \partial_x n + n \partial_{xx}^2 p_{\mathbf{D}}.$



C Proofs for model (12) with perfect conversion in the zygote

C.1 Numerical evidence for the continuity when $r \rightarrow 0$

In Fig. 7, we plot the speed of the traveling wave solutions of model (12) for a range of r and s values, and for r = 0. A positive speed correspond to drive invasion.

We observe continuity in the speed value when $r \to 0$ away from $s = \frac{1}{2}$, meaning that the case r = 0 is relevant to approximate very small intrinsic growth rates.

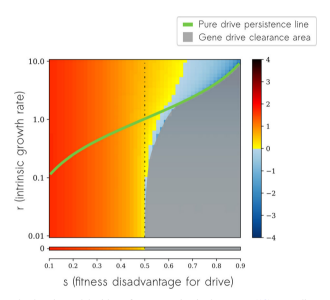


Fig. 7 Wave speed values in model with perfect conversion in the zygote (12), regarding parameters r the intrinsic growth rate (log scale in between 0.01 and 10, plus the exact value r = 0 in the bottom color line) and s the fitness disadvantage for drive (normal scale). Below the pure drive persistence line (light green), a well-mixed population containing only drive homozygous individuals will necessarily go extinct (color figure online)

C.2 Proof of the statements in Tables 3 and 4 when perfect conversion occurs in the zygote

In this section we prove the statements of Tables 3 and 4 on the two models of interest:

$$\mathbf{r} = \infty$$

$$\partial_t p - \partial_{xx}^2 p = \frac{s \ p \ (1-p) \left(p - \frac{2s-1}{s}\right)}{1 - s + s(1-p)^2} = f^{\infty}(p). \tag{47}$$



30 Page 34 of 54 L. Kläy et al.

 $\mathbf{r} = 0$

$$\begin{cases} \partial_{t} n_{\text{DD}} - \partial_{xx}^{2} n_{\text{DD}} &= (1 - s) \frac{n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}} - s n_{\text{DD}} = f^{0}(n_{\text{DD}}, n_{\text{WW}}) \\ \partial_{t} n_{\text{WW}} - \partial_{xx}^{2} n_{\text{WW}} &= \frac{-n_{\text{WW}} n_{\text{DD}}}{n_{\text{WW}} + n_{\text{DD}}}. \end{cases}$$
(48)

Monostable / Bistable

 $\mathbf{r} = \infty$

0 < s < 0.5 The equation admits two admissible steady states 0 and 1. As $(f^{\infty})'(0) > 0$ and $(f^{\infty})'(1) < 0$, the only stable state is p = 1.

0.5 < s < 1 The equation admits three admissible steady states $0, \frac{2s-1}{s}$ and 1.As $(f^{\infty})'(0) < 0$, $(f^{\infty})'(\frac{2s-1}{s}) > 0$ and $(f^{\infty})'(1) < 0$, both p=0 and p=1 are stable states.

 $\mathbf{r} = 0$

0 < s < 0.5

The system admits ($n_{DD} = 0$, $n_{WW} \in [0, 1]$) as admissible steady states. The Jacobian matrix, when switching to n and p_D variables, indicates that the only stable state is (n = 0, $p_D = 1$), i.e. ($n_{DD} = 0$, $n_{WW} = 0$).

0.5 < s < 1 The system admits $(n_{\rm DD} = 0, n_{\rm WW} \in [0, 1])$ as admissible steady states. The Jacobian matrix, when switching to n and $p_{\rm D}$ variables, indicates that the stable states are $(n = 0, p_{\rm D} = 1)$ and $(n \in [0, 1], p_{\rm D} = 0)$, i.e. $(n_{\rm DD} = 0, n_{\rm WW} \in [0, 1])$.

Existence of critical traveling waves

 $\mathbf{r} = \infty$

The existence of traveling waves for the scalar equation (13) in both monostable and bistable cases is a classical result in the theory of reaction-diffusion equations, see for instance the seminal works in Aronson and Weinberger (1975, 1978).

 $\mathbf{r} = 0$

0 < s < 0.5 We apply the results of Appendix D with $\beta_1 = 1$ and $\beta_2 = 1 - s$. Therefore system (48) admits a traveling wave when 0 < s < 0.5.

0.5 < s < 1 There is no drive propagation due to the gene drive clearance: the drive allele density decreases uniformly in space (details in Sect. C.2.1). Regarding the wild-type alleles, their dynamic is given by the heat equation implying only diffusion and no growth. It cannot admit traveling wave solutions.

Pulled/pushed waves and speed values

For both models, the speed of the linearized problem around zero density of drive allele is given by $2\sqrt{1-2s} = 2\sqrt{(f^{\infty})'(0)} = 2\sqrt{\partial_p f^0(0,1)}$.





- $0 < s \lesssim 0.35$ Numerically, we observe that the speed of the wave is equal to the minimal speed of the linearized problem: the wave is pulled (detail in Sect. C.2.2)
- $0.35 \lesssim s < 0.5$ Numerically, we observe that the speed of the wave is strictly above the minimal speed of the linearized problem: the wave is pushed (detail in Sect. C.2.2).
- 0.5 < s < 1 As the system is bistable, the wave is necessarily pushed. The numerical approximation $s \approx 0.70$ indicating whether the drive of the wild-type population will invade the environment was already determined in the work of Tanaka et al. (2017).

 $\mathbf{r} = 0$

- 0 < s < 0.5 We apply the results of Appendix D with $\beta_1 = 1$ and $\beta_2 = 1 s$. Therefore system (48) admits a traveling wave with speed $v = 2\sqrt{1-2s}$ when 0 < s < 0.5. This value corresponds to the KPP speed, by definition the wave is pulled.
- 0.5 < s < 1 No wave (see above, in Existence of critical traveling waves).

C.2.1 Gene drive clearance for $s \in (0.5, 1)$ when r = 0

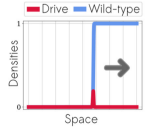
Consider the model with perfect conversion in the zygote (6). The densities n_{WW} and n_{DD} dynamics are qualitatively given in Fig. 8 for r = 0.

When s > 0.5 and r = 0 we observe gene drive clearance (in Fig. 8B). More precisely, we have the following estimate, deduced from (15):

$$\partial_t n_{\rm DD} - \partial_{xx}^2 n_{\rm DD} \le (1 - 2s) n_{\rm DD}, \tag{49}$$

Therefore, $n_{\rm DD}$ is exponentially decaying in time, uniformly in space. The dynamics of the wild type then boil down to the standard heat equation, there cannot exist a traveling wave.

(A) Spreading eradication drive when 0 < s < 0.5. (B) Gene drive clearance when 0.5 < s < 1



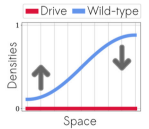


Fig. 8 Qualitative dynamics of the drive homozygotes density $n_{\rm DD}$ (red line) and the wild-type homozygotes density $n_{\rm WW}$ (blue line) in space (color figure online)

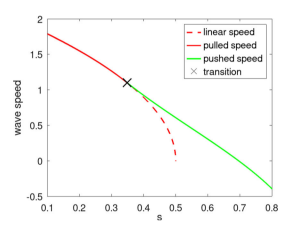


30 Page 36 of 54 L. Kläy et al.

C.2.2 Numerical approximation of s threshold value for the pulled/pushed wave when $r=+\infty$

In order to determine an approximation of the threshold value at which the wave switches from a pulled wave to a pushed wave, we used the recent continuation procedure published in Avery et al. (2022). Figure 9 presents the value of the wave speed obtained via the latter continuation numerical scheme (Holzer 2022), for a wide range of s values. Notice the transition between pulled fronts (plain red) and pushed fronts (plain green). For the sake of clarity, the value of the minimal speed of the linearized problem $v = 2\sqrt{1-2s}$ is shown in red for $s \in (0, \frac{1}{2})$. Notice that the speed of the pushed front changes sign approximately at $s \approx 0.70$, in agreement with the theoretical criterion (9).

Fig. 9 Value of the wave speed when $r = +\infty$ obtained via the numerical scheme in Holzer (2022), for a wide range of s values. The transition between pulled fronts (plain red) and pushed fronts (plain green) is approximately 0.35. For the sake of clarity, the value of the minimal speed of the linearized problem $v = 2\sqrt{1-2s}$ is shown in red for $s \in (0, \frac{1}{2})$ (color figure online)



D Critical travaling wave for an SI similar model.

Consider the following epidemiological model,

$$\begin{cases} \partial_t S - \partial_{xx}^2 S = -\beta \frac{SI}{S+I}, \\ \partial_t I - \partial_{xx}^2 I = \beta \frac{SI}{S+I} - \gamma I. \end{cases}$$
 (50)

where S is the density of susceptible individuals, I is the density of infected individuals, γ is the mortality of infected individuals and β is the transmission coefficient. This model has already been studied in the literature, see (Zhou et al. 2019) and references therein. In particular, the existence of a minimal traveling wave has been established in the latter reference.



Models (15), (25) and (33) are very similar to the above SI model (50), except that the coefficient β is different in the first and the second equation of the system. We write this new system with two coefficients β_1 , β_2 :

$$\begin{cases} \partial_t S - \partial_{xx}^2 S = -\beta_1 \frac{SI}{S+I}, \\ \partial_t I - \partial_{xx}^2 I = \beta_2 \frac{SI}{S+I} - \gamma I. \end{cases}$$
 (51)

D.1 Existence of critical traveling wave solutions

We are able to establish the following Theorem by adapting the proof in Zhou et al. (2019).

Theorem Suppose that $\beta_1 > 0$, and $\beta_2 > \gamma$, then system (51) admits a positive and bounded traveling wave solution with profile (S^*, I^*) , and speed $v = 2\sqrt{\beta_2 - \gamma}$. Furthermore, both S^* and I^* are positive, and bounded by 1 and $\frac{\beta_2 - \gamma}{\gamma}$ respectively.

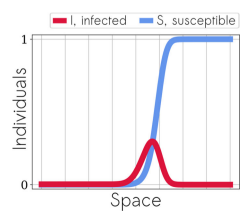
By adapting further the elements of (Zhou et al. 2019), it would be possible to prove that the profile S^* is increasing, whereas the profil I^* is unimodal, as shown in Fig. 10.

D.2 Proof of the theorem

We proceed as follows:

- 1. Although the system does not satisfy the comparison principle due to a lack of monotonicity, the construction of traveling waves is performed through a construction of sub-solutions $(\underline{S}, \underline{I})$ and super-solutions $(\overline{S}, \overline{I})$ for the system.
- 2. Using Schauder's fixed point theorem, we prove the existence of a critical traveling wave solution (S^*, I^*) with speed v such that $\underline{S}(z) \leq S^*(z) \leq \overline{S}(z)$ and $\underline{I}(z) \leq I(z) < \overline{I}(z)$ for all z in \mathbb{R} .

Fig. 10 Qualitative shape of the solution (I^* in red, S^* in blue)





30 Page 38 of 54 L. Kläy et al.

3. Finally, we conclude with the positivity of the critical traveling wave solution thanks to the strong maximum principle.

D.2.1 Construction of sub- and super-solutions

We are seeking sub- and super-solutions, respectively $(\underline{S}, \underline{I})$, (\bar{S}, \bar{I}) . Because of the non-monotonic coupling in the system, the following set of cross-relationships must be satisfied:

1.
$$-v \ \bar{S}' - \bar{S}'' \ge -\beta_1 \frac{\bar{S} I}{\bar{S} + I} \quad \forall \underline{I} \le I \le \bar{I};$$

2. $-v \ \underline{S}' - \underline{S}'' \le -\beta_1 \frac{\underline{S} I}{\underline{S} + I} \quad \forall \underline{I} \le I \le \bar{I};$
3. $-v \ \bar{I}' - \bar{I}'' \ge \beta_2 \frac{S \ \bar{I}}{S + \bar{I}} - \gamma \ \bar{I} \quad \forall \underline{S} \le S \le \bar{S};$
4. $-v \ \underline{I}' - \underline{I}'' \le \beta_2 \frac{S \ \underline{I}}{S + I} - \gamma \ \underline{I} \quad \forall \underline{S} \le S \le \bar{S}.$

Inequalities 1,2,3 and 4 are valid in a weak sense. As \underline{S} is a piece-wise differentiable function, the quantity \underline{S} consists of functions on each sub-interval with a Dirac mass at the point of \mathcal{C}^1 discontinuity. However, the Dirac mass has the good sign in this case (the transition has a convex shape), hence the second derivative \underline{S}'' is non-negative in the sense of a measure. All signs are correct: \underline{I} has a convex transition also, and \overline{I} has a concave transition. This key point is equivalent to the standard principle in the theory of parabolic equations (widely used for reaction-diffusion equations involving comparison techniques): "the maximum of sub-solutions is a sub-solution" (here, \underline{S} , \overline{I}) and "the minimum of super-solutions is a super-solution" (here, \overline{S} , \overline{I}).

To define our sub and super-solutions, it is useful to introduce the the following family of functions $\mathcal{I}(z) = e^{-\lambda^* z}$, where λ^* is solution of the following dispersion equation:

$$(\lambda^*)^2 - v\lambda^* + (\beta_2 - \gamma) = 0.$$
 (52)

They are solutions of the linearized problem

$$v\mathcal{I}' + \mathcal{I}'' + (\beta_2 - \gamma) \mathcal{I} = 0. \tag{53}$$

For the critical speed $v = 2\sqrt{(\beta_2 - \gamma)}$, the corresponding double root is $\lambda^* = \frac{v}{2} = \sqrt{(\beta_2 - \gamma)}$.

Lemma There exist two large enough constants $L_1 > 0$ and $L_2 > 0$, such that the functions \bar{S} , \underline{S} \bar{I} , \underline{I} defined below satisfy the conditions 1. 2. 3. and 4.:

$$\bar{S} = 1.$$

$$\underline{S} = \begin{cases}
0 & \forall z \le L_1 \log(L_1), \\
\frac{z}{L_1 - L_1 e^{-\frac{z}{L_1}}} & \forall z \ge L_1 \log(L_1)
\end{cases}$$
(54)



$$\bar{I} = \begin{cases} M & \forall z \le \frac{1}{\lambda^*}, \\ eM\lambda^* z e^{-\lambda^* z} & \forall z > \frac{1}{\lambda^*}. \end{cases}$$
 (56)

$$\underline{I} = \begin{cases} 0 & \forall z \le \left(\frac{L_2}{eM\lambda^*}\right)^2, \\ \left(eM\lambda^*z - L_2\sqrt{z}\right)e^{-\lambda^*z} & \forall z > \left(\frac{L_2}{eM\lambda^*}\right)^2. \end{cases}$$
 (57)

with
$$M = \frac{\beta_2 - \gamma}{\gamma} = \frac{(\lambda^*)^2}{\gamma}$$
.

Proof Before we proceed with the proof, we introduce the following set of conditions, which are more restrictive than 1,2,3,4, but may appear more useful at some point in the calculations. When satisfied, they clearly imply 1,2,3,4.

- (i) $-v \bar{S}' \bar{S}'' > 0$:
- (ii) $-v \underline{S'} \underline{S''} \leq -\beta_1 \overline{I};$ (iii) $-v \overline{I'} \overline{I''} \geq (\beta_2 \gamma) \overline{I};$

(iii)
$$v I = \sum_{i} (\beta_{2} \gamma_{i}) I$$
,
(iv) $-v \underline{I}' - \underline{I}'' \leq \beta_{2} \frac{\underline{S} \underline{I}}{\underline{S} + \underline{I}} - \gamma \underline{I}$.

We verify each of the four conditions on \bar{S} , \underline{S} \bar{I} , \underline{I} :

Condition 1:
$$-v \ \bar{S}' - \bar{S}'' \ge -\beta_1 \ \frac{\bar{S} \ I}{\bar{S} + I}$$
.

The constant function $\bar{S}=1$ satisfies the more restrictive condition (i) $-v\ \bar{S}'-\bar{S}''$

Condition 2:
$$\underline{S}' - \underline{S}'' \le -\beta_1 \frac{\underline{S} I}{\underline{S} + I}$$
.

Let us take L_1 sufficiently large such that $\frac{1}{\lambda^*} < L_1 \log(L_1)$.

• For $z > L_1 \log(L_1)$:

Since $\bar{I} = e \ M \ \lambda^* \ z \ e^{\lambda^* z}$ and $\underline{S} = 1 - L_1 e^{-\frac{\zeta}{L_1}}$, the condition (ii) $-v \ \underline{S'} - \underline{S''} \le$ $-\beta_1 \bar{I}$ holds for a sufficiently large $L_1 > 0$:

$$-v \underline{S}' - \underline{S}'' = (\frac{1}{L_1} - v) e^{-\frac{z}{L_1}} \le -\beta_1 e M \lambda^* z e^{-\lambda^* z} = -\beta_1 \bar{I}, \qquad (58)$$

$$\iff \beta_1 \ e \ M \ \lambda^* \ z \ e^{\left(\frac{1}{L_1} - \lambda^*\right)z} \le (v - \frac{1}{L_1}). \tag{59}$$

• For $z \leq L_1 \log(L_1)$:

The condition 2. is verified since S = 0.

Condition 3:
$$-v \ \bar{I}' - \bar{I}'' \ge \beta_2 \frac{S \ \bar{I}}{S + \bar{I}} - \gamma \ \bar{I}.$$



30 Page 40 of 54 L. Kläy et al.

• For
$$z \leq \frac{1}{\lambda^*}$$
:

With
$$\bar{I} = M = \frac{\beta_2 - \gamma}{\gamma}$$
:

$$-v \,\bar{I}' - \bar{I}'' = 0 = \beta_2 \, \frac{M}{1+M} - \gamma \, M = \beta_2 \, \frac{\bar{S} \,\bar{I}}{\bar{S} + \bar{I}} - \gamma \, \bar{I} \ge \beta_2 \, \frac{S \,\bar{I}}{S + \bar{I}} - \gamma \, \bar{I} \qquad \forall S \le \bar{S}.$$
(60)

• For $z > \frac{1}{\lambda^*}$:

Since \bar{I} is proportional to $ze^{-\lambda^*z}$, and λ^* is precisely the double root of the characteristic equation (52), we deduce that condition (iii) $-v\ \bar{I}' - \bar{I}'' - (\beta_2 - \gamma)\ \bar{I} \ge 0$ is verified.

Condition 4: $-v \underline{I}' - \underline{I}'' \le \beta_2 \frac{S\underline{I}}{S + \underline{I}} - \gamma \underline{I}$. Let us take L_2 sufficiently large such that $L_2 > eM\lambda^* \sqrt{L_1 \log(L_1)}$.

• For $z \le \left(\frac{L_2}{e \ M \ \lambda^*}\right)^2$:

I = 0 so condition 4. is satisfied.

• For $z > \left(\frac{L_2}{e \ M \ \lambda^*}\right)^2$:

The choice of L_2 implies $z > L_1 \log(L_1)$, which means $\underline{S} = 1 - L_1 e^{-\frac{z}{L_1}}$. We can reformulate condition (iv) as follows:

$$-v \underline{I'} - \underline{I''} \le \beta_2 \frac{\underline{S} \underline{I}}{S + I} - \gamma \underline{I} \iff -v \underline{I'} - \underline{I''} - (\beta_2 - \gamma) \underline{I} \le -\beta_2 \frac{\underline{I}^2}{S + I}.$$
 (61)

With $L_3 = eM\lambda^*$, and:

$$\underline{I} = [L_3 z - L_2 \sqrt{z}] e^{-\lambda^* z}, \tag{62}$$

$$\underline{I}' = [L_3 - L_2 \frac{1}{2\sqrt{z}}] e^{-\lambda^* z} - [L_3 z - L_2 \sqrt{z}] \lambda^* e^{-\lambda^* z}, \tag{63}$$

$$\underline{I}'' = \left[L_2 \frac{1}{4 z \sqrt{z}}\right] e^{-\lambda^* z} - 2 \left[L_3 - L_2 \frac{1}{2 \sqrt{z}}\right] \lambda^* e^{-\lambda^* z} + \left[L_3 z - L_2 \sqrt{z}\right] (\lambda^*)^2 e^{-\lambda^* z}$$
(64)

On the one hand, we obtain the following identities:

$$-v \underline{I}' - \underline{I}'' - (\beta_2 - \gamma) \underline{I} = e^{-\lambda^* z} \Big[L_3 \Big(-v + v \lambda^* z + 2 \lambda^* - (\lambda^*)^2 z - (\beta_2 - \gamma) z \Big)$$

$$+ L_2 \Big(v \frac{1}{2\sqrt{z}} - v \sqrt{z} \lambda^* - \frac{1}{4z\sqrt{z}} - \lambda^* \frac{1}{\sqrt{z}} + (\lambda^*)^2 \sqrt{z} \Big)$$
(65)



$$+ (\beta_{2} - \gamma) \sqrt{z} \Big) \Big],$$

$$= e^{-\lambda^{*}z} \Big[L_{3} \Big((2\lambda^{*} - v) - z \Big(- v\lambda^{*} + (\lambda^{*})^{2} + (\beta_{2} - \gamma) \Big) \Big) \Big)$$

$$+ L_{2} \Big(\sqrt{z} \Big(- v\lambda^{*} + (\lambda^{*})^{2} + (\beta_{2} - \gamma) \Big) + \frac{1}{2\sqrt{z}} (v - 2\lambda^{*})$$

$$- \frac{1}{4z\sqrt{z}} \Big) \Big],$$

$$= -L_{2} e^{-\lambda^{*}z} \frac{1}{4z\sqrt{z}}$$
(69)

On the other hand, we have:

$$-\beta_2 \frac{\underline{I}^2}{\underline{S} + \underline{I}} = -\beta_2 \frac{[L_3 z - L_2 \sqrt{z}]^2 e^{-2\lambda^* z}}{1 - L_1 e^{-\frac{z}{L_1}} + [L_3 z - L_2 \sqrt{z}] e^{-\lambda^* z}}.$$
 (70)

We resume with the reformulation (61), which is now equivalent to the following:

$$-L_{2} e^{-\lambda^{*}z} (1 - L_{1} e^{-\frac{z}{L_{1}}} + [L_{3} z - L_{2} \sqrt{z}] e^{-\lambda^{*}z}) \leq -4\beta_{2} [L_{3} z - L_{2} \sqrt{z}]^{2} e^{-2\lambda^{*}z} z \sqrt{z}$$

$$(71)$$

$$\iff 4\beta_{2} [L_{3} z - L_{2} \sqrt{z}]^{2} e^{-\lambda^{*}z} z \sqrt{z} - L_{2} [L_{3} z - L_{2} \sqrt{z}] e^{-\lambda^{*}z} \leq L_{2} (1 - L_{1} e^{-\frac{z}{L_{1}}})$$

$$(72)$$

$$\iff 4\beta_{2} e^{-\lambda^{*}z} z^{3} \sqrt{z} (L_{3})^{2} + e^{-\lambda^{*}z} \left((1 - 8\beta_{2} z^{2}) L_{3} L_{2} z + (L_{2})^{2} \sqrt{z} (1 - 4\beta_{2} z^{2}) \right)$$

$$\leq L_{2} (1 - L_{1} e^{-\frac{z}{L_{1}}}).$$

$$(73)$$

We may increase L_2 such that $1 - 4\beta_2 \left(\frac{L_2}{eM\lambda^*}\right)^4 \le 0$. Then, since $z > \left(\frac{L_2}{eM\lambda^*}\right)^2$:

$$1 - 8\beta_2 z^2 \le 1 - 4\beta_2 z^2 < 1 - 4\beta_2 \left(\frac{L_2}{eM\lambda^*}\right)^4 \le 0.$$
 (74)

Since $(1 - 8\beta_2 z^2)$ and $(1 - 4\beta_2 z^2)$ are negative terms, we need to show L_2 (1 - $L_1 e^{-L_1}$) $\geq 4\beta_2 e^{-\lambda^* z} z^3 \sqrt{z} (L_3)^2$. Let $g(z) = \beta_2 e^{-\lambda^* z} z^3 \sqrt{z} (L_3)^2$ be a $\mathcal{C}^1([0; -\infty[)$ function. Since $\lim_{z \to 0} (g(z)) = 0$ and $\lim_{z \to +\infty} (g(z)) = 0$ there exists a constant C (which is independent from L_2) such that $g(z) < C \ \forall z \ge 0$. We finally increase L_2 so that condition (iv) is verified.



30 Page 42 of 54 L. Kläy et al.

D.2.2 Existence and positivity of a critical traveling wave solution

Now, exactly as in Zhou et al. (2019), we are in a position to define a set of functions

$$\Gamma = \{ (S, I) \in B_{\mu}(\mathbb{R}, \mathbb{R}^2) \mid \underline{S} \le S \le \overline{S}, \ \underline{I} \le I \le \overline{I} \},$$

where $B_{\mu}(\mathbb{R}, \mathbb{R}^2)$ is the set of two-component continuous functions with each component growing at infinity slower than $e^{\mu|z|}$, as well as an operator $F: \Gamma \to C(\mathbb{R}, \mathbb{R}^2)$ that will satisfy the assumptions of the Schauder fixed point theorem and whose fixed point in Γ will precisely be the solution (S, I) we seek. Note that the inequalities 1., 2., 3., 4. (beginning of Sect. D.2.1) are precisely what we use to prove that $F(\Gamma) \subset \Gamma$. Details can be found in Zhou et al. (2019).

The positivity of both S and I comes from the use of the strong maximum principle, again exactly as in Zhou et al. (2019).

E Study of the reaction term when $r = +\infty$ in section 3.2

We are searching for conditions implying a pulled monostable wave, using criterion (10).

E.1 Conversion occurring in the zygote

We rewrite limit equation (22):

$$\frac{\partial_{t} p_{D} - \partial_{xx}^{2} p_{D}}{\int_{-s}^{\infty} \left[2(1-c)(1-h) - 1 \right] s \ p_{D} - s[1-(1-c)(1-h)] + c(1-s) \right) (1-p_{D}) \ p_{D}}{\int_{-s}^{\infty} \left[2(1-c)(1-h) - 1 \right] p_{D}^{2} - 2s[1-(1-c)(1-h)] p_{D} + 1}.$$
(75)

With $\mathcal{A}_z := s [2(1-c)(1-h) - 1] \in [-s, s]$:

$$\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D} = \frac{\left(-\mathscr{A}_z \ p_{\rm D} + \frac{1}{2}(\mathscr{A}_z - s) + c(1 - s)\right) (1 - p_{\rm D}) \ p_{\rm D}}{-\mathscr{A}_z p_{\rm D}^2 + (\mathscr{A}_z - s) \ p_{\rm D} + 1}.$$
 (76)

Note that the mean fitness $M_z(p_D) = -\mathcal{A}_z p_D^2 + (\mathcal{A}_z - s) p_D + 1 \in [1 - s, 1]^4$. When $\mathcal{A}_z \neq 0$, equation (76) can be rewritten:

$$\partial_{t} p_{\mathrm{D}} - \partial_{xx}^{2} p_{\mathrm{D}} = \frac{-\mathscr{A}_{z} (p_{\mathrm{D}} - p_{\mathrm{D}z}^{*}) (1 - p_{\mathrm{D}}) p_{\mathrm{D}}}{-\mathscr{A}_{z} p_{\mathrm{D}}^{2} + (\mathscr{A}_{z} - s) p_{\mathrm{D}} + 1} \quad \text{with} \quad p_{\mathrm{D}z}^{*} := \frac{1}{2} + \frac{2c (1 - s) - s}{2\mathscr{A}_{z}}.$$
(77)

 $[\]overline{{}^4 M_Z'(p_{\mathrm{D}}) = \mathscr{A}_Z(1 - 2p_{\mathrm{D}}) - s} \le 0 \text{ therefore } M_Z(1) \le M_Z(p_{\mathrm{D}}) \le M_Z(0).$



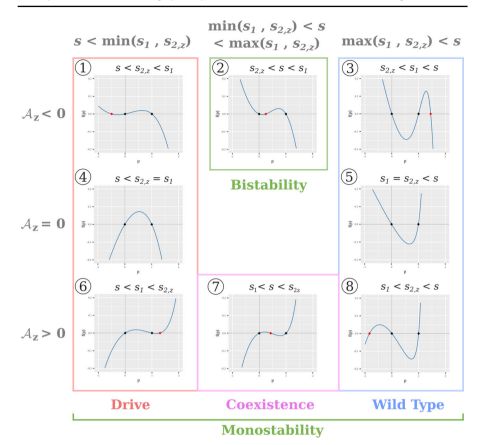


Fig. 11 Reaction term of equation (76) regarding the sign of $\mathcal{A}_z = s[2(1-c)(1-h)-1]$ and the s values. The s threshold values are $s_1 = \frac{c}{1-h(1-c)}$ and $s_{2,z} = \frac{c}{2c+h(1-c)}$. The dots on the x axis correspond to the steady states: $p_D = 0$ and $p_D = 1$ in black, $p_{Dz}^* = \frac{1}{2} + \frac{2c}{2\mathscr{A}_z}$ in red when it exists (color figure online)

Let us introduce $s_1 := \frac{c}{1 - h(1 - c)}$ and $s_{2,z} := \frac{c}{2c + h(1 - c)}$. Note that $\mathscr{A}_z > 0 \iff s_1 < s_{2,z}$. We draw the reaction term regarding the sign of \mathscr{A}_z and the s values in Fig. 11.

When $\mathcal{A}_z < 0$ and $s \in (s_{2,z}, s_1)$, equation (22) admits two stable steady states (bistability). The final proportion will then strongly depend on the initial condition. On the other hand, when $\mathcal{A}_z > 0$ and $s \in (s_1, s_{2,z})$, the only possible equilibrium state is a coexistence state: the final proportion p_D will be strictly in between 0 and 1.

Independently of the sign of \mathcal{A}_z , if $s < \min(s_1, s_{2,z})$ the only stable steady state is $p_D = 1$ meaning that for an initial condition outside of the steady states, we expect that the drive always invades the population. If $s > \max(s_1, s_{2,z})$, the only stable steady state is $p_D = 0$ meaning that for an initial condition outside of the steady states, we expect that the wild-type always invades the population.



30 Page 44 of 54 L. Kläy et al.

In case of bistability, the wave is always pushed (Hadeler and Rothe 1975); we can dismiss condition ② in the research of pulled monostable waves. We use criterion (10) on monostable cases, i.e. drive invasion ③ ④ ⑥, wild type invasion ③ ⑤ ⑧, and coexistence state ⑦ (the numbers refer to the subgraphs in Fig. 11).

E.1.1 Monostable drive invasion

(4) When $\mathscr{A}_z = 0$ and $s < s_1 = s_{2,z} = \frac{2c}{2c+1} \iff s < 2c \ (1-s)$ From equation (76), we have for all $p_D \in [0, 1]$:

$$\sigma(0) - (1 - p_{D}) \sigma(p_{D}) = \left(c(1 - s) - \frac{s}{2}\right) - \frac{\left(c(1 - s) - \frac{s}{2}\right)(1 - p_{D})}{1 - sp_{D}}$$
$$= \left(c(1 - s) - \frac{s}{2}\right) \frac{(1 - s) p_{D}}{1 - sp_{D}} \ge 0, \tag{78}$$

where σ is the selection term defined by equation (8). Criterion (10) is verified.

6 When $\mathcal{A}_z > 0$ and $p_{D_z}^* > 1$

From equation (77), we have for all $p_D \in [0, 1]$:

$$\sigma(0) - (1 - p_{D}) \sigma(p_{D}) = \mathcal{A}_{z} p_{Dz}^{*} - \frac{-\mathcal{A}_{z} (p_{D} - p_{Dz}^{*}) (1 - p_{D})}{-\mathcal{A}_{z} p_{D}^{2} + (\mathcal{A}_{z} - s) p_{D} + 1}$$

$$= \mathcal{A}_{z} p_{D} \frac{-(\mathcal{A}_{z} p_{Dz}^{*} + 1) p_{D} + (\mathcal{A}_{z} + 1 - s) p_{Dz}^{*} + 1}{-\mathcal{A}_{z} p_{D}^{2} + (\mathcal{A}_{z} - s) p_{D} + 1}.$$
(79)

Note that $-\mathscr{A}_z p_D^2 + (\mathscr{A}_z - s) \ p_D + 1 > (1 - s) > 0$ and $\mathscr{A}_z p_D > 0$. The affine term $-(\mathscr{A}_z p_{Dz}^* + 1) \ p_D + (\mathscr{A}_z + 1 - s) \ p_{Dz}^* + 1$ decreases with p_D . In order to show that it is positive for all $p_D \in [0, 1]$, we just need to verify that this it is true for $p_D = 1$:

$$-(\mathscr{A}_z p_{\mathrm{D}z}^* + 1) + (\mathscr{A}_z + 1 - s) \ p_{\mathrm{D}z}^* + 1 = (1 - s) \ p_{\mathrm{D}z}^* \ge 0 \quad \Rightarrow \quad \sigma(0) - (1 - p_{\mathrm{D}}) \ \sigma(p_{\mathrm{D}})$$

$$\ge 0 \forall p_{\mathrm{D}} \in [0, 1]. \tag{80}$$

Criterion (10) is verified.

① When $\mathscr{A}_Z < 0$ and $p_{\mathrm{D}_Z}^* < 0$ and $s < s_{2,z} \iff 0 < c - 2sc - sh + sch$ We consider equation (79) with $-\mathscr{A}_Z p_{\mathrm{D}}^2 + (\mathscr{A}_Z - s) p_{\mathrm{D}} + 1 > 1 - s > 0$ and $\mathscr{A}_Z p_{\mathrm{D}} < 0$. The affine term $-(\mathscr{A}_Z p_{\mathrm{D}_Z}^* + 1) p_{\mathrm{D}} + (\mathscr{A}_Z + 1 - s) p_{\mathrm{D}_Z}^* + 1$ decreases with p_{D} . In order to show that it is negative for all $p_{\mathrm{D}} \in [0, 1]$, we introduce a condition implying the negativity for $p_{\mathrm{D}} = 0$:

$$(\mathscr{A}_{z}+1-s) \ p_{\mathrm{D}z}^{*}+1 = \frac{(\mathscr{A}_{z}+1-s) (\mathscr{A}_{z}+2c(1-s)-s+2\mathscr{A}_{z})}{2\mathscr{A}_{z}} < 0$$

$$\iff \Big(1-2s[1-(1-h)(1-c)]\Big)\Big(c-2sc-sh+sch\Big)+s\Big[2(1-c)(1-h)-1\Big] > 0$$
(82)

Criterion (10) is verified when condition (82) is true.



E.1.2 Monostable wild-type invasion

In case of a monostable wild-type invasion, we need to consider the wild-type proportion $p_W = 1 - p_D \in [0, 1]$ and rewrite the equation (76):

$$-\partial_{t} p_{W} + \partial_{xx}^{2} p_{W} = \frac{\left(-\mathscr{A}_{z} (1 - p_{W}) + \frac{1}{2} (\mathscr{A}_{z} - s) + c(1 - s)\right) (1 - p_{W}) p_{W}}{-\mathscr{A}_{z} (1 - p_{W})^{2} + (\mathscr{A}_{z} - s)(1 - p_{W}) + 1}$$

$$\iff \partial_{t} p_{W} - \partial_{xx}^{2} p_{W} = \frac{\left(-\mathscr{A}_{z} p_{W} + \frac{1}{2} (\mathscr{A}_{z} + s) - c(1 - s)\right) (1 - p_{W}) p_{W}}{-\mathscr{A}_{z} p_{W}^{2} + (\mathscr{A}_{z} + s) p_{W} + (1 - s)}$$
(83)

When $\mathcal{A}_z \neq 0$, equation (83) can be rewritten:

$$\partial_{t} p_{W} - \partial_{xx}^{2} p_{W} = \frac{-\mathscr{A}_{z} (p_{W} - p_{Wz}^{*}) (1 - p_{W}) p_{W}}{-\mathscr{A}_{z} p_{W}^{2} + (\mathscr{A}_{z} + s) p_{W} + (1 - s)} \text{ with } p_{Wz}^{*}$$

$$= \frac{1}{2} - \frac{2c (1 - s) - s}{2\mathscr{A}_{z}} = 1 - p_{Dz}^{*}$$
(84)

(5) When $\mathcal{A}_z = 0$ and $s < s_1 = s_{2,z} = \frac{2c}{2c+1} \iff 2c (1-s) < s$ From equation (83) we have for all $p_W \in [0, 1]$:

$$\sigma(0) - (1 - p_{W}) \sigma(p_{W}) = \left(\frac{s}{2} - c(1 - s)\right) \left(\frac{1}{1 - s} - \frac{1 - p_{W}}{sp_{W} + 1 - s}\right) \ge \frac{p_{W}\left(\frac{s}{2} - c(1 - s)\right)}{1 - s} \ge 0 (85)$$

Criterion (10) is verified.

8 When $\mathcal{A}_z > 0$ and $p_{W_z}^* = 1 - p_{D_z}^* > 1$ From equation (84), we have for all $p_W \in [0, 1]$:

$$\sigma(0) - (1 - p_{W}) \sigma(p_{W}) = \frac{\mathscr{A}_{z} p_{Wz}^{*}}{1 - s} - \frac{-\mathscr{A}_{z} (p_{W} - p_{Wz}^{*}) (1 - p_{W})}{-\mathscr{A}_{z} p_{W}^{2} + (\mathscr{A}_{z} + s) p_{W} + (1 - s)}$$

$$= \mathscr{A}_{z} p_{W} \frac{(-p_{Wz}^{*} + 1 - s) p_{W} + p_{Wz}^{*} (\mathscr{A}_{z} + 1) + (1 - s)}{(1 - s)(-\mathscr{A}_{z} p_{W}^{2} + (\mathscr{A}_{z} + s) p_{W} + (1 - s))}.$$
(86)

Note that $(1-s)(-\mathscr{A}_z p_W^2 + (\mathscr{A}_z + s) \ p_W + (1-s)) > (1-s)^2 > 0$ and $\mathscr{A}_z p_W > 0$. As $p_{Wz}^* > 1$, the affine term $(-p_{Wz}^* + 1 - s) \ p_W + p_{Wz}^* (\mathscr{A}_z + 1) + (1-s)$ decreases with p_W . In order to show that it is positive for all $p_W \in [0, 1]$, we just need to verify that this it is true for $p_W = 1$:

$$(-p_{W_z}^* + 1 - s) + p_{W_z}^* (\mathscr{A}_z + 1) + (1 - s) = 2 (1 - s) + \mathscr{A}_z p_{W_z}^* \ge 0$$

$$\Rightarrow \sigma(0) - (1 - p_W) \sigma(p_W) \ge 0 \quad \forall p_W \in [0, 1]. \tag{87}$$

Criterion (10) is verified.

(3) When $\mathcal{A}_z < 0$ and $p_{W_z}^* = 1 - p_{D_z}^* < 0$



30 Page 46 of 54 L. Kläy et al.

We consider equation (86) with $(1-s)(-\mathscr{A}_z p_W^2 + (\mathscr{A}_z + s) p_W + (1-s)) > (1-s)^2 > 0$ and $\mathscr{A}_z p_W < 0$. The affine term $(-p_{Wz}^* + 1 - s) p_W + p_{Wz}^* (\mathscr{A}_z + 1) + (1-s)$ is strictly positive for $p_W = 1$, therefore criterion (10) is not verified.

E.1.3 Monostable coexistence state

(7) When $\mathcal{A}_z > 0$ and $0 < p_{D_z}^* = 1 - p_{W_z}^* < 1$

In the coexistence case, we have to verify that both waves, the drive invasion wave going to the right and the wild-type invasion wave going to the left, are pulled waves (see Fig. 3).

For the drive invasion wave we consider equation (79) with $0 < p_{\mathrm{D}z}^* < 1$ and $p_{\mathrm{D}} \in [0, p_{\mathrm{D}z}^*]$ (the term drive wave implies that the proportion of wild type increases after the wave passes; therefore the global stable steady state $p_{\mathrm{D}z}^*$ is also the maximum proportion). Once again, we need to prove that the affine term $-(\mathscr{A}_z p_{\mathrm{D}z}^* + 1) p_{\mathrm{D}} + (\mathscr{A}_z + 1 - s) p_{\mathrm{D}z}^* + 1$ is positive. As it decreases with $p_{\mathrm{D}} \in [0, p_{\mathrm{D}z}^*]$, we determine its sign for $p_{\mathrm{D}} = p_{\mathrm{D}z}^*$:

$$-(\mathscr{A}_{z}p_{\mathrm{D}z}^{*}+1)\ p_{\mathrm{D}z}^{*}+(\mathscr{A}_{z}+1-s)\ p_{\mathrm{D}z}^{*}+1 = -\mathscr{A}_{z}\ (p_{\mathrm{D}z}^{*})^{2}+(\mathscr{A}_{z}-s)\ p_{\mathrm{D}z}^{*}+1 \geq 1-s \geq 0 \\ \Rightarrow \ \sigma(0)-(1-p_{\mathrm{D}})\sigma(p_{\mathrm{D}}) \geq 0 \quad \forall p_{\mathrm{D}} \in [0,p_{\mathrm{D}z}^{*}].$$
 (88)

Criterion (10) is verified for the drive wave.

For the wild-type invasion wave, we consider equation (86) with $0 < p_{Wz}^* = 1 - p_{Dz}^* < 1$ and $p_W \in [0, p_{Wz}^*]$ (the term wild-type wave implies that the proportion of wild type increases after the wave passes; therefore the global stable steady state p_{Wz}^* is also the maximum proportion). Once again, we need to prove that the affine term $(-p_{Wz}^* + 1 - s) p_W + p_{Wz}^* (\mathscr{A}_z + 1) + (1 - s)$ is positive. As it decreases with $p_W \in [0, p_{Wz}^*]$, we determine its sign for $p_W = p_{Wz}^*$:

$$(-p_{Wz}^* + 1 - s) \ p_{Wz}^* + (\mathscr{A}_z + 1) \ p_{Wz}^* + (1 - s) = -(p_{Wz}^*)^2 + (\mathscr{A}_z + 2 - s) \ p_{Wz}^* + 1 - s$$

$$\geq \min(1, \mathscr{A}_z + 2(1 - s)) \geq 0 \quad \Rightarrow \quad \sigma(0) - (1 - p_W)\sigma(p_W) \geq 0 \quad \forall p_W \in [0, p_{Wz}^*].$$
(89)

Criterion (10) is verified for the wild-type wave.

E.2 Conversion occurring in the germline

We rewrite limit equation (30):

$$\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D} = \frac{\left(-(1-2h)\,s\,p_{\rm D} + [(1-sh)(1+c)-1]\right)\,p_{\rm D}\,(1-p_{\rm D})}{-s(1-2h)p_{\rm D}^2 - 2shp_{\rm D} + 1} \tag{90}$$



With $\mathcal{A}_g := s (1 - 2h) \in [-s, s]$:

$$\partial_t p_{\rm D} - \partial_{xx}^2 p_{\rm D} = \frac{\left(-\mathscr{A}_g \ p_{\rm D} + \frac{1}{2}(\mathscr{A}_g - s) + c(1 - sh)\right) p_{\rm D} (1 - p_{\rm D})}{-\mathscr{A}_g p_{\rm D}^2 + (\mathscr{A}_g - s) p_{\rm D} + 1}.$$
 (91)

Note that the mean fitness $M_g(p_D) = -\mathcal{A}_g p_D^2 + (\mathcal{A}_g - s) p_D + 1 \in [1 - s, 1]^5$. When $\mathcal{A}_g \neq 0$, equation (91) can be rewritten:

$$\partial_{t} p_{D} - \partial_{xx}^{2} p_{D} = \frac{-\mathscr{A}_{g} (p_{D} - p_{Dg}^{*}) (1 - p_{D}) p_{D}}{-\mathscr{A}_{g} p_{D}^{2} + (\mathscr{A}_{g} - s) p_{D} + 1} = f_{g}(p_{D}) \text{ with}$$

$$p_{Dg}^{*} := \frac{1}{2} + \frac{2c (1 - sh) - s}{2 \mathscr{A}_{g}}.$$
(92)

Let us introduce $s_1 := \frac{c}{1 - h(1 - c)}$ and $s_{2,g} := \frac{c}{2ch + h(1 - c)} = \frac{c}{h(1 + c)}$. We draw the reaction term regarding the sign of \mathscr{A}_g and the s values (in Fig. 12).

In case of bistability, the wave is always pushed (Hadeler and Rothe 1975); we can dismiss condition (2) in the research of pulled monostable waves. We use criterion (10) on monostable cases, i.e. drive invasion (1) (4) (6), wild type invasion (3) (5) (8), and coexistence state (7) (the numbers refer to the subgraphs in Fig. 12).

E.2.1 Monostable drive invasion

(4) When $\mathcal{A}_g = 0 \iff h = \frac{1}{2}$ and $s < s_1 = s_{2,g} = \frac{2c}{c+1} \iff \frac{s}{2}(c+1) < c$ From equation (91), we have for all $p_D \in [0, 1]$:

$$\sigma(0) - (1 - p_{D})\sigma(p) = \left(c - \frac{s}{2}(c+1)\right) - \frac{(c - \frac{s}{2}(c+1))(1 - p_{D})}{1 - sp_{D}}$$

$$= \left(c - \frac{s}{2}(c+1)\right) \frac{(1 - s)p_{D}}{1 - sp_{D}} \ge 0,$$
(93)

where σ is the selection term defined by equation (8). Criterion (10) is verified.

6 When $\mathcal{A}_g > 0$ and $p_{\mathrm{D}g}^* > 1$ From equation (92), we have for all $p_{\mathrm{D}} \in [0, 1]$:

$$\sigma(0) - (1 - p_{D})\sigma(p) = \mathscr{A}_{g} p_{Dg}^{*} - \frac{-\mathscr{A}_{g} (p_{D} - p_{Dg}^{*}) (1 - p_{D})}{-\mathscr{A}_{g} p_{D}^{2} + (\mathscr{A}_{g} - s) p_{D} + 1}$$

$$= \mathscr{A}_{g} p_{D} \frac{-(\mathscr{A}_{g} p_{Dg}^{*} + 1) p_{D} + (\mathscr{A}_{g} + 1 - s) p_{Dg}^{*} + 1}{-\mathscr{A}_{g} p_{D}^{2} + (\mathscr{A}_{g} - s) p_{D} + 1}.$$
(94)



 $[\]overline{\int M'_g(p_{\mathrm{D}}) = \mathscr{A}_g(1 - 2p_{\mathrm{D}}) - s} \le 0 \text{ therefore } M_g(1) \le M_g(p_{\mathrm{D}}) \le M_g(0).$

30 Page 48 of 54 L. Kläy et al.

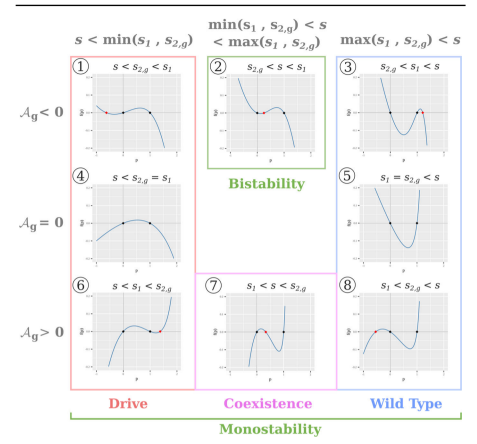


Fig. 12 Reaction term of equation (91) regarding the sign of $\mathscr{A}_g = s \ (1-2h)$ and the s values. The s threshold values are $s_1 = \frac{c}{1-h(1-c)}$ and $s_{2,g} = \frac{c}{2ch+h(1-c)} = \frac{c}{h(1+c)}$. The dots on the x axis correspond to the steady states: $p_D = 0$ and $p_D = 1$ in black, $p_{Dg}^* = \frac{1}{2} + \frac{2c \ (1-sh)-s}{2 \ \mathscr{A}_g}$ in red when it exists (color figure online)

Note that $-\mathscr{A}_g p_D^2 + (\mathscr{A}_g - s) p_D + 1 > (1 - s) > 0$ and $\mathscr{A}_g p_D > 0$. The affine term $-(\mathscr{A}_g p_{Dg}^* + 1) p_D + (\mathscr{A}_g + 1 - s) p_{Dg}^* + 1$ decreases with p_D . In order to show that it is positive for all $p_D \in [0, 1]$, we just need to verify that this it is true for $p_D = 1$:

$$-(\mathscr{A}_g p_{\mathrm{D}g}^* + 1) + (\mathscr{A}_g + 1 - s) \ p_{\mathrm{D}g}^* + 1 = (1 - s) \ p_{\mathrm{D}g}^* \ge 0 \quad \Rightarrow \quad \sigma(0) - (1 - p_{\mathrm{D}})\sigma(p)$$

$$\ge 0 \quad \forall p_{\mathrm{D}} \in [0, 1].$$
 (95)

Criterion (10) is verified.

1 When $\mathscr{A}_g < 0$ and $p^*_{\mathrm{D}g} < 0$ and $s < s_{2,g} \iff 0 < c - sh(1+c)$

We consider equation (94) with $-\mathscr{A}_g p_D^2 + (\mathscr{A}_g - s) p_D + 1 > 1 - s > 0$ and $\mathscr{A}_g p_D < 0$. The affine term $-(\mathscr{A}_g p_{Dg}^* + 1) p_D + (\mathscr{A}_g + 1 - s) p_{Dg}^* + 1$ decreases with p_D . In order to show that it is negative for all $p_D \in [0, 1]$, we introduce a condition



implying the negativity for p = 0:

$$(\mathcal{A}_g + 1 - s) \ p_{\mathrm{D}g}^* + 1 = \frac{(1 - 2sh)(\mathcal{A}_g + 2c(1 - sh) - s) + 2\mathcal{A}_g}{2\mathcal{A}_g} < 0$$
 (96)

$$\iff (1 - 2sh)(c - sh(c + 1)) + s(1 - 2h) > 0$$
 (97)

Criterion (10) is verified when condition (97) is true.

E.2.2 Monostable wild-type invasion

In case of a monostable wild-type invasion, we need to consider the wild-type proportion $p_W = 1 - p_D \in [0, 1]$ and rewrite the equation (91):

$$-\partial_{t} p_{W} + \partial_{xx}^{2} p_{W} = \frac{\left(-\mathscr{A}_{g} (1 - p_{W}) + \frac{1}{2} (\mathscr{A}_{g} - s) + c(1 - sh)\right) (1 - p_{W}) p_{W}}{-\mathscr{A}_{g} (1 - p_{W})^{2} + (\mathscr{A}_{g} - s)(1 - p_{W}) + 1}$$

$$\iff \partial_{t} p_{W} - \partial_{xx}^{2} p_{W} = \frac{\left(-\mathscr{A}_{g} p_{W} + \frac{1}{2} (\mathscr{A}_{g} + s) - c(1 - sh)\right) (1 - p_{W}) p_{W}}{-\mathscr{A}_{g} p_{W}^{2} + (\mathscr{A}_{g} + s) p_{W} + (1 - s)}$$
(98)

When $\mathcal{A}_g \neq 0$, equation (98) can be rewritten:

$$\partial_{t} p_{W} - \partial_{xx}^{2} p_{W} = \frac{-\mathscr{A}_{g} (p_{W} - p_{Wg}^{*}) (1 - p_{W}) p_{W}}{-\mathscr{A}_{g} p_{W}^{2} + (\mathscr{A}_{g} + s) p_{W} + (1 - s)} \text{ with}$$

$$p_{Wg}^{*} = \frac{1}{2} - \frac{2c (1 - sh) - s}{2\mathscr{A}_{g}} = 1 - p_{Dg}^{*}$$
(99)

(5) When $\mathcal{A}_g = 0 \iff h = \frac{1}{2}$ and $s_1 = s_{2,g} = \frac{2c}{c+1} < s \iff c < \frac{s}{2}$ (c+1) From equation (98) we have for all $p_W \in [0, 1]$:

$$\sigma(0) - (1 - p_{W})\sigma(p_{W}) = \left(\frac{s}{2}(c + 1) - c\right)\left(\frac{1}{1 - s} - \frac{1 - p_{W}}{sp_{W} + 1 - s}\right) \ge \frac{p_{W}\left(\frac{s}{2}(c + 1) - c\right)}{1 - s} \ge 0$$
(100)

Criterion (10) is verified.

(8) When $\mathcal{A}_g > 0$ and $p_{\mathrm{W}g}^* = 1 - p_{\mathrm{D}g}^* > 1$ From equation (99), we have for all $p_{\mathrm{W}} \in [0, 1]$:

$$\begin{split} \sigma(0) - (1 - p_{\mathrm{W}})\sigma(p_{\mathrm{W}}) &= \frac{\mathscr{A}_{g} \ p_{\mathrm{W}g}^{*}}{1 - s} - \frac{-\mathscr{A}_{g} \ (p_{\mathrm{W}} - p_{\mathrm{W}g}^{*}) \ (1 - p_{\mathrm{W}})}{-\mathscr{A}_{g} p_{\mathrm{W}}^{2} + (\mathscr{A}_{g} + s) \ p_{\mathrm{W}} + (1 - s)} \\ &= \mathscr{A}_{g} p_{\mathrm{W}} \ \frac{(-p_{\mathrm{W}g}^{*} + 1 - s) \ p_{\mathrm{W}} + p_{\mathrm{W}g}^{*} (\mathscr{A}_{g} + 1) + (1 - s)}{(1 - s)(-\mathscr{A}_{g} p_{\mathrm{W}}^{2} + (\mathscr{A}_{g} + s) \ p_{\mathrm{W}} + (1 - s))}. \end{split}$$

Note that $(1-s)(-\mathscr{A}_g \, p_{\rm W}^2 + (\mathscr{A}_g + s) \, p_{\rm W} + (1-s)) > (1-s)^2 > 0$ and $\mathscr{A}_g \, p_{\rm W} > 0$. As $p_{\rm W}^* \, g > 1$, the affine term $(-p_{\rm W}^* \, g + 1 - s) \, p_{\rm W} + p_{\rm W}^* \, (\mathscr{A}_g + 1) + (1-s)$ decreases



30 Page 50 of 54 L. Kläy et al.

with p_W . In order to show that it is positive for all $p_W \in [0, 1]$, we just need to verify that this it is true for $p_W = 1$:

$$(-p_{Wg}^* + 1 - s) + p_{Wg}^* (\mathcal{A}_g + 1) + (1 - s) = 2 (1 - s) + \mathcal{A}_g p_{Wg}^* \ge 0$$

$$\Rightarrow \sigma(0) - (1 - p_W) \sigma(p_W) \ge 0 \quad \forall p_W \in [0, 1]. \tag{102}$$

Criterion (10) is verified.

(3) When $\mathcal{A}_g < 0$ and $p_{W_g}^* = 1 - p_{D_z}^* < 0$

We consider equation (101) with $(1-s)(-\mathscr{A}_g p_W^2 + (\mathscr{A}_g + s) p_W + (1-s)) > (1-s)^2 > 0$ and $\mathscr{A}_g p_W < 0$. The affine term $(-p_{Wg}^* + 1 - s) p_W + p_{Wg}^* (\mathscr{A}_g + 1) + (1-s)$ is strictly positive for $p_W = 1$, therefore criterion (10) is not verified.

E.2.3 Monostable coexistence state

7 When $\mathcal{A}_g > 0$ and $0 < p_{D_g}^* = 1 - p_{W_g}^* < 1$

In the coexistence case, we have to verify that both sub-traveling waves, the drive invasion wave going to the right and the wild-type invasion wave going to the left, are pulled waves (see Fig. 3).

For the drive invasion wave we consider equation (94) with $0 < p_{\mathrm{D}g}^* < 1$ and $p_{\mathrm{D}} \in [0, p_{\mathrm{D}g}^*]$ (the term drive wave implies that the proportion of wild type increases after the wave passes; therefore the global stable steady state $p_{\mathrm{D}g}^*$ is also the maximum proportion). Once again, we need to prove that the affine term $-(\mathscr{A}_g p_{\mathrm{D}g}^* + 1) p_{\mathrm{D}} + (\mathscr{A}_g + 1 - s) p_{\mathrm{D}g}^* + 1$ is positive. As it decreases with $p_{\mathrm{D}} \in [0, p_{\mathrm{D}g}^*]$, we determine its sign for $p_{\mathrm{D}} = p_{\mathrm{D}g}^*$:

$$-(\mathscr{A}_{g}p_{\mathrm{D}g}^{*}+1) p_{\mathrm{D}g}^{*}+(\mathscr{A}_{g}+1-s) p_{\mathrm{D}g}^{*}+1=-\mathscr{A}_{g} (p_{\mathrm{D}g}^{*})^{2} +(\mathscr{A}_{g}-s) p_{\mathrm{D}g}^{*}+1\geq 1-s\geq 0$$

$$\Rightarrow \sigma(0)-(1-p_{\mathrm{D}})\sigma(p_{\mathrm{D}})\geq 0 \quad \forall p_{\mathrm{D}}\in[0,p_{\mathrm{D}g}^{*}].$$
(103)

Criterion (10) is verified for the drive wave.

For the wild-type invasion wave, we consider equation (101) with $0 < p_{\mathrm{W}g}^* = 1 - p_{\mathrm{D}g}^* < 1$ and $p_{\mathrm{W}} \in [0, p_{\mathrm{W}g}^*]$ (the term wild-type wave implies that the proportion of wild type increases after the wave passes; therefore the global stable steady state $p_{\mathrm{W}g}^*$ is also the maximum proportion). Once again, we need to prove that the affine term $(-p_{\mathrm{W}g}^* + 1 - s) \ p_{\mathrm{W}} + p_{\mathrm{W}g}^* (\mathscr{A}_g + 1) + (1 - s)$ is positive. As it decreases with $p_{\mathrm{W}} \in [0, p_{\mathrm{W}g}^*]$, we determine its sign for $p_{\mathrm{W}} = p_{\mathrm{W}g}^*$:

$$(-p_{\mathrm{W}g}^* + 1 - s) \ p_{\mathrm{W}g}^* + (\mathcal{A}_g + 1) \ p_{\mathrm{W}g}^* + (1 - s) = -(p_{\mathrm{W}g}^*)^2 + (\mathcal{A}_g + 2 - s) \ p_{\mathrm{W}g}^* + 1 - s$$

$$\geq \min(1, \mathcal{A}_g + 2(1 - s)) \geq 0 \quad \Rightarrow \quad \sigma(0) - (1 - p_{\mathrm{W}})\sigma(p_{\mathrm{W}}) \geq 0 \quad \forall p_{\mathrm{W}} \in [0, p_{\mathrm{W}g}^*].$$

$$(104)$$

Criterion (10) is verified for the wild-type wave.



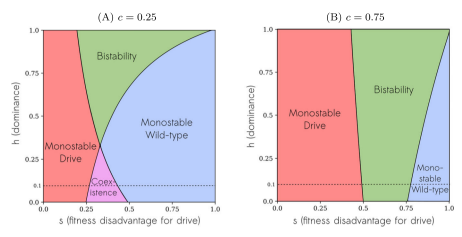
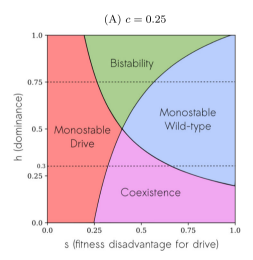


Fig. 13 Effect of fitness disadvantage (s) and dominance coefficient (h) on drive dynamics for system (4) (when conversion occurs in the zygote) when $r=+\infty$. Parameters for Fig. 4 (c=0.25 and h=0.1) and Fig. 5 (c=0.75 and h=0.1) are materialized by dotted lines

Fig. 14 Effect of fitness disadvantage (s) and dominance coefficient (h) on drive dynamics for system (5) (when conversion occurs in the germline) when $r=+\infty$. Parameters for Fig. 6A (c=0.25 and h=0.3) and Fig. 6B (c=0.25 and h=0.75) are materialized by dotted lines



F Heatmap supplementary materials

F.1 Effect of fitness disadvantage (s) and dominance coefficient (h) on drive dynamics, for ${\bf r}=+\infty$

Below, we compute heatmaps indicating the stability regime of systems (4) and (5) when $r = +\infty$, depending on the values of (h, s) and for a fixed value of c.

Such a figure has already been computed in Rode et al. (2019) for c = 0.85, and conversion occurring in the germline.



30 Page 52 of 54 L. Kläy et al.

F.2 Heatmap lines

F.2.1 Pure drive line

Consider model (6) with $n_{\text{WW}} = 0$. A well-mixed population containing only drive homozygous individuals will persist in the environment if its equilibrium state n_{DD}^* is strictly positive, i.e. if:

$$n_{\text{DD}}^* = \min\left(0, 1 - \frac{s}{r(1-s)}\right) > 0 \iff r > \frac{s}{1-s}$$
 (105)

In case of partial conversion, calculations give the same threshold (consider models (4) and (5) with $n_{\text{DW}} = 0$ and $n_{\text{WW}} = 0$).

F.2.2 Composite persistence line

Similarly, in case of coexistence, a well-mixed population will persist in the environment only if its equilibrium state n^* is strictly positive. Using Mathematica, we compute this population density equilibrium when conversion occurs in the zygote (n_7^*) or in the germline (n_p^*) based on systems (21) and (29). We obtain the following:

$$n_z^* = \min\left(0, 1 - \frac{1 - M_z(p_{\mathrm{D}z}^*)}{rM_z(p_{\mathrm{D}z}^*)}\right) \quad \text{and} \quad n_g^* = \min\left(0, 1 - \frac{1 - M_g(p_{\mathrm{D}g}^*)}{rM_g(p_{\mathrm{D}g}^*)}\right), \quad (106)$$

where the mean fitness M_z and M_g (already defined in Appendix E) are given by:

$$M_z(p_D) = -\mathcal{A}_z p_D^2 + (\mathcal{A}_z - s) p_D + 1$$
 and $M_g(p_D) = -\mathcal{A}_g p_D^2 + (\mathcal{A}_g - s) p_D + 1$, (107)

and the proportions $p_{\mathrm{D}z}^*$ and $p_{\mathrm{D}g}^*$ (already defined in Appendix E) are given by:

$$p_{\mathrm{D}z}^* = \frac{1}{2} + \frac{2c(1-s)-s}{2\mathscr{A}_z}$$
 and $p_{\mathrm{D}g}^* := \frac{1}{2} + \frac{2c(1-sh)-s}{2\mathscr{A}_g}$ (108)

with,

$$\mathcal{A}_z = s \left[2(1-c)(1-h) - 1 \right] \text{ and } \mathcal{A}_g = s (1-2h).$$
 (109)

Finally, the threshold values for r are given by:

$$n_z^* > 0 \iff r > \frac{1 - M_z(p_{\rm Dz}^*)}{M_z(p_{\rm Dz}^*)}$$
 (110)

when conversion occurs in the zygote and,

$$n_g^* > 0 \iff r > \frac{1 - M_g(p_{D_g}^*)}{M_g(p_{D_g}^*)}$$
 (111)

when conversion occurs in the germline.



References

- Alphey LS et al (2020) Standardizing the definition of gene drive. Proc Natl Acad Sci 117.49:30864–30867. https://doi.org/10.1073/pnas.2020417117
- Andrew H et al (2015) A CRISPR-Cas9 gene drive system targeting female reproduction in the malaria mosquito vector anopheles gambiae. Nat Biotechnol. https://doi.org/10.1038/nbt.3439
- An J, Henderson C, Ryzhik L (2021) Pushed, pulled and pushmi-pullyu fronts of the Burgers-FKPP equation. https://doi.org/10.48550/arXiv.2108.07861. arXiv: 2108.07861 [math]
- An J, Henderson C, Ryzhik L (2022) Quantitative steepness, semi-FKPP reactions, and pushmi-pullyu fronts. https://doi.org/10.48550/arXiv.2208.02880. arXiv: 2208.02880 [math]
- Aronson DG, Weinberger HF (1975) Nonlinear diffusion in population genetics, combustion, and nerve pulse propagation. In: Goldstein JA (ed) Partial differential equations and related topics. Lecture Notes in Mathematics. Springer, Berlin, pp 5–49. https://doi.org/10.1007/BFb0070595
- Aronson DG, Weinberger Hans F (1978) Multidimensional nonlinear diffusion arising in population genetics. Adv Math 30(1):33–76. https://doi.org/10.1016/0001-8708(78)90130-5
- Avery M, Holzer M, Scheel A(2022) Pushed-to-pulled front transitions: continuation, speed scalings, and hidden monotonicity. https://doi.org/10.48550/arXiv.2206.09989. arXiv: 2206.09989 [nlin]
- Barton NH (1979) The dynamics of hybrid zones. Heredity 43(3):341–359. https://doi.org/10.1038/hdy. 1979.87
- Beaghton A, Beaghton PJ, Burt A (2016) Gene drive through a landscape: reaction-diffusion models of population suppression and elimination by a sex ratio distorter. Theor Popul Biol 108:51–69. https://doi.org/10.1016/j.tpb.2015.11.005
- Buchman A et al (2020) Broad dengue neutralization in mosquitoes expressing an engineered antibody. PLoS Pathog 16(1):e1008103. https://doi.org/10.1371/journal.ppat.1008103
- Burt A (2003) Site-specific selfish genes as tools for the control and genetic engineering of natural populations. Proc R Soc B: Biol Sci 270(1518):921–928. https://doi.org/10.1098/rspb.2002.2319
- Champer J et al (2021) Suppression gene drive in continuous space can result in unstable persistence of both drive and wild-type alleles. Mole Ecol 30(4):1086–1101. https://doi.org/10.1111/mec.15788
- Committee on Gene Drive Research in Non-Human Organisms: Recommendations for Responsible Conduct et al. (2016) Gene drives on the horizon: advancing science, navigating uncertainty, and aligning research with public values. Washington (DC): National Academies Press (US)
- Dahirel M et al (2021) Shifts from pulled to pushed range expansions caused by reduction of landscape connectivity. Oikos 130(5):708–724. https://doi.org/10.1111/oik.08278
- Deredec A, Burt A, Godfray HCJ (2008) The population genetics of using homing endonuclease genes in vector and pest management. Genetics 179(4):2013–2026. https://doi.org/10.1534/genetics.108.089037
- Esvelt KM et al (2014) Concerning RNA-guided gene drives for the alteration of wild populations. Elife. https://doi.org/10.7554/eLife.03401
- Gabriel B, Oskar H, Korolev Kirill S (2018) Fluctuations uncover a distinct class of traveling waves. Proc Natl Acad Sci 115(16):E3645–E3654. https://doi.org/10.1073/pnas.1715737115
- Gantz V et al (2015) Highly efficient Cas9-mediated gene drive for population modification of the malaria vector mosquito anopheles stephensi. Proc Natl Acad Sci. https://doi.org/10.1073/pnas.1521077112
- Gantz VM, Bier E (2015) The mutagenic chain reaction: a method for converting heterozygous to homozygous mutations. Science 348(6233):442–444. https://doi.org/10.1126/science.aaa5945
- Grunwald HA et al (2019) Super-mendelian inheritance mediated by CRISPR-Cas9 in the female mouse germline. Nature 566(7742):105–109. https://doi.org/10.1038/s41586-019-0875-2
- Hadeler KP, Rothe F (1975) Travelling fronts in nonlinear diffusion equations. J Math Biol 2(3):251–263. https://doi.org/10.1007/BF00277154
- Hidenori T, Stone Howard A, Nelson David R (2017) Spatial gene drives and pushed genetic waves. Proc Natl Acad Sci USA 114(32):8452–8457. https://doi.org/10.1073/pnas.1705868114
- Holzer M (2022) Personal communication, conference "Parabolic and Kinetic Models in Population Dynamics", Labex CIMI, Toulouse. https://indico.math.cnrs.fr/event/7589/
- Kyrou K et al (2018) A CRISPR-Cas9 gene drive targeting doublesex causes complete population suppression in caged anopheles gambiae mosquitoes. Nat Biotechnol 36(11):1062–1066. https://doi.org/10.1038/nbt.4245
- Léo G, Florence D (2021) Demographic feedbacks can hamper the spatial spread of a gene drive. J Math Biol 83(6):67. https://doi.org/10.1007/s00285-021-01702-2



30 Page 54 of 54 L. Kläy et al.

Li J et al (2020) Can CRISPR gene drive work in pest and beneficial haplodiploid species? Evol Appl 13(9):2392–2403. https://doi.org/10.1111/eva.13032

- Liu Y, Champer J (2022) Modelling homing suppression gene drive in haplodiploid organisms. Proc R Soc B Biol Sci 289(1972):20220320. https://doi.org/10.1098/rspb.2022.0320
- Male Bias and Female Fertility (2022) Target Malaria. https://targetmalaria.org/what-wedo/our-approach/male-bias-and-female-fertility/ (visited on 02/28/2022)
- Martin J et al (2012) A programmable dual RNA-guided DNA endonuclease in adaptive bacterial immunity. Science 337(6096):816–821. https://doi.org/10.1126/science.1225829
- Martin S, Nicolas V (2016) Reduction to a single closed equation for 2-by-2 reaction- diffusion systems of Lotka–Volterra type. SIAM J Appl Math 76(5):2060–2080. https://doi.org/10.1137/16M1059217
- Michael T, Barton Nicholas H (2017) Deploying dengue-suppressing Wolbachia: robust models predict slow but effective spatial spread in aedes aegypti. Theor Popul Biol 115:45–60. https://doi.org/10.1016/j.tpb.2017.03.003
- Nadin G, Strugarek M, Vauchelet N (2018) Hindrances to bistable front propagation: application to Wolbachia invasion. J Math Biol 76(6):1489–1533. https://doi.org/10.1007/s00285-017-1181-y
- Neve P (2018) Gene drive systems: do they have a place in agricultural weed management? Pest Manage Sci 74(12):2671–2679. https://doi.org/10.1002/ps.5137
- North AR, Burt A, Godfray H, Charles J (2020) Modelling the suppression of a malaria vector using a CRISPR-Cas9 gene drive to reduce female fertility. BMC Biol 18:98. https://doi.org/10.1186/s12915-020-00834-z
- Otto SP, Day T (2011) A biologist's guide to mathematical modeling in ecology and evolution. Princeton University Press, https://doi.org/10.1515/9781400840915
- Peischl S et al (2013) On the accumulation of deleterious mutations during range expansions. Mole Ecol 22(24):5972–5982. https://doi.org/10.1111/mec.12524
- Rode NO et al (2019) Population management using gene drive: molecular design, models of spread dynamics and assessment of ecological risks. Conserv Genet 20(4):671–690. https://doi.org/10.1007/ s10592-019-01165-5
- Roques L et al (2012) Allee effect promotes diversity in traveling waves of colonization. Proc Natl Acad Sci 109(23):8828–8833. https://doi.org/10.1073/pnas.1201695109
- Sumit D, Lloyd Alun L, Fred G (2020) Gene drive dynamics in natural populations: the importance of density dependence, space, and sex. Ann Rev Ecol Evol Syst 51(1):505–531. https://doi.org/10.1146/annurev-ecolsys-031120-101013
- Unckless RL et al (2015) Modeling the manipulation of natural populations by the mutagenic chain reaction. Genetics 201(2):425–431. https://doi.org/10.1534/genetics.115.177592
- Zhou J et al (2019) Critical traveling waves in a diffusive disease model. J Math Anal Appl 476(2):522–538. https://doi.org/10.1016/j.jmaa.2019.03.066

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Springer Nature or its licensor (e.g. a society or other partner) holds exclusive rights to this article under a publishing agreement with the author(s) or other rightsholder(s); author self-archiving of the accepted manuscript version of this article is solely governed by the terms of such publishing agreement and applicable law

