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[Siirry ensimmäiseen muutokseen \(sivu 2\)](#)

Positive fitness effects help explain the broad range of *Wolbachia* prevalences in natural populations

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Article

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Abstract

20 The bacterial endosymbiont *Wolbachia* is best known for its ability to modify its host's
reproduction by inducing cytoplasmic incompatibility (CI) to facilitate its own spread.
22 Classical models predict either near-fixation of costly *Wolbachia* once the symbiont has
overcome a threshold frequency (invasion barrier), or *Wolbachia* extinction if the barrier
24 is not overcome. However, natural populations do not all follow this pattern: *Wol-*
bachia can also be found at low frequencies (below one half) that appear stable over
26 time. *Wolbachia* is known to have pleiotropic fitness effects (beyond CI) on its hosts.
Existing models typically focus on the possibility that these are negative. Here we con-
28 sider the possibility that the symbiont provides direct benefits to infected females (e.g.
resistance to pathogens) in addition to CI. We discuss an underappreciated feature of
30 *Wolbachia* dynamics: that CI with additional fitness benefits can produce low-frequency
($< 1/2$) stable equilibria. Additionally, without a direct positive fitness effect, any stable
32 equilibrium close to one half will be sensitive to perturbations, which make such equi-
libria unlikely to sustain in nature. The results hold for both diplodiploid and different
34 haplodiploid versions of CI. We suggest that insect populations showing low-frequency
Wolbachia infection might host CI-inducing symbiotic strains providing additional (hid-
36 den or known) benefits to their hosts, especially when classical explanations (ongoing
invasion, source-sink dynamics) have been ruled out.

38 Introduction

Maternally inherited endosymbiotic bacteria of genus *Wolbachia* are widespread in arthro-
40 pods and nematodes (Hilgenboecker et al., 2008; Zug and Hammerstein, 2012; Weinert
et al., 2015). Since the discovery of *Wolbachia*-induced cytoplasmic incompatibility (CI,
42 Laven, 1956; Yen and Barr, 1971) – embryonic mortality in the offspring of parents of
different infection status – the symbionts have attained considerable empirical as well as
44 theoretical interest. CI is an obvious and dramatic fitness effect, but *Wolbachia*'s effects
are often pleiotropic, impacting the fitness of its hosts directly and regardless of the type
46 of mates that the host reproduces with (for example Hoffmann et al., 1990). Assuming
that pleiotropy includes direct costs of infection, it becomes at first sight difficult to un-
48 derstand how *Wolbachia* can spread and reach high frequencies. However, theory soon
showed that costly *Wolbachia* infections show positive **frequency dependence**. At low fre-
50 quencies, costs predominate and *Wolbachia* cannot spread, but if the initial frequency of
Wolbachia is sufficiently high (above an invasion barrier), then the frequency-dependent
52 success favours the CI-inducing symbiont sufficiently for it to spread (Fine, 1978; Cas-
pari and Watson, 1959). At equilibrium it will be close to fixation, only prevented from
54 fixing by some infections failing to be passed on to offspring (Hoffmann et al., 1990; En-
gelstädter and Telschow, 2009). This fits in with the general expectation that positive
56 frequency dependence makes it easy to explain either absence or (near) fixation of a trait
(Lehtonen and Kokko, 2012).

58 In many empirical systems, *Wolbachia* are indeed very prevalent in their host popu-
lations (Hoffmann et al., 1990; Jeong et al., 2009; Duplouy and Brattström, 2018; Deng
60 et al., 2021) in line with existing theory. However, the recent accumulation of studies
on the prevalence and penetrance of *Wolbachia* in diverse host species and populations
62 show a wider range of infection prevalences. Sazama et al. (2019) have reported that
infection prevalence of *Wolbachia* is likely to remain below one half in the majority of

64 insect species (Tagami and Miura, 2004; Sun et al., 2007; Arthofer et al., 2009; Hughes
et al., 2011), including systems where the spread of *Wolbachia* is potentially ongoing
66 (Duploux et al., 2010). There are also examples of low prevalence remaining reasonably
stable over time (Duploux et al., 2015). Although some cases of low and apparently
68 stable prevalence might be explicable as a result of migration to sink populations from
sources where symbiont prevalence is high (Flor et al., 2007; Telschow et al., 2007), it
70 appears difficult to consider spatial dynamics a sufficient explanation for widespread
cases of empirically documented low prevalences.

72 Here we show that spatial heterogeneity is not needed for models to produce low stable
frequencies of CI-inducing symbionts, once one allows the effects of infection on the host
74 to be positive. Studies allowing positive fitness effects of CI-inducing symbionts are
rare. However, Zug and Hammerstein (2018) investigated the effect of positive fitness
76 on the invasion dynamics of CI and other types of reproductive parasites into virgin
populations and against each other. They demonstrated the key role of relative fitness
78 in determining invasion conditions of the parasites. The effect is often observed as a
lower (or absent) invasion threshold frequency. While their focus was mostly on the
80 invasibility of the parasites, we aim to widen the understanding of the resulting stable
equilibrium frequencies when positive fitness effects are present.

82 We first revisit classic CI models (following the notation of Engelstädter and Telschow,
2009) and show that the assumption structure, where infected hosts experience either no
84 or negative fitness effects, cannot equilibrate at infection prevalences below one half (as
stated by Turelli, 1994, for a diploid case). We also argue that within the range
86 predicted by the classic model (between one half and one), the lower end of the range
is difficult to maintain in the presence of stochastic fluctuations in infection prevalence,
88 as the invasion barrier will in these cases be too close to the stable equilibrium. We
then show that when we allow infected hosts to benefit from their infection status, the

90 **stable** equilibria can exist below one half in a wide range of cases, including situations
with and without **an** invasion barrier. The results of the classical diplodiploid model of
92 CI are quite similar to **the results of** different haplodiploid models (with an exception
that we discuss). Our observations of low frequency stable equilibria (specifically, below
94 one half) are not entirely novel: the analysis of Zug and Hammerstein (2018) includes
a demonstration of such a situation in a diplodiploid system (their Supplementary Fig.
96 S1), albeit without any written comment as the focus of these authors was elsewhere.
Hence, our aim is to strengthen the appreciation of the dynamic consequences of positive
98 fitness effects in the dynamics of CI-inducing symbionts.

Models and analysis

100 **Diplodiploid model**

Hoffmann et al. (1990) presented a seminal model for the spread of CI-inducing symbionts
102 in diploid-diploid species, building upon a slightly different model of Fine (1978). We
use the discrete time model of Hoffmann et al. (1990) but update it to the notation
104 of Engelstädter and Telschow (2009) (Fig. 1). *Wolbachia*-bearing females have relative
fecundity f compared to the uninfected baseline (which we set to unity). *Wolbachia* is
106 transmitted maternally to a proportion t of eggs. Under rules of CI, an uninfected egg
that is fertilized by a *Wolbachia*-modified sperm (produced by an infected male) dies
108 with probability L , while *Wolbachia*-infected eggs survive irrespective of the status of
the fertilizing sperm (to be precise, this requires sperm to be either from a *Wolbachia*-
110 free father or from a father with the same or a compatible *Wolbachia* strain as the egg;
for simplicity we follow earlier studies and consider only one strain in our model).

112 Our model assumes equal frequency (p) of *Wolbachia* in males and females in the
current generation. We track four types of matings (the mother and the father can each

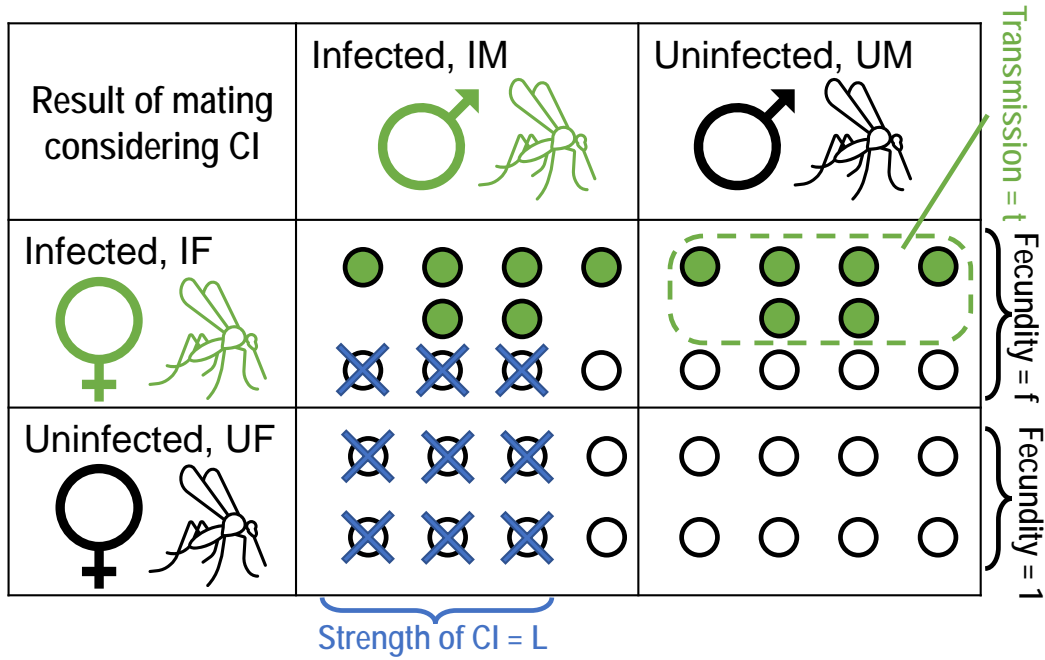


Figure 1: Mating results and key parameters of reproduction when insect population contains *Wolbachia* infected individuals. When an infected male (IM) fertilizes eggs, the *Wolbachia*-modified sperm leads to cytoplasmic incompatibility in all *Wolbachia*-free eggs, regardless of the infection status of their mother (IF, UF). Uninfected males (UM) fertilize all eggs successfully. Note that in the diploid case, the incompatible eggs (that die) could have developed as either males or females. In haplodiploids, where fertilization produces females, the incompatible eggs would all be originally female. In the female-killing type they simply die, while in the masculinization type they develop as males instead.

114 be uninfected or infected) and their outcome in terms of infected and uninfected offspring
 production. Only infected mothers can produce infected offspring, and the total of such
 116 offspring is pft (sperm genotype does not matter for these mothers). The denominator
 consists of all offspring from all matings. The change from the current generation to the
 118 next is

$$\begin{aligned}
 \Delta p &= \frac{pft}{\underbrace{(1-p)^2}_{UF \times UM} + \underbrace{(1-p)p(1-L)}_{UF \times IM} + \underbrace{pf(1-p)}_{IF \times UM} + \underbrace{p^2f(t + (1-t)(1-L))}_{IF \times IM}} - p \\
 &= \frac{pft}{1 - p(1-f) - pL(1-p(1-f+ft))} - p,
 \end{aligned} \tag{1}$$

120 where the first term gives the infection frequency in the next generation. Equation (1)
 has three equilibria:

$$\hat{p}_0 = 0, \hat{p}_1 = \frac{-B - \sqrt{B^2 - 4AC}}{2A}, \text{ and } \hat{p}_2 = \frac{-B + \sqrt{B^2 - 4AC}}{2A},$$

122 where $A = L(1 - f(1 - t))$, $B = f - 1 - L$ and $C = 1 - ft$ (Fig. 2a). As shown by
 Hoffmann et al. (1990), when all three equilibria exist, \hat{p}_0 and \hat{p}_2 are stable while \hat{p}_1
 124 between them is unstable and gives the invasion threshold (in appendix B we rederive
 this result extending it to $f > 1$). Turelli (1994, Eq. 4) and Turelli and Hoffmann (1995,
 126 Eq. 3) indicated that $\hat{p}_2 > 1/2$, although they did not provide an explicit proof for that.
 Below, we show what is required for this result to hold.

128 Before proceeding to understand effects caused by $f > 1$, it is useful to understand
 why the classical result arises, i.e. that $f \leq 1$ predicts the absence of low ($< 1/2$)
 130 and stable frequencies. We outline the reasons both mathematically and biologically.
 Mathematically, \hat{p}_1 and \hat{p}_2 are symmetric around a point $r = -B/(2A)$. Now, for \hat{p}_2 to
 132 remain below $1/2$, it is necessary (but not sufficient) that r is below $1/2$. Solving the
 inequality $r < 1/2$ for f with $0 < t \leq 1$, $0 < L \leq 1$ yields

$$\frac{1 - f + L}{2L(1 - f(1 - t))} < \frac{1}{2} \tag{2}$$

$$\Leftrightarrow f > \frac{1}{1 - L(1 - t)} \geq 1,$$

134 which clearly implies that $r > 1/2$ for any $f \leq 1$. However, the result also suggests
 136 that permitting $f > 1$ can change the results, since $r < 1/2$ becomes possible and thus
 possibly also $\hat{p}_2 < 1/2$. A numerical example shows that this indeed happens with
 138 suitable parameter values (Fig. 2b, c, d).

It is worthwhile to consider the robustness of the equilibria against stochastic variation
 140 in the environment, that may perturb populations away from their current equilibria.

We show that this makes equilibria close to one half unlikely to persist in the long term,
 142 within the classically assumed parameter regime where $f \leq 1$. Consider a situation
 where $f \leq 1$ and the stable equilibrium \hat{p}_2 is close to **one half**. We have a situation $\hat{p}_1 <$
 144 $r < \hat{p}_2$, but **given** the assumption $f \leq 1$ we also know that $r > 1/2$. Since we assumed \hat{p}_2
 is close to one half, it follows that all points \hat{p}_1 , r and \hat{p}_2 must be close to each other (since
 146 r is located exactly midway between \hat{p}_1 and \hat{p}_2). **Hence**, if the stable equilibrium \hat{p}_2 is
 close to one half, then the unstable equilibrium \hat{p}_1 (representing the invasion barrier) is
 148 not far below \hat{p}_2 . Under such circumstances, random fluctuations in infection prevalence
 can easily make the population fall stochastically under the invasion barrier, from where
 150 it deterministically proceeds towards extinction (unless another stochastic event brings
 it back up to above the invasion barrier).

152 Biologically, the above analysis implies that if *Wolbachia* infection has effects that
 go beyond CI, then these additional fitness differences between infected and uninfected
 154 hosts can change the outcome of the interaction significantly. The direction of these
 differences determine whether *Wolbachia* frequencies can only stabilize at high values
 156 (with direct negative fitness effect), or whether the range of prevalence values extends
 to include low frequencies.

158 To develop biological intuition for why this is the case, it is instructive to see why
 the invasion threshold exists in the classical situation ($f \leq 1$), i.e. why in **Fig. 2a**
 160 there is a switch from poor spread of *Wolbachia* at low frequencies to better success
 at moderate frequencies. The impact of *Wolbachia* on uninfected eggs is captured by
 parameter L , but the value of L becomes irrelevant as a determinant of the dynamics
 when the infection is rare (very low p). **A Taylor series expansion of Eq. (1) around**
 164 **$p = 0$ is $(ft - 1)p + f(1 - f + L)tp^2 + \mathcal{O}[p]^3$, showing that parameter L is associated**
with second or higher order terms relative to p . Hence, when p is low, the terms that
 166 **include L become negligible and $\Delta p \approx (ft - 1)p$.**

For the infection to increase in frequency when it is rare, the requirement is that
168 $\Delta p > 0$, creating a simple condition: $ft > 1$ means that the 'leakage' of offspring
into an uninfected state (i.e., only a proportion t of offspring of *Wolbachia*-infected
170 mothers inherit the infection) is more than compensated for by the fecundity advantage
of *Wolbachia*-infected mothers. Assuming this is the case, the equilibrium \hat{p}_0 is unstable
172 and *Wolbachia* can spread from rare. Hence there is no invasion threshold when $ft > 1$
(See also Appendix B). Simultaneously, fixation is not possible if we assume $t < 1$; the
174 'leakage' always maintains a supply of uninfected individuals.

Figure 2a shows an example where ft is below 1 and spread from rare is consequently
176 not possible. When there are not enough males causing the 'competing' egg type to
fail (very low p), the combination of low fecundity ($f < 1$) and imperfect transmission
178 ($t < 1$) together prevent CI-inducing *Wolbachia* from maintaining its frequency in the
population. At higher p , we can no longer ignore the consequences of CI, and therefore
180 the effect of the parameter L . Now, a high proportion of infected males will elevate
the relative success of infected eggs compared to uninfected competitors; the success
182 difference arises from uninfected females' eggs dying. This explains why in classical
examples the CI-inducing symbiont can begin to spread at some threshold value of p :
184 the spread is caused by sufficiently many infected males harming the reproduction of
uninfected individuals.

186 The central role of the product ft provides important insight. Its value exceeds 1
when $f > 1/t$, thus sufficiently high values of f can make the invasion barrier disappear
188 (as stated e.g. by Zug and Hammerstein, 2018). However, this can never happen for any
 $f \leq 1$, since $1/t \geq 1$ for any $t \in (0, 1]$. Thus $f > 1$ is a necessary (but not sufficient)
190 requirement for the invasion barrier to disappear. Note that the above holds assuming
that the stable polymorphic equilibrium exists but it is possible to lose both the threshold
192 and the stable equilibrium also with $ft < 1$ as seen in Fig. 2. Biologically, situations of

$f > 1/t > 1$ are possible, once one considers the full range of possibilities for pleiotropy.
194 If females infected by CI-inducing *Wolbachia* are more fecund than uninfected females,
and if this advantage is sufficiently large, then the invasion barrier disappears. In this
196 case, one might at first sight expect that this additional advantage would simply make
Wolbachia reach fixation. This is not the case, however, since the effect of $t < 1$ causing
198 fixation to be impossible still holds.

Instead, the more interesting finding is that $f > 1$ creates conditions where a low
200 frequency of *Wolbachia* can be stable. This was not possible when $f \leq 1$, because
infected males at low prevalence do not cause a large enough survival difference between
202 eggs. But if $f > 1$, infected females can outcompete uninfected ones with very little
(assuming $ft < 1$) or no ($ft > 1$) CI-related advantages, and as a whole, the infection
204 can spread at values of L that are so low that they would never make *Wolbachia* spread
in situations involving $f \leq 1$ (compare Fig. 2c to Fig. 2e). In other words, the
206 latter condition would not feature an invasion barrier above which positive frequency
dependence would be sufficiently strong to bring *Wolbachia* to higher frequencies.

208 If we now assume that $f > 1/t$, we know that the invasion barrier is absent since
infected females are more fecund than uninfected ones (as the above implies that $f > 1$).
210 Once the infection spreads, so that infected males are no longer negligibly rare, infected
females reap additional benefits (indirectly, by the effect of L killing their competitors).
212 The remaining equilibrium in the $0 < p < 1$ range is stable. Even when $L = 0$, i.e.
the effect of males on eggs is switched off and the aforementioned additional benefits
214 are absent, qualitatively similar dynamics arise: the infection frequency increases from
rare but cannot fix due to imperfect transmission (also shown in Fig. S1 of Zug and
216 Hammerstein, 2018).

Figures 2c and 2d show examples with $ft > 1$ and five values of L . As expected,
218 the $L = 0$ case (lowest curve) exhibits simple dynamics, with the infected strain in-

creasing until imperfect transmission leads to a stable equilibrium which, by solving
 220 $\Delta p = \hat{p}ft / (\hat{p}f + (1 - \hat{p})) - \hat{p} = 0$, is at $\hat{p} = (1 - ft) / (1 - f) \approx 0.06$ with the given numer-
 ical example. Higher values of L lead to equilibria with a higher prevalence of *Wolbachia*
 222 (Figs. 2 and 3). Choosing equivalent values for L in a setting where $ft < 1$ always leads
 to lower growth (all graphs are lower in Fig. 2e, f compared with the equivalent graphs
 224 in Fig. 2c, d). At high L this means introducing an invasion barrier, while at low L the
 result is negative growth regardless of the current value of p , i.e. extinction of *Wolbachia*
 226 (three lowest curves in Fig. 2e).

It is worth clarifying the statement that introducing benefits can allow for a lower
 228 equilibrium frequency of *Wolbachia*, as it is counter-intuitive at first sight. The meaning
 of the statement is not that higher f lowers the equilibrium frequency when other param-
 230 eters are kept constant; this does not happen, instead, higher f increases the frequency
 (Fig. 3). The statement refers to the fact that this very effect (high f improves the
 232 prospects for *Wolbachia*) can, under parameter settings that are a priori unfavourable
 to *Wolbachia*, shift a situation where no *Wolbachia* persist to one where some spread is
 234 possible, and the system finds its equilibrium at low p (the light-coloured curves in Fig.
 2c compared to Fig. 2e; Fig. 3). The interesting findings of equilibria with $p < 1/2$ arise
 236 because $ft > 1$ implies no invasion barrier irrespective of the value of L , thus *Wolbachia*
 can spread from rare even if L is meagre, but the modest value of L then gives very little
 238 ‘boost’ at higher values of p (meagre male effects on uninfected eggs when L is low).

Although our analysis so far appears to suggest that cases with invasion barrier asso-
 240 ciate with high equilibrium frequencies, it should be noted that examining the full range
 of possibilities permitted by $f > 1$ includes cases where an invasion barrier exists, and
 242 the stable equilibrium is below one half. An invasion threshold exists if $ft < 1$ (assum-
 ing existence of the stable polymorphic equilibrium), while the necessary condition for
 244 $\hat{p}_2 < 1/2$ is that $f > 1$. Suitably chosen values for f and t can fulfil both criteria simul-

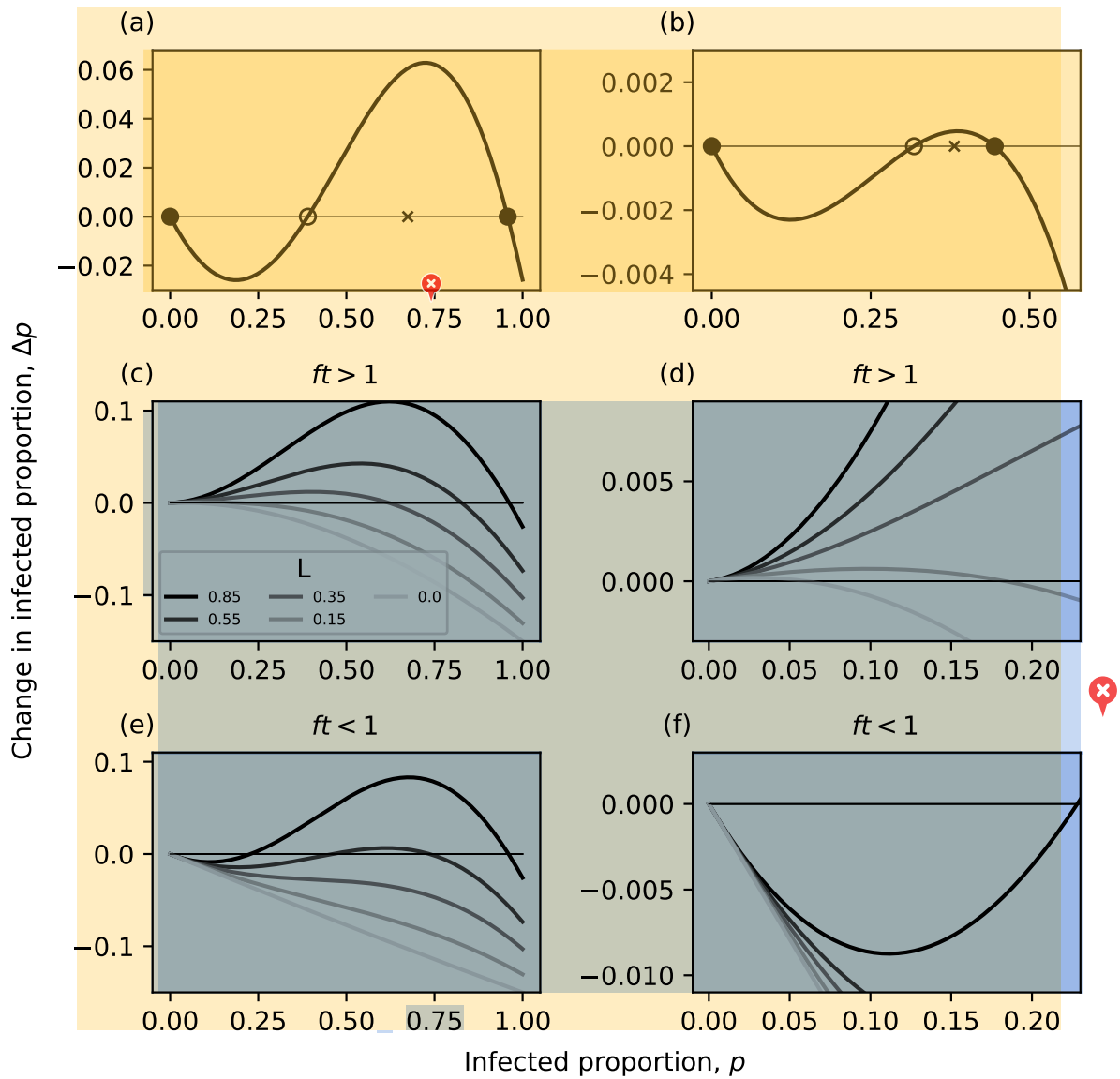


Figure 2: Equilibria of the diploid model (Eq. (1)) shown with plots of Δp as a function of p . Dots: stable equilibria, circle: unstable equilibrium, cross: point r . a) Classical example with an invasion threshold and high-frequency stable equilibrium. Parameters $f = 0.85$, $t = 0.85$, $L = 0.85$. b) It is possible to have a low stable frequency ($p < 1/2$) together with an invasion threshold ($ft = 0.9588 < 1$). ($f = 1.128$, $t = 0.85$, $L = 0.35$.) Panels c), d) and e), f) contrast the effect of the strength of CI (L) on the invasion dynamics when $ft > 1$ or $ft < 1$. In c),d) $f = 1.19$, $t = 0.85$, and no invasion threshold exists. Low value of L leads to low stable infection frequency. In e),f) $f = 0.99$, $t = 0.85$. High L shows the classic case, while low L predicts extinction of *Wolbachia*.

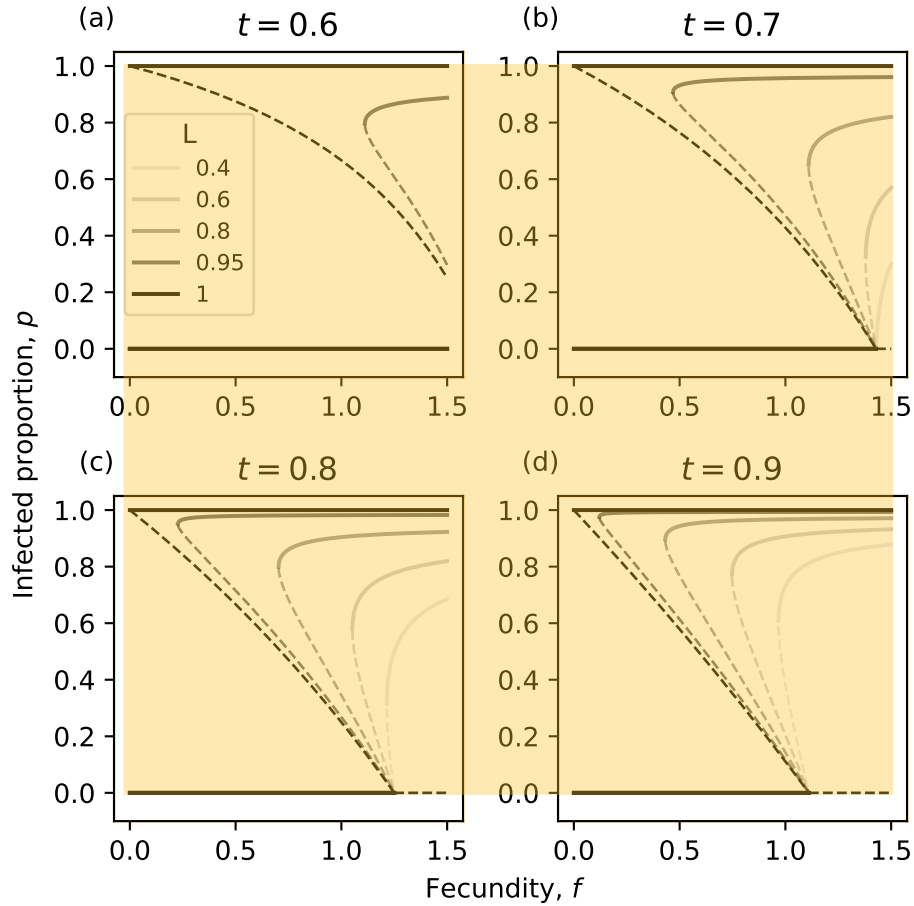


Figure 3: Equilibria of the diploid model (Eq. (1)) as function of f , for different values of t (panel titles) and L (colours of the lines, see legend). Solid lines: stable equilibria. Dashed lines: unstable equilibria.

taneously (example: Fig. 2b). As a summary, increasing f will increase the value of the stable equilibrium, but at the same time open possibilities for less successful symbionts, manifested as lower values of L and t , that will equilibrate at lower frequencies. The effect of parameter f with different combinations of t and L is shown in Fig. 3.

It is of interest to see if equilibria with low p can only exist at very low values of L (mild incompatibility) as in the above examples, or whether other values of other parameters allow low- p equilibria to exist even if L is relatively high. The latter proves to be the case, and as a whole, assuming $f > 1$ reveals a richer set of possible outcomes.

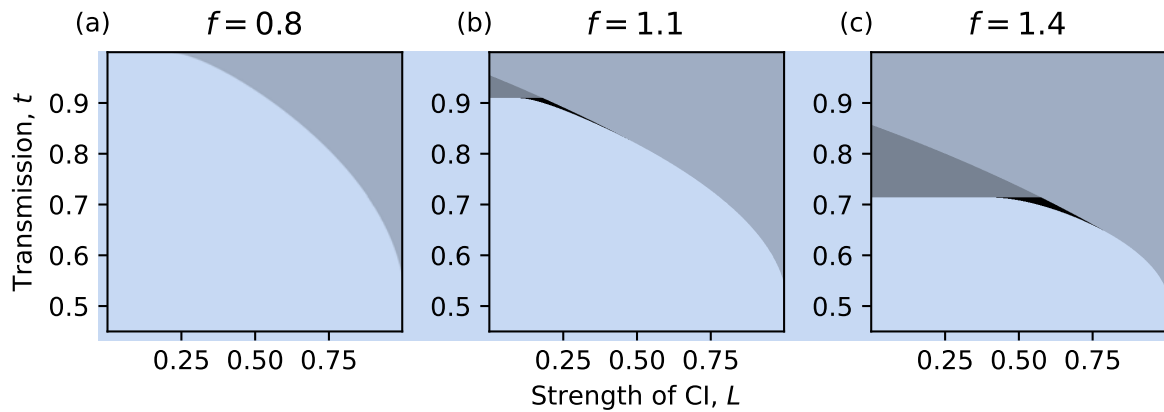


Figure 4: Properties of equilibria of the diploid model shown across parameter space. The parameter regions are categorized as featuring only the trivial equilibrium at $p = 0$ (white), a non-trivial stable equilibrium $\hat{p}_2 > 1/2$ with or without invasion threshold (light grey), and non-trivial stable equilibrium $\hat{p}_2 < 1/2$ without (dark grey) or with (black) an invasion threshold. Values of parameter f (fecundity) are indicated above each panel.

than what can occur if $f \leq 1$ (Fig. 4). The “classical” example with two stable equilibria and an invasion threshold between them (light grey in Fig. 4), as well as extinction of *Wolbachia* (white), can be found whether f indicates costs or benefits to being infected, but $f > 1$ additionally permits cases without invasion barriers and with low stable prevalence of *Wolbachia* when low or moderate L combines with $ft > 1$ (dark grey areas). Finally, the situation with the stable equilibrium below one half co-occurring with an invasion threshold places conflicting demands on the value of f (see above), and while not impossible to fulfil, satisfying them requires rather fine tuned parameter settings with low or intermediate values of L and for a narrow range of t slightly below $1/f$ (black areas in Fig. 4).

Haplodiploid models

Thus far, the presented analysis of low stable frequencies of *Wolbachia* assumed diploid sex determination. In this section, we extend these results to two haplodiploid cases, that differ in the effects of an infection on the host. The first case resembles the classic

diplo-diploid model in that cytoplasmic incompatibility kills fertilized eggs, but since under haplo-diploidy all fertilized eggs develop as females, we now switch to calling it the female-killing effect (and note that it is equivalent to the “*Leptopilina* type” of Vavre et al. (2000)). In the second case, incompatibility instead leads to the loss of paternal chromosomes and haploidization of the zygote, which consequently develops as a male (Breeuwer and Werren, 1990). We call this the masculinization effect (Vavre et al. (2000) call it the “*Nasonia* type”, see also Breeuwer and Werren (1990)). Note that both the female-killing effect and the masculinization effect can lead to a male-biased sex ratio, but the underlying mechanisms differ, and we model them separately.

The existence of a low-value stable equilibrium of *Wolbachia* with $f > 1$ also proves true in these two haplo-diploid systems. We consider the female-killing effect first. Applying our notation from above to the model of Vavre et al. (2000), the dynamics for *Wolbachia* frequency in females and males, p_F and p_M respectively, become

$$\begin{aligned} \Delta p_F &= \frac{p_F f t}{p_F f (1 - (1 - t)p_M L) + (1 - p_F)(1 - p_M L)} - p_F \\ \Delta p_M &= \frac{p_F f t}{p_F f + (1 - p_F)} - p_M \end{aligned} \quad (3)$$

For the masculinization effect, we again adopt the model from Vavre et al. (2000); the dynamics for females and males, respectively, are given as

$$\begin{aligned} \Delta p_F &= \frac{p_F f t}{p_F f (1 - (1 - t)p_M L) + (1 - p_F)(1 - p_M L)} - p_F \\ \Delta p_M &= \frac{p_F f t}{p_F f (1 + (1 - t)\frac{k}{1-k}p_M L) + (1 - p_F)(1 + \frac{k}{1-k}p_M L)} - p_M \end{aligned} \quad (4)$$

The female equations are identical across the two effects (Eqs. (3) and (4)), as females are lost due to infections impacting fertilized eggs in both cases. However, the male equations differ, because the masculinization effect converts females into males instead of killing them. The magnitude of this additional “male production” depends on the

288 fertilization rate k . If k is high, CI has the potential to cause strong effects on the
primary sex ratio, both because the baseline male production is by definition low under
290 high k , and because there are many fertilized eggs available to be masculinized. For the
female equation, the parameter k cancels out.

292 Appendix A gives explicit solutions for *Wolbachia* frequencies at the two non-trivial
equilibria in these models. In Appendix B we explore the stability of the non-trivial
294 equilibria and show numerically that when the two non-trivial equilibria exist, the lower
frequency equilibrium is unstable and the higher one is stable (as claimed by Vavre et al.,
296 2000). In Appendix A, we show analytically that the stable equilibrium frequency for
Wolbachia infection exceeds one half when $f \leq 1$, with the caveat that this applies to
298 both sexes in the female-killing model, but only to the *Wolbachia* frequency in female
hosts in the masculinization model. Regardless of this caveat, numerical examples show
300 that the ability of $f > 1$ to create stable equilibria below one half for both sexes extends
to both models (for examples of equilibria, see Fig. 5). The proofs in Appendix A
302 follow the same logic as for the diplodiploid model: the unstable and stable equilibria are
symmetric around a point r (for each sex), which is never below one half if $f \leq 1$. Hence,
304 again, a stable equilibrium close to one half may be vulnerable to the population falling
under the invasion barrier if stochasticity causes fluctuations in *Wolbachia* frequency, as
306 discussed above with the diplodiploid model.

The infection frequency in males generally does not have to be the same as in females,
308 because of the potential for additional female mortality and of the two different ways
in which males can be produced: the egg not being fertilized in the first place, or via
310 masculinization. In Appendix C, we prove analytically that infection frequency in
females is always higher than or equal to that in males, for both the female-killing
312 model and the masculinization model.

The additional flexibility actually brings about the possibility, in the masculinization

314 case, that the frequency of infection in males can stabilize below one half even with
316 $f = 0.95 < 1$ (Fig. 5c). The parameter range for this type of situation is narrow
318 whenever f is low (Fig. 6), though. In addition, the female infection frequency was
320 very high whenever $f \leq 1$, so that the average infection frequency was above one half
322 even in the few cases where male infection frequency was below one half under the
masculinization model (Fig. 7). Such cases were found with high fertilization rate (k)
yielding a strongly female-biased sex ratio, hence the average infection frequency was
close to that found in females. Overall, low infection frequencies seem very rare unless
 $f > 1$.

Discussion

324 The traditional model of the dynamics of symbionts inducing cytoplasmic incompatibility
326 predicts stable infection frequencies above one half in diploid systems (Turelli, 1994;
Turelli and Hoffmann, 1995). However, the prediction of high stable frequencies is tightly
328 linked to an assumption of direct fitness effects of infection being either neutral or
detrimental. Our findings, that low-frequency (below one half) stable equilibria can
330 exist, are as such not new: examples were presented by Zug and Hammerstein (2018),
but these authors did not elaborate on when exactly low frequencies can be expected to
prevail. We show analytically that they can occur only if the relative fecundity of the
332 infected individuals is higher than one.

The association between negative fitness effects and high frequencies is explained by
334 positive frequency-dependence. CI penalizes uninfected individuals' reproduction, but
cannot do so very efficiently if infections are rare; therefore, infections must overcome
336 an invasion barrier first but can thereafter spread to high frequencies (only counteracted
by leaky transmission). Positive fitness effects remove the difficulties of spreading from
338 very low frequencies, and as a whole this creates conditions for stable equilibria at low

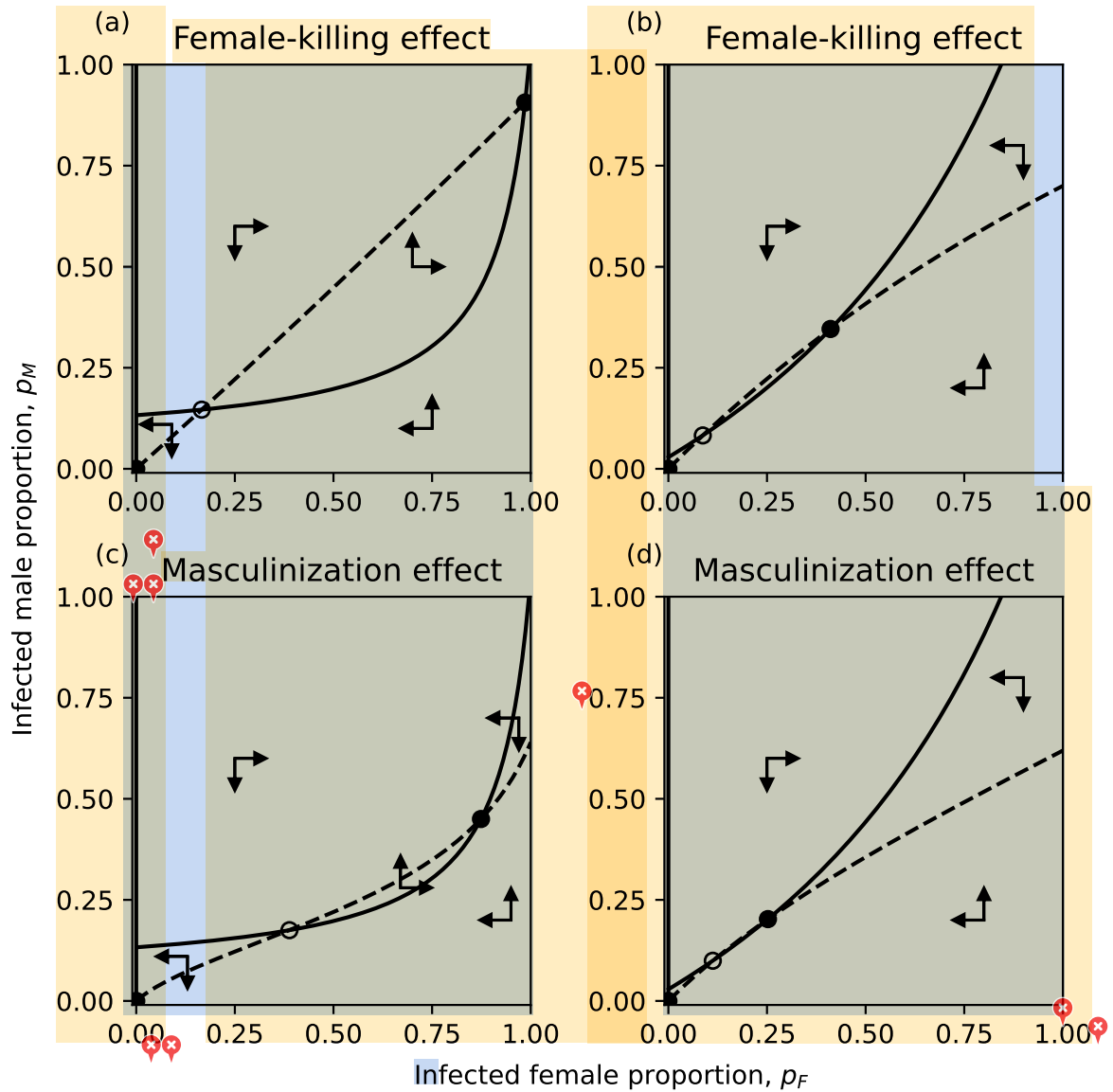


Figure 5: Equilibria of the two haplodiploid models in (p_F, p_M) -space. In left column $f < 1$, right $f > 1$. Dot: stable equilibria, circle: unstable equilibrium. Dashed line: male null-cline ($\Delta p_M = 0$), solid line: female null-cline ($\Delta p_F = 0$). a) The stable equilibrium is close to fixation and invasion threshold exists. b) Low frequency stable equilibrium with invasion threshold. c) The non-trivial stable equilibrium at $\hat{p}_M < 1/2$, although $f < 1$. d) Low frequency stable equilibrium with invasion threshold. Parameters: a) and c) $f = 0.95$, $t = 0.92$, $L = 0.95$, $k = 0.9$ (only c); b) and d) $f = 1.4$, $t = 0.7$, $L = 0.7$, $k = 0.5$ (only d).

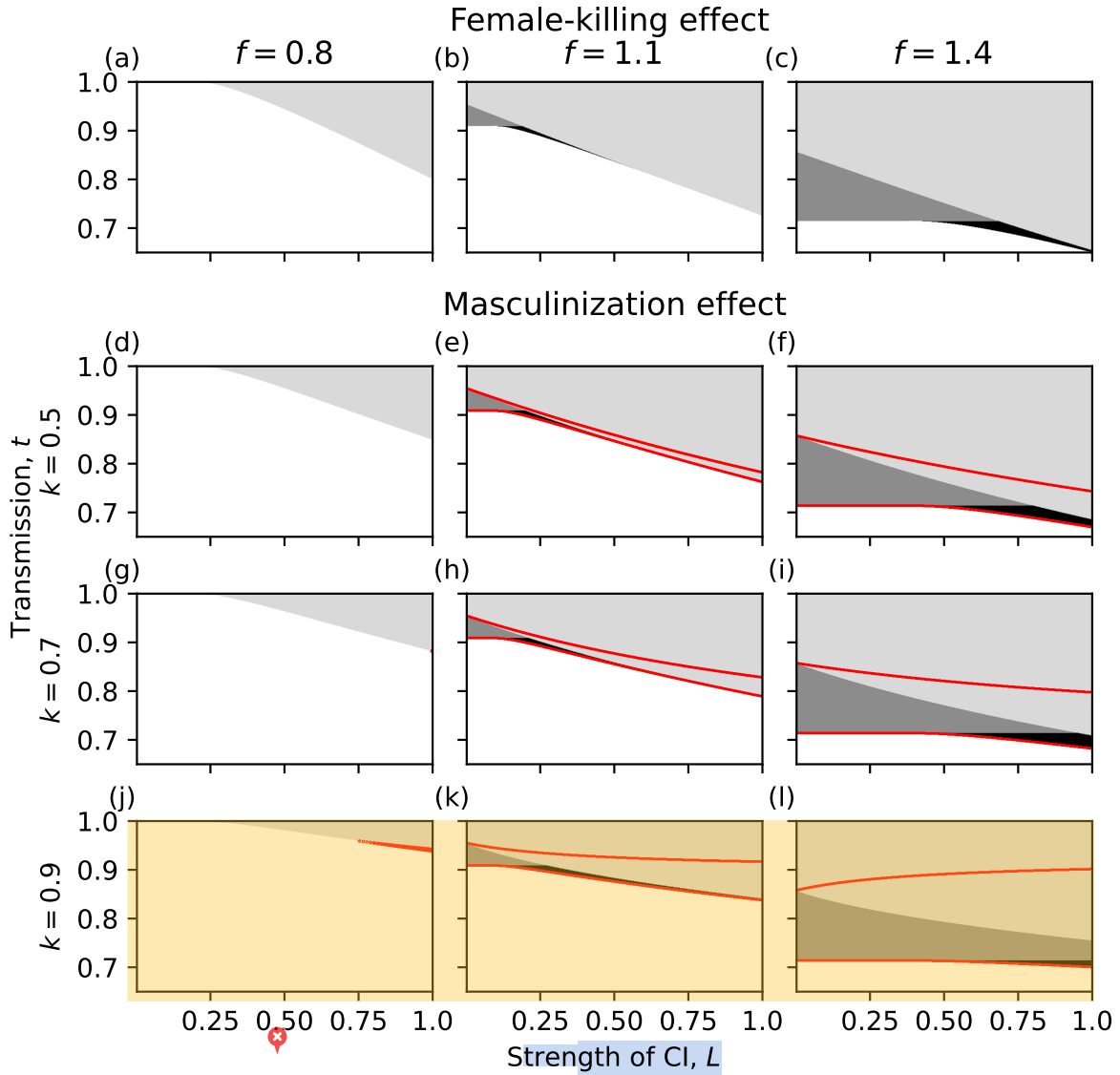


Figure 6: Classification of equilibria of the haplodiploid models across parameter space. Columns from left to right with $f = 0.8$, $f = 1.1$ and $f = 1.4$. First row: female-killing effect; other rows: masculinization effect model with different values of k . White: only the trivial equilibrium at zero. Light grey: non-trivial stable equilibrium with $1/2 < \hat{p}_F \leq 1$. Dark grey: $\hat{p}_F \leq 1/2$ without invasion threshold. Black: $\hat{p}_F \leq 1/2$ with invasion threshold. The red lines outline two thresholds, between which $\hat{p}_M \leq 1/2$. Note that the vertical t -axes span only high values, where non-trivial equilibria exist.

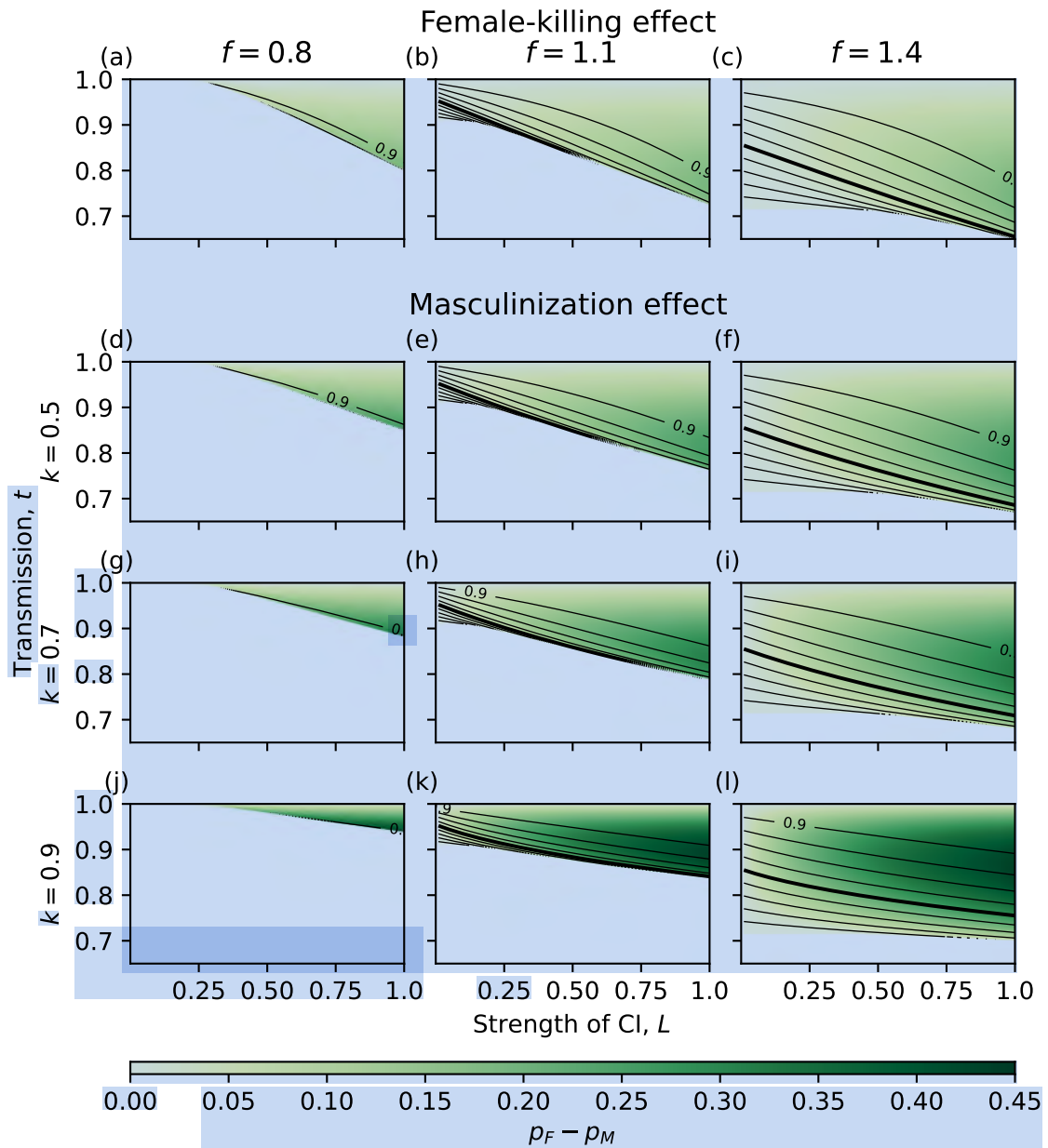


Figure 7: Equilibrium infection frequencies of the haplodiploid models across parameter space. Columns from left to right with $f = 0.8$, $f = 1.1$ and $f = 1.4$. First row: female-killing effect; other rows: masculinization effect model with different values of k . Black contour lines show infection frequency in females (with gaps of 0.1), while the color shows difference between female and male infection frequency according to the colorbar in the bottom. Thick solid contour marks one half frequency. Females always have higher infection frequency than males.

frequency. Similar results hold for haplodiploid systems, with the exception of male
340 infection frequency in masculinization-type CI, which can stay below one half even with
relative fecundity less than or equal to one.

342 The simplest case of low stable frequencies occurs when the effective fitness of rare
infected individuals is higher than that of uninfected individuals ($ft > 1$). In that case,
344 a rare infection can spread as there is no invasion barrier (\hat{p}_0 is unstable; see also Zug and
Hammerstein, 2018), and the infection frequency rises to levels dictated by the strength
346 of CI, efficiency of transmission, and relative fecundity. With very weak CI and leaky
transmission, the stable equilibrium can be very close to zero (e.g. 0.06 in Fig. 2c,
348 d). A more complex situation occurs when the relative fecundity of infected individuals
exceeds one but not by much, so that relative fecundity above one combines with effective
350 fecundity below one ($f > 1$ with $ft < 1$). In that case, the invasion threshold exists,
but it is still possible to have a stable equilibrium below one half, a finding that has
352 not been demonstrated before. As the previous analyses usually assumed equal or lower
fecundity of infected individuals, they consequently observed stable equilibria above one
354 half.

Note that similar results arise when considering a sexual population containing a lin-
356 eage with a wild-type symbiotic strain and a lineage with a fitness-increasing symbiotic
strain with non-perfect transmission. As shown by Zug and Hammerstein (2018), the
358 condition $ft > 1$ is necessary and sufficient for the successful invasion of the fitness-
increasing strain, as it provides the strain with higher effective fecundity than the wild-
360 type lineage. They also mentioned this being analogous to the mutation-selection balance
of haploids (Hoffmann and Turelli, 1997). Along the same lines, when studying the pos-
362 sibility of sex-ratio determination of CI-inducing symbionts, Egas et al. (2002) conclude
that the symbiont can invade “if the proportion [of] infected daughters produced by
364 infected females is bigger than the proportion [of] daughters produced by uninfected fe-

males (when mated with uninfected males)”, again returning to higher effective fecundity
366 of the infected individuals.

Given that the consequences of higher relative fecundity are not only clear but also,
368 in the form of examples, present in earlier analyses, why have positive fitness effects
not been discussed explicitly before? A typical discussion of *Wolbachia* focuses on the
370 fitness deficit caused by infection (Hoffmann et al., 1990), which creates an interesting
paradox: why should a symbiont with negative fitness effects on its host be so common
372 in natural populations? The discovery of reproductive manipulation (such as CI) solved
this conundrum. The possibility of positive fitness effects has thereafter not attracted
374 much attention, despite the relevant equations remaining valid when $f > 1$. As an
exception, Zug and Hammerstein (2018) recently explored positive fitness effects, with
376 a focus on the evolution of the infectious agent and comparison of different symbionts
(non-manipulating, CI, male-killing).

Besides showing that the stable equilibrium almost always exceeds one half when
378 $f \leq 1$, we also argue that infection frequencies slightly above one half are not expected to
be robust in nature. Our model analysis thus suggests that positive fitness effects of CI-
380 inducing *Wolbachia* infection are expected to be found in those natural populations that
show low and stable densities of infection over time. Stable infection frequencies below
382 one half and even slightly above it are likely paired with positive fitness effect unless
another dynamic can explain the situation. Besides the fitness benefit demonstrated
384 here, low frequencies may of course be observable during ongoing invasion or result from
source-sink dynamics (e.g. Flor et al., 2007; Telschow et al., 2007).
386

It would be ideal to complement our findings with concrete empirical examples of
388 stable CI-inducing *Wolbachia* strains that are maintained at low frequencies in their
host populations. Unfortunately, it is difficult to highlight one specific case that perfectly
390 illustrates our findings. Although it is well appreciated that *Wolbachia* is common among

species while not necessarily reaching high frequencies within a species (Sazama et al.,
392 2019), studies typically do not provide much information regarding additional fitness
effects when populations have been screened for *Wolbachia* infection (Russell et al.,
394 2012). However, there is some empirical evidence for *Wolbachia* strains that elevate
host fitness, where the benefit may manifest itself, for example, as higher fecundity (Fry
396 et al., 2004) or protection from viral diseases (Chrostek et al., 2013), and also evidence
of evolution towards mutualism (higher fecundity) in natural *Wolbachia* populations
398 (Weeks et al., 2007). Often, there is no information on whether the symbiont induces CI
or any other phenotype in their hosts (Duplouy et al., 2015). The stability properties of
400 reported low frequencies are rarely evaluated, as most studies report prevalence of the
symbiont at a single time point; recurrent screenings of the same natural populations over
402 the course of several generations are rare (however, see Kriesner et al., 2013; Duplouy
et al., 2015, 2021). Hence, the changes in the infection frequencies through generations
404 remain unknown, and it is difficult to determine stability of the infection frequency, let
alone any existence of invasion barriers or other dynamical features of the system.

406 Another testable prediction from the presented models is that in the haplodiploid
models the female infection frequency seems to be consistently higher than the male
408 infection frequency. Additionally, the masculinization type seems to predict lower equi-
librium frequencies than the female-killing type. The latter observation might be ex-
410 plained, at least partially, by the fact that masculinization allows for excess production
of uninfected males (compared to the zygotes dying), which “dilutes” the infection.

412 Further possibility for extension of this work would be to study the total size of the
host population and the effect of *Wolbachia* on that. Also, understanding the details of
414 dynamics approaching to fixed points would shed light on the diversity of possibilities
that could be observed in nature. Both the host population size and the dynamics before
416 reaching a stable equilibrium would be of interest when applying *Wolbachia* as a control

agent.

418 Our model highlights that when strains are found at low frequencies in their host
populations, there are additional possibilities to them being either in the process of
420 being eliminated or of invading (Duplouy et al., 2010; Kriesner et al., 2013) their host
population: a low frequency can be a stable outcome. As the mitochondrial haplotype
422 will hitchhike a successfully spreading maternally inherited *Wolbachia* infection, the state
of potentially ongoing change can partially be answered with the study of the associated
424 mitochondrial haplotype diversity in the host population, but often even this data is not
available (but see Duplouy et al., 2010; Richardson et al., 2012). Long-term surveys of
426 host populations and their infection status through time, as well as the study of their
ecology are therefore needed. The recent clear results from the effective control strategy
428 against human-born disease such as Dengue (suppressing its vector, *Aedes aegypti*) have
clearly shown that long-term surveys are possible and informative for endosymbionts
430 (e.g. Ryan et al., 2019).

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Conflict of interest disclosure

440 The authors declare they have no conflict of interest relating to the content of this article.
AD is a recommender for PCI Ecology.

442 Data, script and code availability

Matlab (v. R2019b) and Mathematica (v. 12.1.1.0) were used for solving and analysing
444 equations. The images were created and numerical analysis conducted in Python (v.
3.7.4) using packages numpy (v. 1.21.5 Harris et al., 2020) and matplotlib (v. 3.4.2
446 Hunter, 2007). Source code is provided in the Zenodo repository (Karisto et al., 2022).

Appendix A: Limits for CI frequency in haplodiploid systems

Female-killing effect

When CI causes mortality of the female embryo in haplodiploid species, the population dynamics follow the model in Eq. (3). The non-trivial equilibria of the model are

$$\hat{p}_F = \frac{2 - 2f - ft + f^2t + fLt \pm ft\sqrt{(1 + L - f)^2 + 4L(ft - 1)}}{2[1 + f(f - 2 + Lt + fLt(t - 1))]}, \quad (A1)$$

$$\hat{p}_M = \frac{1 + L - f \pm \sqrt{(1 + L - f)^2 + 4L(ft - 1)}}{2L},$$

where pluses belong to the non-trivial stable equilibrium \hat{p}_2 and minuses to the unstable equilibrium \hat{p}_1 when they exist (Vavre et al., 2000). The stability of these is explored numerically in Appendix B.

Note that for the equilibrium \hat{p}_1 we can see that when $ft = 1$, $\hat{p}_M = \hat{p}_F = 0$ and additionally when $ft > 1$, $\hat{p}_M < 0$, hence the “invasion threshold” coincides with \hat{p}_0 at $ft = 1$ and is not in the biologically relevant range when $ft > 1$, similar to the diploid model.

We use the same technique as for diploid model, showing that the “real part” r (the part that is necessarily real, i.e. excluding the square root part) of the equilibrium frequency is below one half if and only if $f > 1$, and hence $\hat{p}_2 > 1/2$ for $f \leq 1$. For the male frequency, it is easy to see that

$$r_M = \frac{1 - f + L}{2L} < \frac{1}{2} \Leftrightarrow 1 - f < 0 \Leftrightarrow f > 1,$$

464 but for the female frequency the proof is somewhat more complicated. In that case

$$r_F = \frac{2 - 2f - ft + f^2t + fLt}{2 \left[1 + f(f - 2 + Lt + fLt(t - 1)) \right]}$$

The denominator of r_F divided by two (i.e. the expression within square brackets) is a
466 quadratic function of f ,

$$(1 + Lt(t - 1))f^2 + (Lt - 2)f + 1,$$

which is always positive, as the intercept is positive and no real roots exist (discriminant
468 $Lt^2(L - 4) < 0$). As the denominator of r_F is always larger than zero, we can multiply
the inequality $r_F < 1/2$ with the denominator, resulting in an equivalent inequality

$$(1 - t)(1 - Lt)f^2 + tf - 1 > 0. \quad (A2)$$

In Eq. (A2) the intercept of the left hand side is negative and the quadratic coefficient
472 $(1 - t)(1 - Lt) > 0$. Hence, the equation holds true for positive f only when f is larger
than the larger root, say f_{lim} , of the quadratic function in the left hand side of Eq. (A2).
474 Only if $f_{\text{lim}} < 1$, can $f \leq 1$ fulfil the inequality in Eq. (A2) and equivalently $r_F < 1/2$.
Let us now investigate in what conditions $f_{\text{lim}} < 1$:

$$\begin{aligned} f_{\text{lim}} &= \frac{-t + \sqrt{t^2 + 4(1-t)(1-Lt)}}{2(1-t)(1-Lt)} < 1 \\ &\Leftrightarrow \sqrt{t^2 + 4(1-t)(1-Lt)} < 2(1-t)(1-Lt) + t \quad | (\cdot)^2, \text{ both sides positive} \\ &\Leftrightarrow t^2 + 4(1-t)(1-Lt) < 4(1-t)^2(1-Lt)^2 + 4t(1-t)(1-Lt) + t^2 \\ &\Leftrightarrow 4(1-t)^2(1-Lt) < 4(1-t)^2(1-Lt)^2 \quad | : 4(1-t)^2(1-Lt) > 0 \\ &\Leftrightarrow 1 < 1 - Lt \\ &\Leftrightarrow Lt < 0. \end{aligned}$$

476 Thus, the lower limit f_{lim} is below one only if $Lt < 0$, which is never true. Hence the
 limit is always at least 1, which implies that $r_F < 1/2$ is true only if $f > 1$. This
 478 concludes the proof that $\hat{p}_{F,2} > 1/2$ for the female-killing type CI when $f \leq 1$.

* Masculinization effect

480 The CI can alternatively lead to deletion of the paternal genome, turning the diploid
 egg into a haploid that develops as male. In that case, the population dynamics follow
 482 the model in Eq. (4). The non-trivial equilibria of the model are

$$\hat{p}_F = \frac{2 + f^2(1+k)t - f \left[2 + t(1+k+L(k-1)) \right]}{2 \left[1 - f \left[2 + Lt(k-1) - f(1 - Lt(k-1)(t-1)) \right] \right]} \pm \frac{ft \sqrt{(k-1) \left[(k-1)(1-f+L)^2 + 4L(k-1+fk(t-1))(ft-1) \right]}}{2 \left[1 - f \left[2 + Lt(k-1) - f(1 - Lt(k-1)(t-1)) \right] \right]} \quad (\text{A3})$$

$$\hat{p}_M = \frac{1 - f - k + fk + L - kL}{2L(1 - k + fk(1 - t))} \pm \frac{\sqrt{(k-1) \left[(k-1)(1-f+L)^2 + 4L(k-1+fk(t-1))(ft-1) \right]}}{2L(1 - k + fk(1 - t))}$$

484 where pluses belong to the non-trivial stable equilibrium \hat{p}_2 and minuses to the unstable
 equilibrium \hat{p}_1 when they exist (Vavre et al., 2000). The stability of these is explored
 486 numerically in Appendix B.

We show that $\hat{p}_{F,2} > 1/2$ whenever $f \leq 1$ using the same logic as before, i.e. we show
 488 by contradiction that for any $f \leq 1$ and valid parameter values, “the real part” of the
 solution, r_F , cannot be less than 1/2. Thus, $r_F \geq 1/2$ and hence also $\hat{p}_{F,2} > 1/2$ as it is

490 larger than r_F , which is

$$r_F = \frac{2 + f^2(1+k)t - f[2 + t(1+k + L(k-1))]}{2\left[1 - f\left[2 + Lt(k-1) - f(1 - Lt(k-1)(t-1))\right]\right]}.$$

The denominator of r_F is a linear function of k :

$$2(f-1)^2 + 2fLt(1+f(t-1)) + k[-2fLt(1+f(t-1))].$$

492 When $f \leq 1$, the slope of that function is negative ($-2fLt(1+f(t-1)) < 0$) and the root satisfies

$$1 + \frac{(f-1)^2}{fLt(1+f(t-1))} > 1.$$

494 Thus, with $f \leq 1$ the denominator is positive for $k < 1$, and the inequality $r_F < 1/2$ simplifies to the equivalent condition

$$496 \quad (1-f)(1+f-ft) + f^2t(1-t)L + k(-ft + f^2t + f^2Lt(t-1)) < 0. \quad (\text{A4})$$

The left hand side is again a linear function of k , which has an intercept that is always positive for $L \in (0, 1]$ as it is linear in L and has positive intercept and slope. The linear function of k in Eq. (A4) has one root at

$$\hat{k} = 1 + \frac{1-f^2 - 2f(1-f)t}{ft(1-f+fL(1-t))} > 1,$$

500 as the second term is positive for $f \leq 1$. The denominator of the second term is obviously positive and the numerator is also positive, as it is linear in t , having negative slope and root $(1+f)/2f \geq 1$ when $f \leq 1$.

Thus, Eq. (A4) has positive intercept and $\hat{k} \geq 1$ holds, which means that the left hand side of Eq. (A4) is positive for all $k \in (0, 1]$ and hence the inequality in Eq. (A4)

is never true for valid values of k . As Eq. (A4) is equivalent with $r_F < 1/2$, this implies
506 that the real part of the female equilibrium frequency is never less than $1/2$ with valid
parameter values (assuming $f \leq 1$). The actual stable equilibrium is even higher than
508 r_F , which concludes the proof that female frequency is never below $1/2$ at the stable
equilibrium.

510 Finally, for the infection frequency in males under the masculinization effect model, it
is easy to prove by example that the frequency at the stable equilibrium indeed can be
512 below $1/2$ even with $f < 1$. In Fig. 5c, the parameters are $f = 0.95$, $t = 0.92$, $L = 0.95$,
 $k = 0.9$ while $\hat{p}_{M,2} < 1/2$.

514 In conclusion, the infection frequency in females follows the same pattern as female-
killing type and diploid systems with symmetry of the stable equilibrium and the
516 unstable equilibrium around a point that is above one half for all $f \leq 1$, but the infection
frequency in males makes an exception as it can go below one half even with $f < 1$.

518 Appendix B: Local stability analysis of the equilibria of CI models

520 In this appendix we will use linear stability analysis to calculate the local stability of
the equilibria of the different models presented in the main text.

522 Diplodiploid model

The **dynamics** of *Wolbachia* **frequency** in both males and females in this model is given
524 by Eq. (1) in the main text, here repeated in slightly different form that specifies the
time T explicitly.

$$526 \quad p_{T+1} = \frac{p_T f t}{p_T f (1 - p_T L (1 - t)) + (1 - p_T)(1 - p_T L)}. \quad (\text{B1})$$

For notational convenience, define a function g as

$$g(p_T) = \frac{f t}{p_T f (1 - p_T L (1 - t)) + (1 - p_T)(1 - p_T L)},$$

528 so that

$$p_{T+1} = g(p_T) p_T. \quad (\text{B2})$$

530 Equilibria of this discrete time dynamical system satisfy

$$531 \quad \hat{p} = g(\hat{p}) \hat{p}. \quad (\text{B3})$$

532 An equilibrium, \hat{p} , is locally stable if and only if the absolute value of the derivative of
 p_{T+1} with respect to p_T is smaller than one (see e.g. Otto and Day, 2007), that is,

$$\left| \frac{dp_{T+1}}{dp_T} \Big|_{\hat{p}} \right| < 1.$$

534 We will denote this derivative evaluated at equilibrium \hat{p} as $\zeta(\hat{p})$.

Therefore to calculate the stability of the equilibria of our diploid model, we need
 536 the derivative of p_{T+1} with respect to p_T . Differentiating Eq. (B2) gives

$$\frac{dp_{T+1}}{dp_T} = g(p_T) + p_T \frac{dg}{dp_T}.$$

Next we want to evaluate this derivative at the equilibria. Equation (B1) has three
 538 equilibria, solved from equation $Ap^2 + Bp + C = 0$, as shown in the main text and
 repeated here for convenience:

$$540 \quad \hat{p}_0 = 0, \hat{p}_1 = \frac{-B - \sqrt{B^2 - 4AC}}{2A}, \text{ and } \hat{p}_2 = \frac{-B + \sqrt{B^2 - 4AC}}{2A}, \quad (\text{B4})$$

where $A = L(1 - f(1 - t))$, $B = -1 + f - L$ and $C = 1 - ft$.

542 The stability of the first equilibrium, \hat{p}_0 , is given by

$$\begin{aligned} \zeta(\hat{p}_0) &= \left. \frac{dp_{T+1}}{dp_T} \right|_{\hat{p}_0} \\ 544 \quad &= g(0) \\ &= ft. \end{aligned}$$

546 ✖ The equilibrium $\hat{p}_0 = 0$ is therefore stable if and only if $ft < 1$. At $ft = 1$, the equilibria
 \hat{p}_0 and \hat{p}_1 coincide, and when $ft > 1$ \hat{p}_0 is unstable. Note that this has implications on
 548 the existence of invasion threshold, since there is no invasion threshold if \hat{p}_0 is unstable
 ($\hat{p}_1 < 0$ when $ft > 1$). In that case the infection frequency will increase from any initial
 550 frequency until it reaches a stable equilibrium.

✖ The stability of the second equilibrium, \hat{p}_1 , is given by

$$552 \quad \zeta(\hat{p}_1) = \left. \left[g(\hat{p}_1) + \hat{p}_1 \frac{dg}{dp_T} \right] \right|_{\hat{p}_1}.$$

* The first term $g(\hat{p}_1) = 1$ when $\hat{p}_1 \neq 0$, because of Eq. (B3). The second term is

$$\begin{aligned} \hat{p}_1 \frac{dg}{dp_T} \Big|_{\hat{p}_1} &= \frac{-\hat{p}_1}{ft} \left[f - 1 - L + 2\hat{p}_1 L(1 - f(1 - t)) \right] \\ &= -\frac{\hat{p}_1}{ft} (B + 2\hat{p}_1 A), \end{aligned}$$

and hence

$$\zeta(\hat{p}_1) = \left| 1 - \frac{\hat{p}_1}{ft} (B + 2\hat{p}_1 A) \right|,$$

or using Eq. (B4),

$$\zeta(\hat{p}_1) = \left| 1 + \frac{\hat{p}_1}{ft} \sqrt{B^2 - 4AC} \right|.$$

Stability requires that $\zeta(\hat{p}_1) < 1$. Since $ft > 0$ always holds, equilibrium \hat{p}_1 is unstable

* whenever $\hat{p}_1 > 0$. The first non-trivial equilibrium is therefore always unstable when it is positive, in agreement with Hoffmann et al. (1990).

* Similarly, the stability of the second equilibrium, \hat{p}_2 , is given by

$$\begin{aligned} \zeta(\hat{p}_2) &= \left| g(\hat{p}_2) + \hat{p}_2 \frac{dg}{dp_T} \Big|_{\hat{p}_2} \right| \\ &= \left| 1 - \frac{\hat{p}_2}{ft} \sqrt{B^2 - 4AC} \right|. \end{aligned}$$

The equilibrium \hat{p}_2 is therefore stable if and only if

$$0 < \frac{\hat{p}_2}{ft} \sqrt{B^2 - 4AC} < 2. \quad (\text{B5})$$

If $\hat{p}_2 > 0$, then the left-hand side of this inequality is true. To check the right-hand side

of the inequality, we substitute the expression of \hat{p}_2 in Eq. (B4) into Eq. (B5) to obtain:

$$\frac{B^2 - 4AC - B\sqrt{B^2 - 4AC}}{2A(1 - C)} < 2,$$

570 or equivalently

$$B \left(\frac{B - \sqrt{B^2 - 4AC}}{4A} \right) < 1,$$

which can be written as

$$572 \quad \frac{-B}{2} \hat{p}_2 < 1. \quad (\text{B6})$$

Now, if $B < 0$ (i.e. $f < 1 + L$) we can multiply both sides of inequality by $-2/B$ to get
574 an equivalent stability condition

$$\hat{p}_2 < \frac{2}{1 + L - f},$$

which is always true since $0 < 1 + L - f < 2$ and hence the right-hand side is larger than
576 1. Therefore as long as $\hat{p}_2 \leq 1$ this condition is fulfilled and \hat{p}_2 is a stable equilibrium.
On the other hand, if $B \geq 0$ (i.e. $f \geq 1 + L$), the left hand side of Eq. (B6) is zero
578 or negative since $-B < 0$ and the inequality is true for all $\hat{p}_2 \geq 0$. Hence, \hat{p}_2 is always
stable if it exists within the biologically relevant range between 0 and 1.

580 Haplodiploid models

The *Wolbachia* models with haplodiploid sex determination in the main text are both
582 frequency-dependent models that in general we can write as follows:

$$\mathbf{p}_{T+1} = \mathbf{B}(\mathbf{p}_T)\mathbf{p}_T, \quad (\text{B7})$$

584 where \mathbf{p}_T is a vector with the frequency of *Wolbachia* infection in females and males
as its entries. Moreover, both the female-killing effect model, and the masculinization
586 effect model have the following general shape:

$$\begin{pmatrix} p_F \\ p_M \end{pmatrix}_{T+1} = \begin{pmatrix} g(p_{F,T}, p_{M,T}) & 0 \\ h(p_{F,T}, p_{M,T}) & 0 \end{pmatrix} \begin{pmatrix} p_F \\ p_M \end{pmatrix}_T, \quad (\text{B8})$$

where the functions g and h are different for the female-killing model and the masculinization
588 model, and they are presented in the sections below.

Equilibria, $\hat{\mathbf{p}}$, of this discrete time dynamical system satisfy

$$\hat{\mathbf{p}} = \mathbf{B}(\hat{\mathbf{p}})\hat{\mathbf{p}}.$$

590 The stability of an equilibrium to small perturbations is determined by the absolute value
of the dominant eigenvalue of the Jacobian matrix of the model evaluated at the equilib-
592 rium. The equilibrium is stable if the absolute value of the dominant eigenvalue of the
Jacobian matrix is smaller than one. The Jacobian matrix is obtained by differentiating
594 Eq. (B7) and when evaluated at an equilibrium it is

$$\mathbf{M} = \left. \frac{d\mathbf{p}_{T+1}}{d\mathbf{p}_T^T} \right|_{\hat{\mathbf{p}}} = \begin{pmatrix} \frac{\partial p_{F,T+1}}{\partial p_{F,T}} & \frac{\partial p_{F,T+1}}{\partial p_{M,T}} \\ \frac{\partial p_{M,T+1}}{\partial p_{F,T}} & \frac{\partial p_{M,T+1}}{\partial p_{M,T}} \end{pmatrix} \Big|_{\hat{\mathbf{p}}}.$$

As both models have similar structure (Eq. B8) their Jacobian matrices have the general
596 form

$$\mathbf{M} = \begin{pmatrix} g(\hat{\mathbf{p}}) & 0 \\ h(\hat{\mathbf{p}}) & 0 \end{pmatrix} + \hat{p}_F \begin{pmatrix} \frac{\partial g}{\partial p_F} & \frac{\partial g}{\partial p_M} \\ \frac{\partial h}{\partial p_F} & \frac{\partial h}{\partial p_M} \end{pmatrix} \Big|_{\hat{\mathbf{p}}}. \quad (\text{B9})$$

Female-killing effect

We consider the female-killing effect first (Eq. (3) from the main text in recursion form with explicit time T):

$$\begin{aligned} p_{F,T+1} &= \frac{p_{F,T} ft}{p_{F,T} f (1 - p_{M,T} L (1 - t)) + (1 - p_{F,T}) (1 - p_{M,T} L)}, \\ p_{M,T+1} &= \frac{p_{F,T} ft}{p_{F,T} f + (1 - p_{F,T})}. \end{aligned} \tag{B10}$$

Therefore, if we define,

$$\begin{aligned} g(p_{F,T}, p_{M,T}) &= \frac{ft}{p_{F,T} f (1 - p_{M,T} L (1 - t)) + (1 - p_{F,T}) (1 - p_{M,T} L)}, \\ h(p_{F,T}, p_{M,T}) &= \frac{ft}{p_{F,T} f + (1 - p_{F,T})}, \end{aligned}$$

then we can write the recursion equations (B10) in the matrix form given above by equation (B8).

For the trivial equilibrium, $\mathbf{p}_0 = (0, 0)$, $g(\hat{\mathbf{p}}) = h(\hat{\mathbf{p}}) = ft$ and we have

$$\mathbf{M} = \begin{pmatrix} ft & 0 \\ ft & 0 \end{pmatrix}.$$

The eigenvalues of this Jacobian matrix are 0 with eigenvector $(p_F, p_M) = (0, 1)$, and ft with eigenvector $(p_F, p_M) = (1, 1)$. The zero eigenvalue is associated with perturbations in the directions of males only, which makes sense biologically: these perturbations decay back to the zero equilibrium since males do not transmit *Wolbachia*. The largest eigenvalue of the $\mathbf{p}_0 = (0, 0)$ equilibrium is ft . Similarly to the diplo-diploid model, the trivial equilibrium is therefore stable if $ft < 1$, and unstable if $ft > 1$.

The non-trivial equilibria of this model are given in Eq. (A1) and repeated below,

$$\hat{p}_F = \frac{2 - 2f - ft + f^2t + fLt \pm ft\sqrt{(1+L-f)^2 + 4L(ft-1)}}{2\left[1 + f(f-2 + Lt + fLt(t-1))\right]},$$

$$\hat{p}_M = \frac{1 + L - f \pm \sqrt{(1+L-f)^2 + 4L(ft-1)}}{2L}.$$

612 To calculate the Jacobian matrix for the non-trivial equilibria, we need to calculate
the derivatives in Eq. (B9). It will be useful to note that for the non-trivial equilibria,
614 $\hat{\mathbf{p}} = \mathbf{B}(\hat{\mathbf{p}})\hat{\mathbf{p}}$ implies that $g(\hat{p}_F, \hat{p}_M) = 1$, and $h(\hat{p}_F, \hat{p}_M) = \frac{\hat{p}_M}{\hat{p}_F}$. It follows that

$$\begin{aligned} \left. \frac{\partial h}{\partial p_F} \right|_{\hat{\mathbf{p}}} &= \frac{1 - f \frac{\hat{p}_M^2}{\hat{p}_F^2}}{ft \frac{\hat{p}_F^2}{\hat{p}_F^2}}, \\ \left. \frac{\partial h}{\partial p_M} \right|_{\hat{\mathbf{p}}} &= 0, \end{aligned}$$

616

and

$$618 \quad \left. \frac{\partial g}{\partial p_F} \right|_{\hat{\mathbf{p}}} = \frac{1}{ft} g(p_F, p_M)^2 [1 - p_M L - f(1 - (1-t)p_M L)],$$

and therefore

$$620 \quad \left. \frac{\partial g}{\partial p_F} \right|_{\hat{\mathbf{p}}} = \frac{1}{ft} [1 - \hat{p}_M L - f(1 - (1-t)\hat{p}_M L)]. \quad (\text{B11})$$

The equilibrium condition $g(\hat{p}_F, \hat{p}_M) = 1$ can be rewritten as

$$\hat{p}_F f(1 - (1-t)\hat{p}_M L) = ft - 1 + \hat{p}_M L + \hat{p}_F - \hat{p}_F \hat{p}_M L.$$

622 Multiplying Eq. (B11) with \hat{p}_F as in Eq. (B9), and substituting the above expression

yields

$$\frac{\partial g}{\partial p_F} \Big|_{\hat{p}} = \frac{1}{ft} (1 - \hat{p}_M L - ft).$$

Similarly,

$$\frac{\partial g}{\partial p_M} = \frac{1}{ft} g(p_F, p_M)^2 (p_F f L (1 - t) + L (1 - p_F)),$$

and

$$\begin{aligned} \frac{\partial g}{\partial p_M} \Big|_{\hat{p}} &= \frac{1}{ft} (\hat{p}_F f L (1 - t) + L (1 - \hat{p}_F)) \\ &= \frac{1}{\hat{p}_M ft} (\hat{p}_F f + (1 - \hat{p}_F) - ft) \\ &= \frac{1}{\hat{p}_M} \left(\frac{\hat{p}_F}{\hat{p}_M} - 1 \right), \end{aligned}$$

where we have used $g(\hat{p}_F, \hat{p}_M) = 1$ and $h(\hat{p}_F, \hat{p}_M) = \frac{\hat{p}_M}{\hat{p}_F}$ for the last two lines, respectively.

Putting all of this together, we get the following Jacobian matrix,

$$\begin{aligned} \mathbf{M} &= \begin{pmatrix} 1 & 0 \\ \frac{\hat{p}_M}{\hat{p}_F} & 0 \end{pmatrix} + \begin{pmatrix} \frac{1}{ft} (1 - \hat{p}_M L - ft) & \frac{\hat{p}_F}{\hat{p}_M} \left(\frac{\hat{p}_F}{\hat{p}_M} - 1 \right) \\ \frac{(1-f) \hat{p}_M^2}{ft \hat{p}_F} & 0 \end{pmatrix} \\ &= \begin{pmatrix} \frac{1}{ft} (1 - \hat{p}_M L) & \frac{\hat{p}_F}{\hat{p}_M} \left(\frac{\hat{p}_F}{\hat{p}_M} - 1 \right) \\ \frac{\hat{p}_M}{\hat{p}_F} \left(1 + \frac{(1-f) \hat{p}_M}{ft} \right) & 0 \end{pmatrix}. \end{aligned}$$

The eigenvalues of this Jacobian, denoted by λ , are given by the following quadratic equation

$$\lambda^2 - \frac{\lambda}{ft} (1 - \hat{p}_M L) - \left(1 + \frac{1-f}{ft} \hat{p}_M \right) \left(\frac{\hat{p}_F}{\hat{p}_M} - 1 \right) = 0.$$

If the absolute value of the dominant eigenvalue of the Jacobian is below one, then the

638 equilibrium is stable. The eigenvalues are given by

$$\lambda = \frac{1}{2ft} (1 - \hat{p}_M L) \pm \frac{1}{2ft} \sqrt{(1 - \hat{p}_M L)^2 + 4(ft)^2 C},$$

where

$$C = \left(1 + \frac{1-f}{ft} \hat{p}_M \right) \left(\frac{\hat{p}_1}{\hat{p}_M} - 1 \right).$$

640 Since $L \leq 1$, the first term of λ is positive and the eigenvalue with the largest absolute value is

$$\lambda_1 = \frac{1}{2ft} (1 - \hat{p}_M L) + \frac{1}{2ft} \sqrt{(1 - \hat{p}_M L)^2 + 4(ft)^2 C}.$$

642 We could not determine stability of the equilibria analytically. Instead, we used
 Python to explore the parameter space $\{0.01 \leq L, t \leq 1\} \times \{0.01 \leq f \leq 3\}$ sam-
 644 pling values for each parameter uniformly with 0.01 interval, which resulted in 3 000 000
 parameter combinations. Whenever an equilibrium (either \hat{p}_1 or \hat{p}_2) appeared in the
 646 biologically meaningful range $0 < \hat{p}_M, \hat{p}_F \leq 1$, we analysed its stability. For all cases
 where $|\lambda_1| \neq 1$ (i.e. linear stability analysis did not fail) the linear stability analysis
 648 concluded that \hat{p}_1 was unstable and \hat{p}_2 was stable.

Masculinization effect

650 For the masculinization effect, the dynamics for females and males, respectively, are
 given by Eq. (4) in the main text (here in recursion form):

$$\begin{aligned} p_{F,T+1} &= \frac{p_{F,T} f t}{p_{F,T} f (1 - (1-t)p_{M,T} L) + (1 - p_{F,T})(1 - p_{M,T} L)}, \\ p_{M,T+1} &= \frac{p_{F,T} f t}{p_{F,T} f (1 + (1-t)\frac{k}{1-k} p_{M,T} L) + (1 - p_{F,T})(1 + \frac{k}{1-k} p_{M,T} L)}. \end{aligned} \tag{B12}$$

652 Therefore, if we define,

$$g(p_{F,T}, p_{M,T}) = \frac{ft}{p_{F,T}f(1 - (1-t)p_{M,T}L) + (1 - p_{F,T})(1 - p_{M,T}L)},$$

$$h(p_{F,T}, p_{M,T}) = \frac{ft}{p_{F,T}f(1 + (1-t)\frac{k}{1-k}p_{M,T}L) + (1 - p_{F,T})(1 + \frac{k}{1-k}p_{M,T}L)},$$

then we can write the recursion equations (B12) in the matrix form given above by
 654 equation (B8).

The non-trivial equilibria of this model are given by Eq. (A3) and repeated here:

$$\hat{p}_F = \frac{2 + f^2(1+k)t - f[2 + t(1+k + L(k-1))]}{2\left[1 - f\left[2 + Lt(k-1) - f(1 - Lt(k-1)(t-1))\right]\right]}$$

$$\pm \frac{ft\sqrt{(k-1)\left[(k-1)(1-f+L)^2 + 4L(k-1 + fk(t-1))(ft-1)\right]}}{2\left[1 - f\left[2 + Lt(k-1) - f(1 - Lt(k-1)(t-1))\right]\right]}$$

$$\hat{p}_M = \frac{1 - f - k + fk + L - kL}{2L(1 - k + fk(1-t))}$$

$$\pm \frac{\sqrt{(k-1)\left[(k-1)(1-f+L)^2 + 4L(k-1 + fk(t-1))(ft-1)\right]}}{2L(1 - k + fk(1-t))}$$

656 As before, at the trivial equilibrium, $\mathbf{p}_0 = (0, 0)$, $g(\hat{\mathbf{p}}) = h(\hat{\mathbf{p}}) = ft$, and therefore

$$\mathbf{M} = \begin{pmatrix} ft & 0 \\ ft & 0 \end{pmatrix}.$$

Again the eigenvalues of this Jacobian matrix are 0 with eigenvector $(p_F, p_M) = (0, 1)$,
 658 and ft with eigenvector $(p_F, p_M) = (1, 1)$. The zero eigenvalue is associated with per-
 turbations in the directions of males only. The largest eigenvalue of the $\mathbf{p}_0 = (0, 0)$

660 equilibrium is ft . Similarly to the diplodiploid model, the trivial equilibrium is **there-**
fore stable if $ft < 1$, and unstable if $ft > 1$.

662 For the non-trivial equilibrium, we need to calculate the derivatives in the formula for
the Jacobian again (Eq. (B9)):

$$\begin{aligned}
664 \quad \left. \frac{\partial g}{\partial p_F} \right|_{\hat{\mathbf{p}}} &= \frac{1}{ft} (1 - \hat{p}_M L + f(1-t)\hat{p}_M L - f), \\
\left. \frac{\partial g}{\partial p_M} \right|_{\hat{\mathbf{p}}} &= \frac{1}{ft} L (1 - \hat{p}_F + \hat{p}_F f(1-t)), \\
666 \quad \left. \frac{\partial h}{\partial p_F} \right|_{\hat{\mathbf{p}}} &= \frac{1}{ft} \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \left(1 + \frac{k}{1-k} \hat{p}_M L - f - (1-t) \frac{k}{1-k} f \hat{p}_M L \right), \\
\left. \frac{\partial h}{\partial p_M} \right|_{\hat{\mathbf{p}}} &= \frac{1}{ft} \frac{kL}{1-k} \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 (\hat{p}_F - \hat{p}_F f(1-t) - 1).
\end{aligned}$$

668 Using $g(\hat{p}_F, \hat{p}_M) = 1$ and $h(\hat{p}_F, \hat{p}_M) = \frac{\hat{p}_M}{\hat{p}_F}$, we can simplify

$$\begin{aligned}
\left. \frac{\partial g}{\partial p_F} \right|_{\hat{\mathbf{p}}} &= \frac{1}{ft} (1 - ft - \hat{p}_M L), \\
670 \quad \left. \frac{\partial h}{\partial p_F} \right|_{\hat{\mathbf{p}}} &= \frac{1}{ft} \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \left(1 - \frac{\hat{p}_F}{\hat{p}_M} ft + \frac{k}{1-k} \hat{p}_M L \right).
\end{aligned}$$

672 Putting it all together gives the following expression for the Jacobian matrix,

$$\mathbf{M} = \begin{pmatrix} \frac{1}{ft} (1 - \hat{p}_M L) & \frac{1}{ft} L \hat{p}_F (1 - \hat{p}_F + \hat{p}_F f(1-t)) \\ \frac{1}{ft} \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \left(1 + \frac{k}{1-k} \hat{p}_M L \right) & \frac{1}{ft} \frac{k}{1-k} \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 L \hat{p}_F (\hat{p}_F - \hat{p}_F f(1-t) - 1) \end{pmatrix}.$$

The eigenvalues of this Jacobian are given by the solution to

$$\lambda^2 - \lambda \frac{1}{ft} \left(1 - \hat{p}_M L - \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \frac{kB}{1-k} \right) - \frac{1}{(ft)^2} \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \frac{B}{1-k} = 0,$$

674 where

$$B = L\hat{p}_F(1 - \hat{p}_F + \hat{p}_F f(1 - t))!$$

The eigenvalues are therefore equal to

$$\lambda = \frac{1}{2ft} \left(1 - \hat{p}_M L - \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \frac{kB}{1-k} \right) \pm \frac{1}{2ft} \sqrt{\left(1 - \hat{p}_M L - \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \frac{kB}{1-k} \right)^2 + 4 \left(\frac{\hat{p}_M}{\hat{p}_F} \right)^2 \frac{B}{1-k}}$$

676 As with the female-killing model, we could not determine the stability of the equilibria analytically. Instead, we conducted similar numerical analysis in Python with
 678 100 values for $\{0.01 \leq L, t \leq 1\}$, 300 values for $\{0.01 \leq f \leq 3\}$, and three values of
 $k \in \{0.5, 0.7, 0.9\}$, resulting in 9 000 000 parameter combinations in total. We calculated
 680 the values of the equilibria and determined their stability based on the larger absolute
 value of the eigenvalues above, whenever $0 < \hat{p}_M, \hat{p}_F \leq 1$. Again, the equilibria were
 682 unstable (\hat{p}_1) and stable (\hat{p}_2) as expected, whenever they occurred within the biologically
 meaningful range and the leading eigenvalue had absolute value different from 1.

684 **Appendix C: *Wolbachia* frequency is higher in**
females than in males in the haplodiploid models

686 Here we show that in the haplodiploid models presented in the manuscript, the infection frequency in females is never lower than infection frequency in males.

688 **Female killing effect**

We start from the remark that $g(\hat{p}_F, \hat{p}_M) = 1$. Hence,

690
$$g(\hat{p}_F, \hat{p}_M) = \frac{ft}{\hat{p}_F f (1 - \hat{p}_M L (1 - t)) + (1 - \hat{p}_F)(1 - \hat{p}_M L)} = 1 \quad (C1)$$

$$\Leftrightarrow \hat{p}_F f + 1 - \hat{p}_F - \hat{p}_F \hat{p}_M f (1 - t) L - \hat{p}_M L (1 - \hat{p}_F) = ft.$$

Next, we use $h(\hat{p}_F, \hat{p}_M) = \frac{\hat{p}_M}{\hat{p}_F}$, namely

$$h(\hat{p}_F, \hat{p}_M) = \frac{ft}{\hat{p}_F f + (1 - \hat{p}_F)} = \frac{\hat{p}_M}{\hat{p}_F}$$

$$\Leftrightarrow \hat{p}_F f + 1 - \hat{p}_F = \frac{\hat{p}_F}{\hat{p}_M} ft.$$

692 Substituting the latter row into Eq. (C1) yields

$$\frac{\hat{p}_F}{\hat{p}_M} ft - \hat{p}_F \hat{p}_M f (1 - t) L - \hat{p}_M L (1 - \hat{p}_F) = ft,$$

and therefore

694
$$\frac{\hat{p}_F}{\hat{p}_M} = 1 = \frac{1}{ft} (\hat{p}_F \hat{p}_M f (1 - t) L + \hat{p}_M L (1 - \hat{p}_F)). \quad (C2)$$

The right-hand side of Eq. (C2) cannot be negative with valid parameter values (including $f > 1$). Therefore $\frac{\hat{p}_F}{\hat{p}_M} \geq 1$, which means that the infection frequency in females is always higher than or equal to male infection frequency.

698 Masculinization effect

We start again from $g(\hat{p}_F, \hat{p}_M) = 1$, and $h(\hat{p}_F, \hat{p}_M) = \frac{\hat{p}_M}{\hat{p}_F}$. The first equality gives

$$\begin{aligned}
 g(\hat{p}_F, \hat{p}_M) &= \frac{ft}{\hat{p}_F f (1 - (1-t)\hat{p}_M L) + (1 - \hat{p}_F)(1 - \hat{p}_M L)} = 1 \\
 &\Leftrightarrow \hat{p}_F f (1 - (1-t)\hat{p}_M L) + (1 - \hat{p}_F)(1 - \hat{p}_M L) = ft,
 \end{aligned}
 \tag{C3}$$

and the second gives

$$\begin{aligned}
 h(\hat{p}_F, \hat{p}_M) &= \frac{ft}{\hat{p}_F f \left(1 + (1-t)\frac{k}{1-k}\hat{p}_M L\right) + (1 - \hat{p}_F) \left(1 + \frac{k}{1-k}\hat{p}_M L\right)} = \frac{\hat{p}_M}{\hat{p}_F} \\
 &\Leftrightarrow \hat{p}_F f \left(1 + (1-t)\frac{k}{1-k}\hat{p}_M L\right) + (1 - \hat{p}_F) \left(1 + \frac{k}{1-k}\hat{p}_M L\right) = ft \frac{\hat{p}_F}{\hat{p}_M}.
 \end{aligned}
 \tag{C4}$$

Subtracting the last row of Eq. (C3) from the last row of Eq. (C4) yields

$$\begin{aligned}
 ft \left(\frac{\hat{p}_F}{\hat{p}_M} - 1 \right) &= \hat{p}_F \hat{p}_M f (1-t)L \left(1 + \frac{k}{1-k}\right) + \hat{p}_M L (1 - \hat{p}_F) \left(1 + \frac{k}{1-k}\right) \\
 &= \frac{1}{1-k} (\hat{p}_F \hat{p}_M f (1-t)L + \hat{p}_M L (1 - \hat{p}_F)),
 \end{aligned}$$

and therefore

$$\frac{\hat{p}_F}{\hat{p}_M} - 1 = \frac{1}{ft} \frac{1}{1-k} (\hat{p}_F \hat{p}_M f (1-t)L + \hat{p}_M L (1 - \hat{p}_F)).$$

The right-hand side above is never below zero for any valid parameter values, and therefore $\frac{\hat{p}_F}{\hat{p}_M}$ is never less than one. Hence, the female infection frequency is always higher than or equal to the male infection frequency.

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
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