A Re-Examination of Wolbachia-Induced Cytoplasmic Incompatibility in California Drosophila simulans

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Abstract

Background: In California Drosophila simulans, the maternally inherited Riverside strain Wolbachia infection (wRi) provides a paradigm for rapid spread of Wolbachia in nature and rapid evolutionary change. wRi induces cytoplasmic incompatibility (CI), where crosses between infected males and uninfected females produce reduced egg-hatch. The three parameters governing wRi infection-frequency dynamics quantify: the fidelity of maternal transmission, the level of cytoplasmic incompatibility, and the relative fecundity of infected females. We last estimated these parameters in nature in 1993. Here we provide new estimates, under both field and laboratory conditions. Five years ago, we found that wRi had apparently evolved over 15 years to enhance the fecundity of infected females; here we examine whether CI intensity has also evolved.

Methodology/Principal Findings: New estimates using wild-caught flies indicate that the three key parameters have remained relatively stable since the early 1990s. As predicted by our three-parameter model using field-estimated parameter values, population infection frequencies remain about 93%. Despite this relative stability, laboratory data based on reciprocal crosses and introgression suggest that wRi may have evolved to produce less intense CI (i.e., higher egg hatch from incompatible crosses). In contrast, we find no evidence that D. simulans has evolved to lower the susceptibility of uninfected females to CI.

Conclusions/Significance: Evolution of wRi that reduces CI is consistent with counterintuitive theoretical predictions that within-population selection on CI-causing Wolbachia does not act to increase CI. Within taxa, CI is likely to evolve mainly via pleiotropic effects associated with the primary targets of selection on Wolbachia, i.e., host fecundity and transmission fidelity. Despite continuous, strong selection, D. simulans has not evolved appreciably to suppress CI. Our data demonstrate a lack of standing genetic variation for CI resistance in the host.

Introduction

Wolbachia are maternally inherited intracellular bacteria found in many – and perhaps most – arthropods [1]. The Riverside strain of Wolbachia (wRi) was identified in Drosophila simulans in the 1980s [2] in southern California, USA. The infection was initially identified by a reduced hatch rate in crosses between males from southern California (Riverside, CA) and females from central California (Watsonville, CA). The reciprocal cross produced significantly more progeny, comparable to the number produced by infected pairs or uninfected pairs. This cytoplasmically inherited incompatibility was associated with Wolbachia, first microscopically [3], then via PCR [4]. In laboratory tests, the infection produced no statistically significant fitness consequences other than a reduction in fecundity relative to uninfected flies [5]. Through the late 1980s and early 1990s, wRi spread rapidly into central and northern California [2,4–7]. The wRi infection is now pervasive in California D. simulans populations, and its frequency has remained relatively stable after the initial spread [8].

The reduced egg-hatch produced by matings between infected males and uninfected females is termed cytoplasmic incompatibility (CI), and it seems to be the most common reproductive manipulation caused by Wolbachia [9,10]. CI and long-distance dispersal produced the rapid spread of wRi within and among California D. simulans populations [2]. A simple population model of CI explained the observed within-population dynamics and equilibrium frequency of wRi in terms of three parameters (see Table 1 of [11]): the maternal transmission rate, $1−μ$, where $μ$ is the fraction of uninfected ova produced by infected females; the relative fecundity of infected females compared to uninfected females, $F$; and the average hatch rate from incompatible fertilizations, $H$, i.e., the relative hatch rate when uninfected ova are fertilized by sperm from infected males. Studies in the early 1990s used repeated experiments with wild-caught flies to estimate these parameters in nature; the average values were $μ=0.045$, $F=0.97$, and $H=0.73$. The wRi infection produces incompatibility among crosses when maternal transmission fidelity is high, but lower transmission fidelities result in less incompatibility. Here we re-examine the nature of wRi infection-frequency dynamics and the underlying variation in CI.
F=1.00, and H=0.55 [4]. While estimates of fecundity in the field generally showed no statistically significant difference between infected and uninfected females, laboratory studies demonstrated a 10–20% fecundity reduction in infected females [5]. Although maternal transmission of aRi in perfect in our laboratory stocks, studies of transmission in the field demonstrated heterogeneity among infected females, with many producing all infected progeny, but many others producing a significant fraction of uninfected progeny [4].

Theory predicts evolution towards a more mutualistic symbiosis, involving changes in one or more of the three key parameters [11]. Selection acts on both the host and Wolbachia to foster mutualism. For Wolbachia variants that remain mutually compatible, natural selection acts to increase the average number of infected progeny produced by infected females, \( F(1-\mu) \) [11,12]. Under Wolbachia evolution, the intensity of cytoplasmic incompatibility, as measured by \( H \), might also change, but via pleiotropic effects rather than direct selection [11–13]. The effects of natural selection on the host are more complex; but in the absence of pleiotropy, it should act to increase maternal transmission, 1–\( \mu \); increase female fecundity, \( F \); and suppress CI, i.e., increase \( H \). The CI phenotype has been described in terms of two bacterial components: the modification of sperm that induces embryo death in incompatible crosses, and the rescue capability of infected eggs, restoring the modification [14]. Both can be influenced by host background effects and Wolbachia [10]. Given that both infected and uninfected individuals share the same nuclear genes, a host-mediated evolutionary shift in hatch rate might be the product of changes in males or females. This could involve suppression of CI by the male host genome, or increased resistance to CI by the female genome. We examine both.

Experiments conducted by Weeks et al. [8] indicated that aRi has evolved to increase fecundity. Isofemale lines collected between 2002 and 2004, but maintained in the lab for several generations, showed significant variation in fecundity. Moreover, several showed statistically significant decreases in fecundity when their Wolbachia were removed with tetracycline (whereas consistent increases in fecundity with Wolbachia removal had been observed in previous studies [5]). One line in which Wolbachia increased fecundity, IR2, was chosen for further investigation. Reciprocal introgressions moved Wolbachia from Riv84 and Riv88 into the alternative nuclear backgrounds. The results demonstrated that IR2 Wolbachia significantly raised fecundity on both nuclear backgrounds. Weeks et al. [8] found no evidence for CI between the alternative Wolbachia, suggesting that aRi had evolved in the field to become more mutualistic. This interpretation was also supported by the complete sequence identity between the IR2 and Riv88 Wolbachia at the rapidly evolving wsp locus, which is routinely used for phylogenetic placement of Wolbachia strains [15]. CI induction levels across three male ages (5, 10 and 15 days old) in the new flies were not significantly different from those previously reported [4].

In this study, we estimated the current aRi infection frequency in northern California and re-estimated the three key population dynamic parameters: the intensity of cytoplasmic incompatibility (\( H \)), the fidelity of maternal transmission (\( \mu \)), and the relative fecundity of infected females (\( F \)). We also asked whether there are differences in CI induction and resistance levels between newly collected flies from the field versus infected and uninfected reference stocks and determined whether observed changes were attributable to the host and/or Wolbachia. Our results are discussed in light of theoretical predictions [11].

### Materials and Methods

#### Fly stocks and PCR

Four reference *D. simulans* mass-bred laboratory stocks were used: two aRi infected stocks, Riv84 and Riv88, and two naturally uninfected stocks, W88 and Coffs80. Riv84 and Riv88 originated in 1984 and 1988 from Riverside, in southern California, the type location for the aRi infection [2]. W88 originated in 1988 from Watsonville, in central California, our type location for uninfected flies before the aRi infection spread northward [2,4]. Uninfected Coffs80 originated in 2008 from Coffs Harbour, near the center of the Australian east coast distribution of *D. simulans*.

Collections of *D. simulans* were obtained in 2008 and 2009 from three locations in the Central Valley of California. All three are rural orchards within 10 km of each other, between the towns of Davis and Winters, and all are within 500 m of a riparian corridor, Putah Creek, that separates Yolo and Solano counties. The aRi infection spread northward into this region in the late 1980s [4]. These collections have been named according to their location and year (in subscript). In 2008, we pooled flies from all three orchards and denoted them D157; in 2009, we collected only from a Yolo county orchard and denote the sample Y09. When referring to a specific line from these collections, the line number is given in place of the collection year, e.g., line 157 collected in Winters in 2008 is D157. A letter following the line number indicates *Wolbachia* transmission characteristics, ‘P’ indicates partial transmission (i.e., some uninfected isofemale sublines were obtained), and ‘U’ indicates an uninfected line. For all field assays, flies were crossed to reference stocks or placed individually in vials to create isofemale lines within hours of field collection.

#### Infection status via PCR

To screen for *Wolbachia*, we used a rapid method of DNA isolation for PCR amplification. A 120 µl aliquot of buffer (10 mM TrisHCl, 1 mM EDTA, 25 mM NaCl, with 0.005% Protease K) was used for grinding individual flies for two minutes in 1.1 ml deep-well plates (Axygen, USA) using 3 mm glass beads in either a

| Assay                  | Collection month (2008) | N  | Infection frequency | 95% binomial confidence interval |
|------------------------|-------------------------|----|---------------------|---------------------------------|
| Maternal Transmission  | June/July               | 146| 0.918               | (0.861, 0.957)                  |
| CI (assay 1)           | July                    | 176| 0.938               | (0.891, 0.968)                  |
| CI (assay 2)           | August                  | 222| 0.937               | (0.896, 0.965)                  |
| Fecundity              | September               | 538| 0.931               | (0.906, 0.951)                  |
| **Total**              |                         | 1082| **0.932**           | **(0.915, 0.946)**             |

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CI levels induced by lab-reared D08 males were compared to those induced by our laboratory stock, Riv88. Five D08 isofemale lines, assayed in this study for maternal transmission of Wolbachia, were tested. Three of the five showed "perfect" transmission (i.e., produced no uninfected sublines) (D157, D351 and D352), and two showed "partial" (<100%) transmission (D29P and D287P). The ability of five uninfected D08 isofemale lines (D350U, D362U, D365U, D372U and D373U) to resist CI was compared to that of W88 females.

Our experimental design for CI induction involved mating males from Riv88 and each of the five D08 infected lines to W88 females. We also mated males from each of the infected D08 lines to females from one of the five uninfected D08 lines. Conversely, females from W88 and each of the D08 uninfected lines were mated to Riv88 males to test for variation in susceptibility to CI. Crosses within each of the five infected and uninfected D08 lines were included as controls.

For these CI assays, flies were reared at a controlled density prior to testing. Groups of 25 eggs were collected from treacle-based media with a yeast suspension and transferred to fresh vials containing 10 ml of cornmeal medium. CI was tested in virgin males aged 7 and 14 days. Females were between three and five days of age. We performed 20 replicates of each incompatible cross, and 10 replicates of each compatible cross. The D08 flies had been reared in the lab for a maximum of four generations. As in the field CI assays, females were allowed to deposit eggs on spoons for 48 hours (spoons changed after 24 hours), and hatch for a further 24 hours, before being frozen and later counted to determine hatch rate.

**Assay 2: CI induction using Y09 isofemales.** We also assessed variation in male-associated CI induction using our Y09 lines. A total of 100 isofemale lines, established from wild-caught females, were tested. The F1 males were collected as virgins from each line and aged seven days. One generation later, we retested the level of CI produced by 14 of these lines, 7 that initially showed the highest levels of CI and 7 that initially showed the lowest levels of CI. The experimental design followed our lab CI assays, with between 5 and 10 replicate crosses per line.

**Assay 3: retest CI induction of Y09 isofemales in Melbourne.** The 14 lines initially identified as high versus low CI in Assay 2 were shipped to Melbourne for further testing. One of the seven initial low-CI lines was lost. A subset of the Y09 isofemale lines was retested with increased replication in Melbourne to assess whether the differences observed in Davis were repeatable and attributable to host versus Wolbachia effects. Again, these males were placed with virgin females of the W88 line. The flies were tested about ten generations after being removed from the field. As elaborated below, three lines from the initial “high” CI group and three lines from the initial “low” CI group were used in reciprocal crosses with Riv88 to test for maternally inherited variation in CI induction, while controlling for host nuclear genetic effects.

**Host versus Wolbachia contributions to CI variation**

We performed two assays to determine whether variation among males from D08 and Y09 lines in CI induction and differences from our old Riverside stocks were most likely attributable to Wolbachia (cytoplasmic) or Drosophila (nuclear) effects. Each assay involved introgression or reciprocal crosses. As discussed below, these designs do not directly distinguish Wolbachia effects from other maternally inherited factors.

**Introgression assays between Riv88 and D157.** Our assay of male-induced CI variation in D08 resulted in the identification of line D157, which produced only infected sublines, yet induced less intense CI than Riv88. Reciprocal backcrossing was used to

**CI assays using field-collected males**

To quantify CI intensity, field-collected males were brought into the laboratory and mated to reference laboratory stocks within hours of collection. Over two independent assays (initiated 23 July 2008 and 5 August 2008), 500 wild-caught males were tested. Each assay began with 250 field-collected males. The four sets of crosses undertaken in both assays were identical: (1) 200 field males mated to W88 females on the first day, then mated to R88 females on the second day; (2) 50 field males mated to W88 females on two successive days to test the effects of remating on CI [17,18]; (3) 20 W88 males mated to W88 females on the first day and Riv88 females on the second day to determine whether egg hatch from compatible crosses depended significantly on the origin of the uninfected male (and to look for inbreeding effects); and (4) a second group of 10 W88 males mated to W88 females on two successive days (to further check for inbreeding effects). After 24 hours with the second female, males were removed and frozen at −80°C for later testing of infection status.

Females were aged between 3–5 days after eclosion before mating. Females were allowed to lay eggs on spoons for 48 hours, with fresh spoons provided after 24 hours. The spoons contained molasses medium with a live yeast suspension to encourage egg-laying. After each 24-hour laying period, all eggs laid on the spoons were left 24 hours for hatching before the spoons were frozen. The number of eggs laid by each female was counted, along with the number of hatched eggs (or larvae). Hatch rates were compared between incompatible and compatible crosses and among classes of compatible controls. We excluded from the analyses egg data from females that laid fewer than 10 eggs over two days and data from wild-caught males whose infection status could not be confirmed.

**CI assays comparing newly collected stocks to Riv88 and W88**

We performed three sets of assays to determine whether CI levels had evolved.

**Assay 1: CI induction and susceptibility using D08 isofemales.** Field-collected females were used in 2008 to assess maternal transmission of Wolbachia. Some of the resulting infected and uninfected isofemale lines were subsequently assayed for CI to determine: (1) whether infected males from D08 stocks produced CI levels that differed from our reference Riv88 stock, and (2) whether females from uninfected D08 stocks differed from our reference W88 stock in their susceptibility to CI.
place the *Wolbachia* from Riv88 versus D157 onto one another’s nuclear genetic background. The reciprocal replacements were denoted D157\(^{Kw88}\) and Riv88\(^{D157}\), where the superscript denotes the *Wolbachia* source. Four generations of backcross resulted in a predicted replacement of over 93% of the host nuclear background. The backcross lines and the parentals (i.e., Riv88\(^{Kw88}\) and D157\(^{D157}\)) were assayed together.

CI assays used seven-day-old males. As with the other lab-based CI assays, single virgin males and females were placed together in a vial to mate, and the female was allowed to deposit eggs for 24 hours. The females were three to four days old. For each line, males were crossed to both W88 and Riv88 females. There were eight to ten replicates per cross. A second assay using these same lines was performed three generations after the first, with 16 to 30 replicates for each incompatible cross. **Reciprocal cross assays using Y\(_0\) lines and Riv84.** We assayed CI with F1s from reciprocal crosses between six of the initially screened Y\(_0\) lines and Riv84. There were seven to eight replicates of each cross type, and only the replicates that produced 10 or more eggs were included in the analyses. F1 males were mated to uninfected Coffs88 females to produce incompatible crosses.

**Maternal Transmission**

Maternal transmission of *Wolbachia* was assayed twice using wild-caught D88 females, following an improved version of the Turelli and Hoffmann [4] protocols. More than 100 *D. simulans* females were collected over one month, from mid-June to mid-July 2008. The frequency of uninfected individuals produced by infected females was determined by establishing multiple F1 sublines from the wild-caught females, then scoring the F2, as described below. Each field-collected female was allowed to lay eggs for around six days in the lab prior to mating her to an uninfected reference male (W88), and then allowing her to lay eggs for another six days. By remating field-collected females to W88 males, we retested the hypothesis that uninfected ova produced by an infected mother show the same susceptibility to CI as the ova from uninfected mothers [4]. This susceptibility must be considered to accurately assess *μ* because the high elevation in California means that most fertilizations involve sperm from infected males. If an infected female produces uninfected ova, these ova may be subject to CI and not hatch. This would lead to underestimation of *μ* from the fraction of uninfected sublines. Mating wild-caught females to uninfected males should produce more accurate estimates.

Each field female’s F1 progeny were aged and allowed to mate with each other, both before and after the field-caught female was mated to the W88 male, and up to ten already mated F1 females were placed individually in fresh vials to establish F2 sublines. Initially, one F2 female from each subline was tested for infection status. A subline was scored as infected if the individual tested positive for *Wolbachia*. Sublines were scored as uninfected only after four additional females from the same subline were found to be uninfected. The percentage of infected F2 progeny was estimated for each field female based on the fraction of uninfected versus infected sublines produced before and after mating with a W88 male.

After the initial maternal transmission experiment, ten D88 lines, five infected and five uninfected, were selected for continued maintenance and used in the CI comparisons discussed previously under “Assay 1: CI induction and susceptibility using D88 isofemales”. These five infected lines consisted of three *Wolbachia*-infected lines that produced 100% infected sublines and two that produced <100% infected sublines.

**Fecundity in the field**

Relative fecundity of infected and uninfected wild-caught females was estimated using flies collected in August 2006. A total of 538 *D. simulans* females from the field was obtained and assayed for fecundity over four days. Single females were placed in vials containing spoons, replaced daily. Spoons were frozen until the eggs could be counted. The F1 progeny of these lines were used to test the infection status of the field females. We discarded data from females who died during the four-day egg-laying period.

Body size and fecundity are typically positively correlated in *Drosophila* (e.g. [19]). Thorax measurements were made for 441 field females, as an index of body size. The thorax of each female was photographed and measured using the program AutoMontage, after storage in 70% ethanol. Data on thorax length and fecundity for all field females were used to look for associations between them and for effects of *Wolbachia* on size or fecundity.

**Infection frequency**

We estimated the incidence of *Wolbachia* in the field in northern California in 2008. We tested 398 field-collected males from the field CI assay and 538 field-collected females from the fecundity assay. We also assessed the infection status of 146 field females collected for our maternal transmission study by examining their F1 male progeny. In this case, when a negative PCR result was initially obtained for an isofemale line, four more males were tested.

**Statistical analyses**

For all CI assays, hatch rates were computed only when females produced at least 10 eggs. For the initial assays involving field males and comparisons to laboratory males, data were often not normally distributed, so nonparametric tests were used in all initial surveys of lines. In experiments where the same males were mated to different females on successive days, we undertook pairwise comparisons of percentage hatch rates using Wilcoxon paired tests to test for differences in CI levels in assays on day 1 versus day 2. For the remaining comparisons of CI, we considered different crosses and these were compared with nonparametric Mann-Whitney tests (for two-group comparisons) or Kruskal-Wallis tests (for more than two groups). For estimates of CI intensity (*H*) and failure of maternal transmission (*μ*), bias-corrected and accelerated (BC\(_a\)) bootstrap confidence intervals were obtained based on 10,000 bootstrap replicates [see 4] and Ch. 14 of [20].

In CI assays following crosses between different strains to determine the effects of *Wolbachia* or nuclear background, hatch rate data were normally distributed; and we used parametric tests to compare host and *Wolbachia* background effects. ANOVAs were used to separate these effects.

To compare infection frequencies across collections, we used contingency tables and computed the chi-square statistic. Exact confidence intervals based on the binomial distribution were computed for the resulting frequency estimates [4].

**Results**

**Cytoplasmic incompatibility (CI) assays with field-collected males**

**Field assay 1.** As noted above, there were three classes of males: (1) 200 field males mated to W88 females on the first day, then mated to R88 females on the second day; (2) 50 field males mated to W88 females on two successive days; (3) 20 W88 males mated to W88 females on the first day and Riv88 females on the second day. We included only individual crosses that produced at least 10 eggs. The first two groups of males were used to estimate
levels of CI, controlling for male fertility and examining the effects of male age and/or remating, while the last cross tested the fitness of the W88 line, looking for possible inbreeding effects. We also included crosses between W88 males and W88 females on both days, but hatch rates from these crosses did not differ significantly in either assay and are not presented.

As expected for the first class of males, matings on the first day, between infected males and uninfected females, produced a lower hatch rate (38%, N = 41) than matings on the second day between the same infected males and infected females (75%). Considering only males who produced at least 10 offspring from each cross, the difference is significant in a paired Wilcoxon test on hatch rates (Z = 4.20, P < 0.001). In contrast, for uninfected field males, there was no significant difference (Z = 0.674, P = 0.50) between hatch rates from day 1 matings with uninfected females (65%) and day 2 matings with infected females (75%), although there were only five data points for comparison. When all crosses are compared by a Kruskal-Wallis test including those that produced sufficient offspring on either day 1 or day 2, there was no difference in CI between the three sets of compatible crosses involving field males (X² = 0.37, df = 2, P = 0.831); these consisted of uninfected males mated to uninfected (hatch rate 71%, N = 6) or infected females (73%, N = 9) and infected males mated to infected females (80%, N = 131). We combined all three classes of compatible crosses involving field males to estimating field CI levels. However, for the incompatible crosses, we considered only those from day 1, because CI levels decline with age [4] and remating [17], and a difference between days was detected in assay 2 in paired crosses (below).

Using our pooled compatible crosses, we obtain a point estimate of H, the relative hatch rate from incompatible crosses, of 0.42, with a bias corrected and accelerated (BCα) [20], Ch. 14) 95% bootstrap confidence of (0.33, 0.53).

In the third set of crosses, a Mann-Whitney U test indicated an unexpected statistically significant difference (Z = 2.098, P = 0.036) between hatch rates for W88 males mated to W88 females on days 1 (93%, N = 12) and Riv88 females on day 2 (83%, N = 15). We assign no biological significance to this anomalous result, which was not repeatable in our second field assay.

**Field assay 2.** For the first set of males, infected males produced a lower hatch rate (Z = 9.74, P < 0.001) when mated to uninfected females on day 1 (26%, N = 130) than when mated to infected females on day 2 (86%), as expected with CI. For the uninfected field males, there was no difference (Z = 1.57, P = 0.116) between days 1 (75%, N = 7) and 2 (86%), as expected for compatible crosses. For the second set of infected field males mated to uninfected females on days 1 and 2, there was a significant difference by paired Wilcoxon test (Z = 3.79, P < 0.001) between days due to a lower hatch rate on day 1 (37%, N = 40) compared to day 2 (49%). This increase may be attributable to either male age or remating. We also had five uninfected males in the second set of crosses that produced offspring on both days, and these did not differ significantly between days (Z = 0.67, P = 0.50).

As for assay 1, we combined all three classes of compatible crosses in estimating field CI rates. However, there was a significant difference among the classes (X² = 29.83, df = 3, P < 0.001), because there was a lower hatch rate in crosses between uninfected field males and uninfected females on day 1 (66%, N = 12) and day 2 (59%, N = 5) compared to crosses on day 2 involving infected field males and infected females (84%, N = 156) and uninfected field males and infected females (87%, N = 9). The reason for these differences is unclear unless some field males scored as uninfected carried a low level Wolbachia infection that was not detected.

For the compatible cross, we considered only males that produced at least 10 offspring on day 1. These led to a point estimate for the intensity of CI in the field of H = 0.35 and a 95% BCα bootstrap confidence interval of (0.30, 0.40).

For the third set of crosses with W88 males, the Mann-Whitney U test was not significant (Z = 0.260, P = 0.793), with a high hatch rate after crosses to both W88 females on day 1 (81%, N = 14) and Riv88 females on day 2 (80%, N = 14).

**CI assays comparing newly collected stocks to Riv88 and W88**

We performed three sets of assays; the first assay addressed two separate issues, discussed in succession below.

**Assay 1a: CI induction by infected males, D08 lines versus Riv88.** We compared levels of CI induced at ages 7 and 14 days by Riv88 males versus males from five infected D08 lines. The D08 lines were selected based on their maternal transmission characteristics (see section on Maternal Transmission below). We used three lines showing perfect transmission and two showing partial transmission, as assessed using isofemale lines established directly from the field. Figure 1 presents the data from all of our incompatible crosses, distinguishing those that involved W88 females (grey bars) versus uninfected D08 females (white bars). None of the adjacent grey and white bars (reflecting a specific infected male line mated to two different uninfected female lines) differ significantly (statistics not shown). Hence, our comparisons below pool the results involving W88 and uninfected D08 females. This lack of variation in uninfected female susceptibility to CI is elaborated below.

For day 7, as expected, incompatible crosses produced consistently lower hatch rates (average 29%) than compatible crosses (73%) [Mann-Whitney: Z = −11.308, P < 0.001]. More interestingly, Figure 1A shows that Riv88 males had a lower average hatch rate, 16.9% (N = 102), than males from the five new infected lines, which averaged 37.8% (N = 178). There was a significant effect of male line (Kruskal-Wallis: X² = 72.27, df = 5, P < 0.001) and significant heterogeneity among the five infected D08 lines (Kruskal-Wallis: X² = 47.03, df = 4, P < 0.001).

Two of the five infected D08 lines assayed, D29P and D287P, showed incomplete maternal transmission. Assuming the males inherited relatively few or no Wolbachia [see [4]], we expected them to induce less intense CI than did the lines with perfect transmission. As expected, these two lines produced the highest hatch rates of the five D08 lines tested (Figure 1A). The perfectly transmitting and partially transmitting D08 lines are different from each other. Overall, the perfect lines have higher CI than the partial lines (Z = 6.028, P < 0.001). However, even the perfectly transmitting lines produced higher hatch rates than Riv88. Because D157 produced the lowest level of CI among the perfect transmission lines, it was chosen for further study. At age 7, D157 had a mean hatch of 0.335, with a 95% bootstrap confidence interval of (0.276, 0.403), versus 0.188 (0.150, 0.224) for Riv88.

At day 14, CI induced by infected males declined significantly, as expected [2]. The hatch rate of the incompatible crosses increased (mean 0.586, 95% confidence interval (0.556, 0.615)), but remained significantly lower than that observed for compatible crosses (0.721, (0.665, 0.770)) (Mann-Whitney: Z = −5.59, P < 0.001). Male lines still differ in their CI intensities (Kruskal-Wallis: X² = 39.88, df = 5, P < 0.001).

**Assay 1b: CI susceptibility of uninfected females, D08 lines versus W88.** The susceptibility to CI of recently collected, uninfected isofemale lines was assessed relative to W88 females.
Figure 2 compares the levels of CI induced by Riv88 males, aged 7 and 14 days, on W88 and all five uninfected D08 lines used in Figure 1 (elaborating the comparison shown in the Riv88 columns of Figure 1). As expected from Figure 1, Figure 2 shows that the female line did not influence the hatch rate in incompatible crosses at age 7 (Kruskal-Wallis: \(X^2 = 5.55, \text{df} = 5, \ P = 0.352\)) or 14 (\(X^2 = 4.92, \text{df} = 5, \ P = 0.425\)). Thus, Figure 2 indicates that females from the new uninfected isofemale lines did not differ from W88 in their susceptibility to CI induced by Riv88 males; whereas Figure 1 shows that new (D08) and old (W88) uninfected lines are equally susceptible to newly collected infected males (D08).

Assay 2: CI induction by males from Riv88 versus infected Y09 isofemales. We repeated CI assays in 2009 using newly established Y09 isofemale lines. Virgin F1 males from 100 Y09 lines were screened for CI induction at age seven days. After our initial screening, we maintained 14 isofemale lines for further testing: 7 that indicated the highest level of CI (with mean egg hatch ranging from 0.043 to 0.155, overall mean 0.091) and 7 that indicated the lowest level of CI (mean egg hatch ranging from 0.28 to 0.585, overall mean 0.433). These 14 lines were tested again for seven-day-old CI induction the next generation. The average hatch rates for the seven “low” lines now ranged from 0.102 to 0.354, with an overall mean of 0.254; whereas the seven “high” lines now ranged from 0.232 to 0.637, with an overall mean of 0.344. These two groups differed significantly (one-sided Wilcoxon, \(P = 0.014\)). However, there was considerable variation in the point estimates of CI intensity across the two replicates. If we rank lines as “high” or “low” based on whether the mean hatch rates are in the bottom six or top six, only lines Y6, Y30 and Y41 were consistently high across both tests; and only Y47 and Y83 were consistently low across both tests.

Dissection of variation in male CI-induction

Introggression assays between Riv88 and D157. This assay addressed whether differences in the host, the Wolbachia, or both genomes produced the lower levels of CI for D157 in comparison to the Riv88 (Figure 4). An ANOVA showed a significant effect of Wolbachia (and/or other cytoplasmically inherited factors) \(F_{1, 97} = 6.874, \ P = 0.010\). The mean hatch rate for males carrying Riv88 Wolbachia, irrespective of their host background, was 0.327 (95% bootstrap confidence interval (0.273, 0.386)), while males carrying Wolbachia from the D157 line showed a mean of 0.455 (0.387, 0.523). No effect of host genetic background \(F_{1, 97} = 0.274, \ P = 0.602\) was found. The Riv88 background had a mean hatch rate of 0.409 (0.350, 0.470), and the D157 background had a mean hatch of 0.356 (0.289, 0.428).
Reciprocal-cross assays using \( Y_{09} \) lines and \( \text{Riv84} \). We further explored the effects of maternally inherited factors while controlling for host nuclear background by setting up a single generation of reciprocal crosses between six \( Y_{09} \) lines and Riv84. When all the new lines were considered together and compared to Riv84, we found a significant difference between the reciprocal crosses (\( F_{1,100} = 9.404, P = 0.003 \)), with lower hatch rates associated with Riv84 matrilines. For five of the six lines, means indicate higher hatch rates when the female parent was from a \( Y_{09} \) line versus Riv84; while for the remaining line (\( Y_{54} \)), means were essentially identical between reciprocal crosses (Figure 5). When reciprocal crosses were compared with one-tailed \( t \) tests, there were significant differences between crosses involving \( Y_6 \) (\( t = 2.37, \text{df} = 17, P = 0.015 \)) and \( Y_{30} \) (\( t = 1.95, \text{df} = 15, P = 0.035 \)). The \( Y_6 \) strain had previously shown consistently low CI levels compared to the old Riv strains (see above). There were marginally non-significant differences for crosses with \( Y_{47} \) (\( t = 1.46, \text{df} = 17, P = 0.081 \)) and \( Y_{83} \) (\( t = 1.59, \text{df} = 12, P = 0.079 \)), while the differences were not significant for \( Y_{23} \) (\( t = 1.067, \text{df} = 14, P = 0.150 \)) or \( Y_{54} \) (\( t = 0.76, \text{df} = 15, P = 0.225 \)).

Three of these lines were identified in our initial screen of CI-induction assay 2 as producing high levels of CI (\( Y_6, Y_{30} \) and \( Y_{54} \)) and three were initially identified as producing low levels of CI (\( Y_{23}, Y_{47}, Y_{83} \)). However, when compared in this reciprocal-cross assay, the two groups did not differ significantly. Only \( Y_6 \) showed consistently high CI across both tests in CI-induction assay 2 and CI-induction assay 3 (Figure 3); whereas both \( Y_{47} \) and \( Y_{83} \) were consistently in the lower CI groups in all three tests. Overall, our data indicate a high level of variation in CI intensities across replicate assays.

Maternal transmission

From 477 wild-caught females, 146 isofemale \( D. \text{simulans} \) lines were established. The sample included uninfected females, infected females that produced only infected progeny (“perfect” transmission) and those that produced both infected and uninfected progeny (“imperfect” transmission). Maternal transmission was initially assessed using F1 progeny produced by females directly from nature. Maternal transmission was reassessed after the field females were aged and mated to W88, our reference...
uninfected stock. Only isofemale lines that produced eight or more F2 sublines were used in our analyses.

Using F1 produced from field matings, 77 infected females produced a total of 712 F2 sublines, of which 11 were uninfected. Because most of these field females will have mated with infected males, many of their uninfected ova would be subject to CI. Let \( n \) denote the fraction of uninfected F2 sublines produced by wild-mated, infected females. Our data produce the estimate \( n = 0.015 \), a lower bound for \( m \). Only 3 of the 77 wild-caught, infected females (3.8% of total) produced the 11 uninfected sublines. A more careful estimate involves weighting the data from each female by the number of F2 sublines she produced. This also produces \( r = 0.015 \). Bootstrapping the data from the collections of sublines produced by individual wild-caught females produces a 95% BCa bootstrap confidence interval of (0.003, 0.070). Note that this confidence interval is much broader than the 95% binomial confidence interval, (0.008, 0.027), based on 11 uninfected F2 sublines out of 712. The much larger BCa confidence interval more appropriately accounts for sampling variance because of the extraordinary heterogeneity of maternal transmission among females. Out of 77 wild-caught, infected females that produced at least 8 F2 sublines, 74 females produced only infected F2 sublines. In contrast, of the remaining three females, one produced only two infected sublines out of ten, the other two produced 8/10 and 8/9 infected sublines.

If uninfected ova from infected females are subject to CI, \( v \) underestimates \( m \), the fraction of uninfected ova produced by infected mothers. To estimate \( m \) more accurately, the field-caught females were aged and remated to W88 males in the lab, then F2 sublines were established from the resulting F1. Using F1 produced from W88 matings, 65 infected females produced a total of 588 F2 sublines, of which 29 were uninfected. Only 6 of the 65 W88-remated, infected females (9.2%) produced the 29 uninfected sublines. These data produce a weighted-average estimate of \( \mu = 0.048 \) (unweighted, 0.049) and a 95% BCa bootstrap confidence interval of (0.016, 0.116), based on 10,000 bootstrap replicates (over the groups of sublines produced by individual field-caught females). Again, the bootstrap confidence interval is much larger than the corresponding 95% binomial confidence interval of (0.033, 0.070), because of the degree of heterogeneity in transmission fidelity across W88-remated females. Of the six females that produced both infected and uninfected F2 sublines, two produced only one infected subline out of nine, a third produced three infected out of ten, a fourth produced five infected out of nine, and the remaining two produced 7/8 and 9/10 infected. Hence 4 of 65 wild-caught, infected females accounted for most of the imperfect maternal transmission. The F2 data from infected females remated to W88 males are illustrated in Figure 6.
The most comparable data from Turelli and Hoffmann [4] are from their fourth maternal transmission experiment, reported in their Figure 4. It shows that 15 out of 59 females from nature, remated to uninfected males, produced both infected and uninfected progeny. Those data involved eliminating one female who produced only 1 infected subline out of 11, so overall 16 of 60 females collected in 1993 produced infected and uninfected sublines. In contrast, only 6 of 65 females in 2008 produced both infected and uninfected F2. Using a χ² test, this difference is statistically significant (P<0.05). However, as elaborated in our Discussion, much greater differences were observed between two of the other maternal transmission studies reported in Turelli and Hoffmann [4]. Despite significant heterogeneity among estimates, our new point estimate, μ = 0.048, is similar to the composite estimate, μ = 0.042, produced by Turelli and Hoffmann [4] from four separate experiments.

Although the binomial confidence intervals for our new estimates of υ and μ, from wild-mated versus W88-remated females respectively, do not overlap, the valid BCα confidence intervals do. Hence, even though the corresponding point estimates, 0.015 versus 0.048, are appreciably different, they are not statistically significantly so. (This assertion is supported by a permutation-based test, analogous to that in [21], which asks whether the observed difference in the estimates is consistent with a null distribution of differences obtained by randomly partitioning the subline data pooled from both experiments.) Nevertheless, as discussed below, our data indicate that uninfected ova from infected females are subject to CI, as demonstrated by Figure 4 of [4].

Fecundity

The field assay for fecundity assessed relative egg numbers produced by infected and uninfected females from nature. The fecundity of 538 D. simulans females was tested, over four days. This sample contained 93.12% infected females. During the assay, 35 infected females died, leaving 466 infected and 37 uninfected females. The fecundity of infected and uninfected females did not differ significantly (F1, 501 = 0.96, P = 0.75). Infected females laid on average 13.56 (+0.27 SE) eggs per day, while uninfected females laid 13.87 (+1.02) eggs, producing a point estimate for F, the relative fecundity of infected females, of 0.98, and a 95% BCα bootstrap confidence interval of (0.85, 1.15), based on 10,000 bootstrap replicates.

Thorax length was measured for 441 females. This was significantly correlated with fecundity (r² = 0.294, P<0.001). When thorax length was included as a covariate, it significantly influenced egg production (F1, 439 = 41.71, P<0.001), as expected; but there was no significant difference in the mean thorax lengths of infected and uninfected females (F1, 439 = 0.786, P = 0.376).

Infection Frequency

For our 2008 collections, the infection frequencies were homogeneous across assays (χ² test, 3 df, P>0.8). In a sample of over 1000 (Table 1), we found 93.2% infected with aRi, with a binomial confidence interval of (0.913, 0.946). These data are obviously consistent with the overall infection frequency of 0.93 reported by Weeks et al. [8] from 654 individuals sampled in 2004 from four populations throughout California, including one near Davis and two from southern California.

Discussion

Cytoplasmic Incompatibility

Field estimates. Our new estimates of hatch rates from incompatible fertilizations in the field, H = 0.42 (0.33, 0.53) and H = 0.35 (0.30, 0.40), are lower than many estimates previously reported in California D. simulans, which ranged from 0.32 to 0.71 [4]. However, the one field estimate from Turelli and Hoffmann [4] derived solely from wild-caught males, the procedure used in this study, produced a very similar result: H = 0.43 (0.35, 0.56).

Field studies, male age and mating number cannot be controlled and this contributes to variation in H estimates. A likely factor contributing to our relatively low estimates is that the flies were collected in a commercial peach orchard, in which the fruit was just ripening, and the D. simulans population was expanding. Therefore, the population age structure may have been skewed towards young males, enhancing CI. Our lab studies, discussed below, suggest that recently collected strains of aRi may actually cause less intense CI than found previously.

Lab estimates: Uninfected female susceptibility. Our lab studies tested whether either the host or Wolbachia had evolved to significantly alter CI intensity. Given imperfect maternal transmission, uninfected females persist in all natural populations; and imperfect transmission quickly breaks down associations between nuclear alleles and infection status [22]. Hence, in polymorphic populations, all D. simulans nuclear alleles will find themselves in both infected and uninfected females. Obviously, under these conditions, selection will favor genotypes that suppress CI [11]. Our data (Figures 1 and 2) demonstrate that recently collected females are not able to resist CI induction any more or less than W88 females. Our W88 lab stock has been sheltered from CI for over 20 years, while populations near Davis have been subject to selection to reduce CI since 1990, over 200 generations, assuming that generation times in nature are roughly one month, as estimated in [4].

Our finding that CI-resistance has not evolved in California D. simulans is consistent with the data of Hoffmann and Turelli [6] who found no heritable variation in CI-susceptibility among hundreds of isofemale lines collected from Melbourne, Australia in 1985. Jaenike and Dyer [23] present evidence that D. annabida has not evolved resistance to Wolbachia-induced male killing (MK), despite appreciable selection favoring suppressors for at least 15,000 – and possibly over 500,000 – generations. In contrast, transinfection experiments, introducing the MK Wolbachia that naturally infects D. reeves into its infected sister species D. subpiniwara, reveal that D. subpiniwara is polymorphic for dominant alleles conferring...
resistance to MK [24]. Even more striking is the evolution of resistance to MK over the past century in south Pacific and southeast Asian populations of the butterfly *Hypolimnas bolina* [25,26]. As in *D. subquinaria*, MK resistance in *H. bolina* is dominant [23].

How strong is the selection on *D. simulans* in CA to suppress CI? As shown in [4] and our new data, about 7% of the *D. simulans* females in CA are uninfected and they are likely to lose about half of their potential offspring to CI. Thus, a dominant autosomal suppressor of CI in *D. simulans* would experience a selective advantage on the order of 2%. The fixation probability of a new mutation with this phenotype would be about 4%. Once such an allele reached an appreciable frequency, say about 5%, its near-deterministic trajectory of fixation would take it from 5% to about 50% within 200 generations, the time scale of our field observations in California. These calculations support the arguments of Jaenike and Dyer [23] that dominant suppressors of *Wolbachia*-induced embryo death arise in some taxa extremely rarely or may have strongly deleterious pleiotropic effects that prevent their spread.

**Lab estimates: CI-induction by infected males.** Like Hoffmann and Turelli [6], we found maternally inherited variation in the intensity of CI induced by males from both our D08 and Y09 collections. The D157 line, which produced no uninfected sublines when initially collected, induced lower CI levels at day 7 than did Riv88. As shown by reciprocal intercrosses (Figure 4), the difference in CI intensity was attributed to *Wolbachia* or other maternally inherited factors. Reciprocal crosses also showed that maternal factors caused several Y09 lines to produce less intense CI than our older reference stock, Riv84 (Figure 5).

Changes in fecundity have also been associated with evolution of *wRi* rather than its host [8]. Contrary to the intuition that *Wolbachia* should evolve to increase CI [27,28], theoretical analyses indicate that *Wolbachia*-produced changes in CI should largely be pleiotropic by-products of changes in fecundity and/or transmission fidelity [11–13]. In principle, the generally lower intensity of CI produced by newly collected stocks may be attributable to either upward evolution of *H* in nature or decreases in *H* associated with selection and/or inbreeding in our laboratory stocks. This is currently being investigated.

**Maternal Transmission**

Our best estimate of maternal transmission from wild-caught females followed the protocol of the fourth maternal transmission experiment of Turelli and Hoffmann [4]. This involved remating wild-caught females to uninfected males and testing several F2 from each putatively uninfected F1 derived from a wild-caught infected female. Our goal was to estimate in nature *µ*, the fraction of uninfected ova produced by infected females. We first determined *v*, the fraction of uninfected F2 sublines produced by a wild-caught female. Some uninfected ova from these females may be lost due to CI. This initial experiment produced an estimate *v* = 0.015, with 95% BCα bootstrap confidence interval (0.003, 0.070), based on bootstrapping over the groups of isofemale sublines produced by individual wild-caught females. After mating these females to uninfected lab males, we obtained *µ* = 0.048 (0.016, 0.116) in our second experiment.

As noted in [4], if we assume that uninfected ova from infected females are as susceptible to CI as are the ova from uninfected females, then

\[
v = \frac{\mu(1 - psh)}{1 - \mu psh} \leq \mu,
\]

where *p* denotes the infection frequency among mating adults in nature, and *H*= 1−*s* is the relative hatch rate of incompatible crosses in nature. (The numerator in (1) is proportional to the number of uninfected surviving progeny from an infected female, and the denominator is proportional to the total number of surviving progeny for both infected and uninfected females.) Our point estimates from this study are *µ* ≈ 0.048, *p* = 0.932 and *s* = 0.615. Using (1), these produce an estimate of *v* = 0.021, which is very close to the observed value (0.015) and certainly within the estimated 95% confidence interval (even without accounting for the variation in our individual point estimates).

The new estimate *µ* ≈ 0.048 is essentially identical to the composite estimate *µ* = 0.042 produced in [4], indicating that the mean rate of maternal transmission has not changed substantially in nature. However, several aspects of our data and those in [4] merit comment. First, different experiments in [4] gave very different results. Their first and third maternal transmission experiments estimated *v* in Eq. (1) from females collected in April 1993 and November 1993 at Ivanhoe in central California. These females were not remated to uninfected males in the lab. In the April sample, of 44 infected wild-caught females that produced 8–10 F2 sublines, 23 produced both infected and uninfected F2 progeny (420 F2 sublines were assayed in total). In the November sample, all 49 infected wild-caught females produced only infected F2 (490 sublines were examined). Using a $\chi^2$ contingency test, these experiments gave significantly different results concerning the fraction of infected wild-caught females that show imperfect maternal transmission (**P** < 10−7). Given this extreme temporal variation from a single population, changes in *µ* over decades involving different populations will be essentially impossible to detect.

A striking aspect of our new data is that 4 of 65 wild-caught females accounted for most of the imperfect maternal transmission. Indeed, one female produced only one infected F2 subline out of ten. Our new estimate of *µ* is based on the experimental design used to produce the data in Figure 4 of [4]. In that experiment, 15 of 60 infected females produced both infected and uninfected F2 sublines, and one produced only one infected subline out of 11. Turelli and Hoffmann [4] conjectured that this female may have experienced rare paternal transmission of *Wolbachia*, as described in Hoffmann and Turelli [6]. However, it is equally plausible that females in nature harbor dramatically different levels of *Wolbachia*, which affect maternal transmission rates of *Wolbachia* and susceptibility to CI (see Figure 4 of Turelli and Hoffmann [4]). Unckless et al. [29] documented 20,000-fold variation in *Wolbachia* density among wild-caught *D. innubila*. This variation is associated with both penetrance of the MK phenotype and fidelity of maternal transmission. Similar data are needed for the *D. simulans*-wRi system [4]. In light of plausible extreme variation in *Wolbachia* titers among wild-caught flies, one might question the validity of our PCR assay for infection status, upon which our parameter estimates critically depend. Indirect support for the reliability of our assay comes from a new study by Casper-Lindley et al. [30] that reports a recent survey of *Wolbachia* infection frequencies in central California based on high-resolution microscopic examination of fluorescent-stained ovarioles. In a pooled sample of 106 females collected over three years, they report an infection frequency of 93.4%, consistent with our PCR-based estimate in Table 1.

**Fecundity**

Our point estimate of *p*†, the relative fecundity of infected females, is 0.98. However, even with over 500 wild-caught females, the fact that only about 7% are uninfected limits the accuracy of this estimate, as reflected by the wide 95% BCα bootstrap confidence interval (0.85, 1.15). Weeks et al. [8] used data from recently collected (2003–2005) versus old (1988) laboratory stocks to argue...
that aRi had evolved over about 15 years to increase female fecundity. They found that aRi seemed to increase fecundity in the laboratory by about 10%, but argued for much smaller effects in nature, on the order of a few percent. Our new field data are consistent this prediction. Differences in fecundity effects for laboratory versus field populations are analogous to the differences we observe in maternal transmission. In our long-held laboratory stocks, we observe perfect maternal transmission of aRi, whereas we have repeatedly found infected females in nature who produce infected and uninfected progeny [4,5]. To test the robustness of the Weeks et al. [8] fecundity finding and our new results concerning CI levels, we are now screening large samples of recently collected isofemale lines for variation in both CI and aRi-induced fecundity effects under laboratory conditions.

**Infection Frequency**

Sampling over a thousand individuals in 2008, we estimated the aRi infection frequency near Davis, California to be 0.932 (0.915, 0.946). Essentially the same value was obtained by Weeks et al. [8] sampling four populations from southern to northern California in 2003 and by Turelli and Hoffmann [4] from similar sites in late 1993. Is this apparent equilibrium frequency, denoted \( \hat{p} \), consistent with the simple three-parameter model proposed in Hoffmann et al. [5]? Using our point estimates from the field, \( \mu = 0.048 \), \( s_f = 0.62 \) and \( s_h = 0.0 \), we obtain \( \hat{p} = 0.966 \), which is significantly higher than the observed value. However, relatively slight changes in the parameter estimates, consistent with their variation over assays, can produce the observed value. For instance, if we set \( \mu = 0.05 \) and \( s_f = 0.0 \), we obtain \( \hat{p} = 0.932 \) with \( s_h = 0.47 \), which is consistent with many of our field estimates, including one of the two reported here. Conversely, if we set \( s_f = 0.6 \) and \( s_h = 0.0 \), we obtain \( \hat{p} = 0.932 \) with \( \mu = 0.079 \), which is statistically consistent with our latest estimate, \( \mu = 0.048 \) (0.016, 0.116). Holding \( s_f = 0.6 \) and introducing a 5% fecundity cost (\( s_h = 0.05 \)) reduces the required imperfect transmission rate to \( \mu = 0.053 \). Hence, the simple model with field-estimated parameters is consistent with the observed infection frequencies. However, this test is not particularly stringent. As discussed by Weeks et al. [8] and emphasized by Jaenike [31], the surface of equilibrium infection frequencies as a function of \( \mu \), \( s_f \) and \( s_h \) is relatively flat near our field-estimated values.

Despite the apparent quantitative adequacy of our simple model, our data demonstrate that it provides an incomplete description of aRi dynamics in nature. The model ignores age structure, even though the intensity of CI is known to be strongly age dependent [2,4]. However, given that there are no appreciable effects of aRi on viability, fecundity or development time, an age-structured model with many more parameters is unlikely to help us understand our infection frequency data [32]. Our model also uses a binary description, individuals are either infected or uninfected; whereas our new and old data indicate a continuum of infection states that correspond to different levels of maternal transmission and at least partial incompatibility of some infected females with some aRi males (see Figure 4 of [4]). Only one field test has directly addressed this. Turelli and Hoffmann [4] collected virgin females as they emerged from fruit collected and held in nature, then mated them to wild-caught males from the same population. They found no statistically significant difference in the hatch rate of eggs from aRi-infected females mated with infected or uninfected males. However, the sample sizes were relatively small.

A more elaborate model, with an additional parameter describing partial CI between infected males and infected ova, will clearly lower the predicted equilibrium. In addition to the three parameters \( \mu \), \( s_f \) and \( s_h \), let \( H_1 = 1 - s_h \) denote the average relative hatch rate when infected ova are fertilized by sperm from infected males. If we assume that \( \mu = 0.05 \), \( s_f = 0.0 \), \( s_f = 0.5 \) and \( s_h = 0 \), we obtain \( \hat{p} = 0.941 \) from the standard equilibrium formula (Eq. 3 of [4]). Introducing \( s_h = 0.05 \), i.e., only one-tenth the relative embryo loss attributable to CI between uninfected ova and sperm from infected males, lowers the stable equilibrium to \( \hat{p} = 0.933 \).

**Implications and Future Directions**

Our laboratory analyses of CI intensity are strongly suggestive of aRi evolution in California over the past 20 years, consistent with the data of Weeks et al. [8]. Future analyses will focus on characterizing many aRi-infected lines for Wolbachia load in nature, maternal transmission, fecundity and CI induction, relative to the reference Riv88 and Riv04 stocks. We also hope to associate CI and fecundity phenotypes with aRi genomic variation. The published aRi genome sequence [33] greatly facilitates the assembly of other aRi genomes.

This study produced four primary findings. First, we found relative stability in the parameter values governing the dynamics and equilibria of the Californian D. simulans-aRi system after about 15 years and probably around 200 generations. Second, consistent with the results from Hoffmann and Turelli [6], we found no variation among uninfected isofemale lines in their susceptibility to aRi-induced CI. Third, like [6], we found abundant maternally inherited variation in CI induction, presumably associated with variation among the aRi segregating in nature. Fourth, we have tentative evidence that aRi has evolved to produce lower levels of CI. One 2006 isofemale line (D157) has repeatedly shown lower CI intensity than Riv06. Five additional lines from our 2009 collection showed lower CI intensity than Riv04, which in all cases could be attributed to a cytoplasmically inherited factor. Further screening is required to rule out inbreeding artifacts associated with our laboratory stocks, to document the repeatability of the proposed evolutionary changes, and to document the spatial and temporal dynamics of the aRi variants we have described.

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**Author Contributions**

Conceived and designed the experiments: AAH MT LBC. Performed the experiments: LBC JRL MT. Analyzed the data: LBC AAH MT JRL. Contributed reagents/materials/analysis tools: MT AAH. Wrote the paper: MT LBC AAH.

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