**Wolbachia-induced cytoplasmic incompatibility as a means for insect pest population control**

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Biological control is the purposeful introduction of parasites, predators, and pathogens to reduce or suppress pest populations. **Wolbachia** are endosymbiotic bacteria that have recently attracted attention for their potential as new biocontrol agents. **Wolbachia** manipulate host reproduction by using several strategies, one of which is cytoplasmic incompatibility (CI) [Stouthamer, R., Breeuwer, J. A. J. & Hurst, G. D. D. (1999) *Annu. Rev. Microbiol.* 53, 71–102]. We established **Wolbachia**-infected lines of the medfly *Rhagoletis cerasi* using the infected cherry fruit fly *R. cerasi* as donor. **Wolbachia** induced complete CI in the novel host. Laboratory cage populations were completely suppressed by single releases of infected males, suggesting that **Wolbachia**-induced CI could be used as a novel environmentally friendly tool for the control of medfly populations. The results also encourage the introduction of **Wolbachia** into pest and vector species of economic and hygienic relevance to suppress or modify natural populations.

Cytoplasmic incompatibility (CI) is the most widespread and, perhaps, the most prominent feature that **Wolbachia** endosymbionts impose on their hosts (1, 2). CI results in embryonic mortality (EM) in matings between insects of the same species with different **Wolbachia** infection status (3, 4). It can be either unidirectional or bidirectional. Unidirectional CI is typically expressed when an infected male mates with an uninfected female. The reciprocal mating is fully compatible, as are matings between infected individuals. Bidirectional CI usually occurs in matings between infected individuals harboring different strains of **Wolbachia**. Although the mechanism of CI has not yet been elucidated on the molecular level, several lines of evidence suggest that **Wolbachia** somehow modifies the paternal chromosomes during spermatogenesis (mature sperm does not contain the bacteria). This modification influences their behavior during the first mitotic divisions and results in loss of mitotic synchrony (5, 6). Even before the etiological connection between **Wolbachia** and CI was revealed in mosquitoes (7), attempts were made to exploit CI as a method to suppress natural populations of arthropod pests in a way analogous to the sterile insect technique (S.I.T.) (8). These early attempts involved the mass production and release of incompatible male insects to control wild populations of disease vectors such as the mosquito *Culex pipiens* (9) and of agricultural pests such as the European cherry fruit fly, *Rhagoletis cerasi* (10) and, at a smaller scale, the almond moth *Cadra cautella* (11).

The Mediterranean fruit fly (medfly) *Ceratitis capitata* is a worldwide pest that infests >250 fruit varieties of economic importance (12). Extensive screenings of both laboratory and natural populations of medfly have shown that this insect pest is not infected with **Wolbachia** (13). However, at the time of this writing, ongoing experimental work suggests that some Brazilian natural populations of medfly may be infected with **Wolbachia** (D. Selivon, personal communication). To determine whether the medfly can support **Wolbachia** infections and express CI, we used conventional embryonic cytoplasmic injections (14–17) for transfer of natural bacterial symbionts from a related species, *R. cerasi* (Diptera, Tephritidae) (18), to an uninfected laboratory strain of medfly *C. capitata* (Diptera, Tephritidae), the Benakcion strain.

Previous studies have demonstrated high levels of incompatibility between natural populations of *R. cerasi* (10, 19), the basis of which was recently shown to be **Wolbachia** (18). Populations of *R. cerasi* are either infected by a single **Wolbachia** variant, wCer1, or coinfected by two variants, wCer1 and wCer2. Incompatibility occurs between males from doubly infected populations and females from singly infected populations, suggesting the wCer2 infection as the cause of CI (18). Additionally, transfer of wCer2 in *Drosophila simulans* also resulted to the induction of CI (17). An additional, yet uncharacterized, **Wolbachia** strain (wCer3) has been recently found in Sicilian populations of *R. cerasi* (M.R. and C. Stauffer, unpublished results).

The aim of our study was to establish **Wolbachia**-infected lines of the medfly by using the infected cherry fruit fly as donor. Previous attempts to transfer **Wolbachia** strains from species such as *Drosophila melanogaster, D. simulans* (infected with *wRi*), and *Cadra cautella* into medfly failed (K.B., unpublished data), due to either an unsupportive host background for **Wolbachia**, or the lack of adaptation of the transferred **Wolbachia** strains to the new host. A transfer of **Wolbachia** strains from the more closely related host species *R. cerasi* would have a greater potential. Furthermore, we were interested in whether transferred **Wolbachia** could induce CI in the novel host, also in an attempt to use the mechanism of **Wolbachia**-induced CI as a novel environmentally friendly tool for the control of medfly populations. The results presented here are very supportive and encourage the introduction of **Wolbachia** into pest and vector species of economic and health relevance to suppress or modify natural populations.

**Materials and Methods**

**Insects.** *Ceratitis capitata*. Benakcion is an uninfected laboratory strain. A71 is an uninfected white eye mutant laboratory strain. Both strains are kept in mass in population cages at 24°C on standard medfly diet (13).

*Rhagoletis cerasi*. Two natural populations of cherry fruit flies were collected from Austria and from Sicily (Italy). The first was doubly infected with wCer1 and wCer2 and the second infected with wCer1 (18) and wCer4 (this study). The Sicilian population had previously been found to carry additional **Wolbachia** strains (M.R. and C. Stauffer, unpublished results).

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Embryonic Cytoplasm Transfer. Wolbachia was transferred from naturally infected cherry fruit fly populations into the Benakeion laboratory strain of C. capitata. Microinjections were carried out by using a microcapillary needle (Bohringer Femtotips). Medfly embryos were collected for 60 min, dechorionated, and slightly desiccated. The cytoplasm was taken from the posterior region of donor mature oocytes and injected at the posterior pole of the recipient preblastoderm embryos (16, 17).

Establishment of the Infected C. capitata Lines. Females deriving from injected embryos represent the generation G0 postinjection. G0 females were each crossed with two Benakeion males. The progeny of transfected G0 females were kept in mass culture, establishing a line.

DNA Extraction, PCR, and Sequencing. Total DNA was extracted from single individuals following the STE boiling method (20). Wolbachia was detected by PCR using the 16S rDNA Wolbachia-specific primers 99F and 994R (20). At least 60 individuals from the WolMed 88.6 line and 40 individuals from the WolMed SI0.3 line were screened for infections every generation. Distinction between different Wolbachia variants was done with strain-specific wsp primers (18). For sequencing, PCR products were amplified by using the wsp primers 81F and 691R (21) and were subjected to direct sequencing. At least three individuals from each established line were sequenced.

CI Assays and Statistical Analysis. CI levels were measured in two different ways, in single pair matings and in cage populations. All of the crosses were performed at 24°C. For single-pair matings, 2-day-old virgin females were individually mated with 1-day-old virgin males. The egg-laying plates were removed daily, and all eggs were scored for a period of 6–8 days. Cage populations consisted of 100 2-day-old virgin females mated with 100 1-day-old virgin males. A random sample of 500 eggs was kept every day. Hatching rates were scored 72 h after egg collection. EM was determined as the percentage of unhatched eggs. The standard error (SD) for EM, was determined according to Ott (22). EM in the progeny of infected males and uninfected females was complete, which let us assume that CI is complete.

Suppression of C. capitata Populations. The population suppression experiments were performed in six population cages at 24°C. Each cage contained equal numbers of 2-day-old virgin uninfected females and 1-day-old virgin uninfected males (1:1), plus 1-day-old virgin transfected males at different ratios 1:1:0, 1:1:1, 1:1:10, 1:1:20, 1:1:30, and 1:1:50. The first five cages contained ~300 flies and the last one (ratio 1:1:50) ~520 flies. Egg-laying plates were removed every day for a period of 6–8 days. In the first two cages, a random sample of 500 eggs was kept daily, whereas, in the rest of them, all laid eggs were collected. Hatching rates were scored 72 h after egg collection. Survival was determined as percentage of hatched eggs.

Detection of Wolbachia by Immunofluorescence. Embryos were collected and dechorionated in 50% commercial bleach for 5 min. After a quick rinse with washing buffer (0.7% NaCl, 0.3% Triton X-100), they were transferred to 1:1 heptane-methanol solution and shaken vigorously for a couple of minutes. They were then briefly washed three times with methanol and three times with TBST (50 mM Tris-HCl/150 mM NaCl/0.1% Tween 20/0.05% NaN₃, pH 7.5), 15 min each; were then blocked in 1% BSA in TBST; and incubated with the Wolbachia surface protein (WSP) antibody with a 1:500 dilution overnight at 4°C. After three washes with TBST, the eggs were incubated for 1 h at room temperature with a 1:500 dilution of Alexa Fluor 488 goat anti-rabbit IgG-labeled antibody (Molecular Probes) and 2 mg/ml Rhose A (Sigma) in TBST. After several washes in TBST, eggs were stained with 5 μg/ml propidium iodide (Molecular Probes) for 20 min, rinsed, and mounted with ProLong Antifade kit (Molecular Probes).

Ovaries from 2- to 3-day-old females and testes from 1-day-old males were removed in TBST and further dissected on glass slides. The ovaries were flattened under a cover glass and stored in liquid nitrogen. Cover glasses were removed by using a razor blade, and the slides were placed in ice-cold ethanol for 3 min and fixed in 4% paraformaldehyde for 12 min. Slides were rehydrated in TBST, blocked, and incubated with antibodies and propidium iodide as described (23).

Optical sections were taken by using a confocal laser-scanning microscope (TCS-NT, Leica, Deerfield, IL), and they were projected onto single images. Images were further processed by using photoshop 6.0 (Adobe Systems, San Jose, CA).

Results and Discussion

Establishment of Wolbachia-Infected Medfly Lines. To determine whether the medfly can support Wolbachia infections and express CI, we used conventional embryonic cytoplasmic injections (14–17) to transfer natural bacterial symbionts from a related species, R. cerasi (Diptera, Tephritidae) (18), to an uninfected laboratory strain of medfly C. capitata (Diptera, Tephritidae), the Benakeion strain. Infected lines of R. cerasi from Sicily (Italy) and Austria were used as donors of Wolbachia-infected embryonic cytoplasm. Eighty-eight G0 isofemale lines were established and were monitored at each generation for the presence of Wolbachia by using a specific PCR assay (20). After three generations of monitoring, 2 of initially 11 positive transfected isofemale lines remained positive for the presence of Wolbachia, namely WolMed 88.6 and WolMed SI0.3. Analysis of PCR-amplified sequences of bacterial wsp gene showed that each of the 2 lines was infected by a different Wolbachia strain. Line WolMed 88.6 was found to be infected with the wcer2 strain, which originated from the R. cerasi Austrian population (accession no. AF418557), whereas line WolMed SI0.3 was found to be infected with wcer4. This Wolbachia strain was previously undetected in its original host, but originated from an island population (Sicily) that has characteristic infection types (18), including previously nondescribed Wolbachia strains (M.R. and C. Stauffer, unpublished results). The wcer4 strain was found to be 100% identical with the wIr-A1 strain (accession no. AF217714) based on a partial wsp gene sequence. At the time of this writing, 26 generations (~23 months) postinjection, both transfected lines are stably infected, with infection rates of 100%.

Confocal microscopy analysis was performed in embryos, ovaries, and testes of both transfected medfly lines by using an anti-WSP (Wolbachia surface protein) antiserum (24) (Fig. 1). Both lines present high infection levels and distribution patterns as is characterized Drosophila–Wolbachia associations (23, 25–27). In early medfly embryos, Wolbachia show a clear association with the host cytoskeleton interacting with the mitotic spindle apparatus (Fig. 1A). The bacteria then comigrate with the nuclei toward the periphery of the embryo and at the same time get incorporated into pole cells (Fig. 1B). It is notable that, in medfly preblastoderm embryos, the bacteria are concentrated at the basal side of the nuclei, in contrast with most Drosophila–Wolbachia symbioses where the bacterium is associated with the apical side of precellular nuclei (23). At later stages, Wolbachia are evenly distributed throughout the developing embryo, and this pattern remains constant until late gastrulation (Fig. 1C). In the transfected medfly lines, Wolbachia were present in the ovaries, mostly within the egg chamber surrounding the nuclei of the nurse cells, probably infecting the oocyte at later stages of the oogenesis through the cytoplasmatic dumping process (Fig. 1D). Wolbachia density is quite high throughout spermatogenesis, the bacteria being present at high levels within the sperm cysts until their removal to the waste bags during the individualization process of spermatooza (Fig. 1E and F).

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Fig. 1. Presence of *Wolbachia* in transinfected *C. capitata* embryos, ovaries, and testes. (A) Transinfected medfly embryo undergoing synchronous mitotic divisions showing *Wolbachia* localization at the mitotic spindles. (B) The posterior part of a transinfected medfly embryo, where pole cells (the precursors of gonads) are being formed, showing incorporation of *Wolbachia* in the pole plasm. (C) Uniform distribution of *Wolbachia* bacteria in a postgastrulation transinfected embryo. (D) Distribution of *Wolbachia* during oogenesis, when oocytes start to form. The bacteria are mostly concentrated in nurse cells, presumably infecting the oocyte at later stages of oogenesis. (E) Large numbers of *Wolbachia* are present in adult testes of transinfected medfly. (F) A sperm cyst is shown with the distal end toward the left and nuclei to the right. After elongation is complete, most cytoplasmic components, including *Wolbachia*, are stripped away from the sperm during individualization and sequestered in the waste bag (shown as a “green ball” in the left). Bacteria are visualized green-yellow and nuclei red. [Scale bars: 40 μm (A, B, and D–F) and 100 μm (G).]

**Wolbachia-Infected Medfly *C. capitata* Lines and Expression of CI.** To determine whether the two transinfected lines were capable of inducing CI, test crosses were performed between each transinfected line and the parental naturally uninfected Benakeion strain. Crossings in experiments in cages with 100 uninfected females and equal numbers of infected males from each transinfected line (seventh generation post injection) resulted in 100% egg mortality; the reciprocal crosses resulted in <32% egg mortality under identical experimental conditions (Table 1, No tetracycline). Similar results were obtained in cage experiments between uninfected females from the medfly strain A71 (a white eye mutant *C. capitata* line) and infected males from each transinfected line (sixth generation postinjection, data not shown).

Persistence of *Wolbachia*-induced CI was also tested by single-pair matings. Twenty-six single-pair crosses between WolMed 88.6 (wCer2) transinfected males (fifth generation postinjection) and uninfected females resulted in 100% egg mortality (no hatched eggs out of 2164). Similarly, 17 single-pair crosses between WolMed S10.3 (wCer4) transinfected males (fifth generation postinjection) and uninfected females resulted again in 100% egg mortality (no hatched eggs out of 1,325). The same experiments were repeated again in the tenth generation postinjection, providing the same results, 100% egg mortality (data not shown). It is notable that complete CI has been observed only in very few *Wolbachia*-infected species such as *C. pipiens* (9). To our knowledge, there have been no previous reports that a novel transinfected host species presents high stability of the infection and, at the same time, expresses 100% CI.

The two transinfected medfly lines WolMed 88.6 (wCer2) and WolMed S10.3 (wCer4), each infected with a different *Wolbachia* strain, were 100% bidirectionally incompatible in appropriate genetic cross experiments performed in cages (Table 1).

Control crosses between infected males and females resulted in high EM, ~65%, in both transinfected lines (Table 1, No tetracycline). This effect is caused by *Wolbachia*, as shown by eliminating the bacteria by treatment with antibiotics: Crosses between individuals of the tetracycline cured lines, three and five generations after curing, resulted in normal rates of EM (Table 1, Tetracycline). One explanation for these high EM could be inefficient transmission of both wCer2 and wCer4 in medfly. However, at least 60 individuals from the WolMed 88.6 line and 40 individuals from the WolMed S10.3 line were tested by PCR in every generation for 26 generations (~23 months), and all individuals have been found positive for *Wolbachia*. A more likely cause for the high mortalities observed in the crosses between infected individuals could be additional fertility effects of wCer2 and wCer4 on medfly females, effects other than CI. Alternatively, it is possible that infected medfly females can only

Table 1. *Wolbachia*-induced cytoplasmic incompatibility in two transinfected lines of the medfly *C. capitata*

<table>
<thead>
<tr>
<th>Cross (females × males)</th>
<th>Embryos scored</th>
<th>Embryonic mortality, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>No tetracycline*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Uninfected × WolMed 88.6 (wCer2)</td>
<td>3,000</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>Uninfected × WolMed S10.3 (wCer4)</td>
<td>3,000</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>WolMed 88.6 (wCer2) × Uninfected</td>
<td>3,000</td>
<td>16.73 ± 0.68</td>
</tr>
<tr>
<td>WolMed S10.3 (wCer4) × Uninfected</td>
<td>3,000</td>
<td>32.03 ± 0.85</td>
</tr>
<tr>
<td>WolMed S10.3 (wCer4) × WolMed 88.6 (wCer2)</td>
<td>3,000</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>WolMed 88.6 (wCer2) × WolMed S10.3 (wCer4)</td>
<td>3,000</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>WolMed 88.6 (wCer2) × WolMed 88.6 (wCer2)</td>
<td>3,000</td>
<td>64.77 ± 0.87</td>
</tr>
<tr>
<td>WolMed S10.3 (wCer4) × WolMed S10.3 (wCer4)</td>
<td>3,000</td>
<td>67.25 ± 0.87</td>
</tr>
<tr>
<td>Uninfected × Uninfected</td>
<td>3,000</td>
<td>12.17 ± 0.60</td>
</tr>
<tr>
<td>Tetracycline†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WolMed S10.3 tet × WolMed S10.3 tet†</td>
<td>1,890</td>
<td>23.44 ± 0.97</td>
</tr>
<tr>
<td>WolMed S10.3 tet × WolMed S10.3 tet†</td>
<td>3,000</td>
<td>11.80 ± 0.59</td>
</tr>
<tr>
<td>WolMed 88.6 tet × WolMed 88.6 tet†</td>
<td>2,283</td>
<td>25.10 ± 0.91</td>
</tr>
</tbody>
</table>

*Wolbachia*-induced cytoplasmic incompatibility is expressed as percentage of unhatched eggs ± SE. Egg-laying plates were removed daily for a period of 6 days. Hatching rates were scored 72 h after egg collection.

*Test crosses between each transinfected line and the parental naturally uninfected Benakeion strain as well as between the two transinfected lines. Crosses between 100 females (2–3 days old) and equal numbers of males (1 day old) were performed in cages.

†Test crosses between tetracycline-treated individuals of the line WolMed S10.3. Crosses between 30 females and equal numbers of males were performed in cages, three generations (†) and five generations (‡) after the tetracycline treatment.
partially rescue the modification induced by the bacteria in infected medfly males. The last two hypotheses will require further investigation.

**Wolbachia-Induced CI as a Potential Tool for Medfly Population Suppression.** To determine whether cytoplasmic incompatibility expressed by the *Wolbachia*-infected medfly lines could be used for population suppression, we set up cage populations containing different ratios of uninfected females to uninfected males to transinfected males (1:1.0, 1:1.1, 1:1.10, 1:1.20, 1:1.30, 1:1.50). These experiments were performed by using transinfected WolMed 88.6 (wCM1) males from generations 8 to 11 postinfection. As shown in Fig. 2, the laboratory cage medfly populations were suppressed by these single “releases” of incompatible males in a ratio-dependent manner. Under these conditions, population suppression reached levels higher than 99% in releases of incompatible males at a ratio 1:1.50. Similar results were obtained in cage experiments by using as target population the medfly white eye mutant strain A71 (data not shown). These

data clearly demonstrate that *Wolbachia*-induced cytoplasmic incompatibility could be used as a tool for the population control of this major agricultural pest.

In recent years, there has been increasing interest in the biology of *Wolbachia* and in its application as an agent for control or modification of insect populations. It has been proposed that *Wolbachia* might be used (i) as a vector for the expression of genes of interest, (ii) as a tool to drive desirable genotypes into arthropod populations, and (iii) to directly suppress arthropod populations (28–31). The recent publication of the genome sequence of a CI-inducing strain (32) should accelerate progress in elucidating, at the molecular level, the interactions between *Wolbachia* and its hosts that lead to CI. Our present study clearly shows that *Wolbachia* endosymbionts can be experimentally transferred over genus barriers into a novel host, forming associations that express complete CI. The population cage experiments strongly suggest that *Wolbachia*-induced CI (unidirectional and, importantly, bidirectional) could be used as a means for control of natural medfly populations. The observed increased lethality in crosses between infected males and females will have some negative impact in mass rearing; however, *Wolbachia* transinfection experiments into other host insects have shown that deleterious fitness effects can attenuate rapidly in consecutive selection processes (33). Alternatively, the system may be improved by testing additional bacterial strains that may not be causing this lethality effect. For effective *Wolbachia*-based population suppression, an efficient genetic sexing system producing males only is necessary. Such systems are available in medfly (34). However, because accidental release of even one female can theoretically result in the replacement of the target population by a second, the sexing system has to be 100% effective. Altogether, our results should encourage efforts to transfer *Wolbachia* into other insect pests or disease vectors for suppression or modification of natural populations.

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