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Sexual functionality of *Leptopilina clavipes* (Hymenoptera: Figitidae) after reversing *Wolbachia*-induced parthenogenesis

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**Keywords:**
antibiotic curing; arrenotoky; *Leptopilina clavipes*; parthenogenesis; sexual function decay; thelytoky; *Wolbachia*.

**Abstract**
Females infected with parthenogenesis-inducing *Wolbachia* bacteria can be cured from their infection by antibiotic treatment, resulting in male production. In most cases, however, these males are either sexually not fully functional, or infected females have lost the ability to reproduce sexually. We studied the decay of sexual function in males and females of the parasitoid *Leptopilina clavipes*. In western Europe, infected and uninfected populations occur allopatrically, allowing for an investigation of both male and female sexual function. This was made by comparing females and males induced from different parthenogenetic populations with those from naturally occurring uninfected populations. Our results indicate that although males show a decay of sexual function, they are still able to fertilize uninfected females. Infected females, however, do not fertilize their eggs after mating with males from uninfected populations. The absence of genomic incompatibilities suggests that these effects are due to the difference in mode of reproduction.

**Introduction**
*Wolbachia* bacteria are cytoplasmic endosymbionts (z-proteobacteria) that infect a wide range of arthropod and nematode hosts. They are maternally inherited and enhance their transmission by altering the reproductive system of their host in various ways, such as cytoplasmic incompatibility, male killing, leminization and parthenogenesis induction (PI) (Stouthamer et al., 1999). All these phenomena lead to an increase in infected females.

Parthenogenesis-inducing *Wolbachia* are mainly found in the arthropod group Hymenoptera (Huigens & Stouthamer, 2003), and sporadically in other groups, such as Coleoptera (Werren et al., 1995), Thysanoptera (Arakaki et al., 2001) and in mites of the genus *Bryobia* (Weeks & Breeuwer, 2001). PI *Wolbachia* are restricted to hosts with haplodiploid modes of reproduction (Huigens & Stouthamer, 2003) in which males are haploid and females are diploid. The most common mode of haplodiploid reproduction is arrenotoky, where males develop from unfertilized eggs and females from fertilized eggs (White, 1973; Luck et al., 1993). Females infected by PI *Wolbachia* produce all-female offspring (thelytoky) through gamete duplication (Stouthamer & Kazmer, 1994; Gottlieb et al., 2002; Pannebakker et al., 2004b).

Parthenogenesis-inducing *Wolbachia*-infected females can be cured from their infection by high temperature or antibiotic treatment (Stouthamer et al., 1990a), which results in male production. In most cases however, either these males are sexually not fully functional (e.g. Zchori-Fein et al., 1992, 1995; Gottlieb & Zchori-Fein, 2001), or the cured females have lost the ability to reproduce sexually (Pijs et al., 1996; Arakaki et al., 2000). Only in several *Trichogramma* species did the removal of PI *Wolbachia* results in the production of arrenotokously reproducing lines (Stouthamer et al., 1990a,b).
In parthenogenetic populations, genes involved in sexual reproduction are not actively maintained by selection, and random mutations in those genes are thus not removed by selection (Muller, 1949; Carson et al., 1982). Mutations can accumulate in these genes or, by means of antagonistic pleiotropy, even be actively selected for if they improve the parthenogenetic performance of females (Pijls et al., 1996; Werren, 1998). The decay of sexual function in PI Wolbachia-infected systems has been studied extensively in species where infection has gone to fixation (e.g. Zchori-Fein et al., 1992, 1995; De Barro & Hart, 2001; Gottlieb & Zchori-Fein, 2001; Weeks & Breeuwer, 2001). However, the decay of sexual function in both males and females can only be studied in species where infection status is polymorphic.

Here we study the decay of sexual function in males and females of Leptopilina clavipes (Hartig) (Hymenoptera: Figitidae), a parasitoid wasp of fungi-breeding Drosophila larvae that occurs in western Europe (Nordlander, 1980). In north-western Europe, populations of L. clavipes are thelytokous (Driessen et al., 1990; Pannebakker et al., 2004c), which is Wolbachia induced (Werren et al., 1995; Schidlo et al., 2002). Recently, uninfected arrhenotokous individuals were found in northern Spain (Pannebakker et al., 2004c). In this study, we compare the sexual function of antibiotic curing-induced (ACI) males and Wolbachia-infected females from different thelytokous populations with that of material from natural arrhenotokous populations. We make the comparison at different levels: (i) male courtship behaviour (we include a description of L. clavipes courtship behaviour), (ii) male fertilization capacity, and (iii) sperm use by Wolbachia-infected females. In addition, we tested for the presence of genomic incompatibilities between the two modes of reproduction. Our results indicate that males show a decay of sexual function, but are still able to fertilize arrhenotokous females. Infected females however, do not use the sperm they receive after mating with arrhenotokous males. Genomic incompatibilities were absent between the two modes of reproduction and hence do not provide an explanation for our results.

Material and methods

Insect cultures

Leptopilina clavipes stocks

Twelve L. clavipes lines were used in the different experiments. The infected thelytokous lines BBH-NL00, DB23/9-NL99, DBK-NL00, GBW-NL00, HOD-NL00, KBH-NL00, VOSB-NL00 originated from the Netherlands, NEUVIC-F01 and RENNES-F01 originated from France. The uninfected arrhenotokous lines CCAP-E00, DC-E00 and Moll1-E00 originated from Spain. Collection details can be found in Pannebakker et al., 2004c. Cultures were maintained in the laboratory on Drosophila phalerata larvae at 20 °C, L : D = 16 : 8 and 65% relative humidity (RH). Drosophila phalerata were reared on a medium containing mushroom (Agaricus bisporus), water, dry yeast (Saccharomyces cerevisiae Hansen) and agar. The fungicides nipagin and propionic acid were added to prevent moulding of the medium.

Curing experiments and crosses were carried out using D. subobscura as a host. This species was reared on a patch of live bakers yeast (S. cerevisiae) suspension on a medium of water, dry yeast (S. cerevisiae) and agar at 20 °C, L : D = 16 : 8 and 65% RH.

Antibiotic treatment

To induce male offspring from the thelytokous line, infected females were cured from their Wolbachia infection using antibiotics applied in the honey (0.5% rifampicin) and in the host medium (0.2% rifampicin) as described by (Schidlo et al., 2002). Curing was done using D. subobscura as a host on a live bakers yeast patch on agar at 25 °C, L : D = 16 : 8 and 65% RH.

Crossing experiments

Courtship behaviour

The courtship behaviour of ACI males from eight different thelytokous lines (BBH-NL00, DB17/9-NL99, DBK-NL00, GBW-NL00, HOD-NL00, KBH-NL00, NEUVIC-F01 and VOSB-NL00) was compared with that of arrhenotokous males (DC-E00). One virgin female (DC-E00) was placed in a glass mating arena (9 mm high × 32 mm in diameter) closed off with a thin glass plate. A single virgin male was introduced and the couple was observed under a dissecting microscope at 10× magnification. If no copulation attempt occurred within the first 15 min after the introduction of the male, a ‘no copulation event’ was scored. If copulation began within 15 min, the couple was observed until the mating stopped. Successful insemination following copulation was confirmed by allowing each mated female to oviposit on a patch with approximately 110 D. subobscura larvae for 24 h, and scoring the sex of the offspring. Age of the mating pairs was not strictly controlled for; mean age ± SD (days) males: 3.05 ± 3.46; females: 2.74 ± 3.33.

Observations were made at an ambient temperature of 20 °C and 65% RH, from mid-July to late September 2001. The male courtship behaviour was recorded using specialized computer software (Observer 3.0; Noldus Information Technology, Wageningen, The Netherlands, 1993). Descriptive statistics were calculated on the number of, and duration of the behavioural elements in courtship.

Behavioural data were analysed by fitting generalized linear models (GLM) with the appropriate error structures and link functions (i.e. transition probability data: binomial error structure and a logit link; latencies and durations: Gaussian error structure and identity link; count data: Poisson distribution and a log link). Moderate levels of overdispersion in the binomial and Poisson GLMs were corrected for by rescaling the deviance by the
heterogeneity factor (HF), the ratio of the residual deviance to the degrees of freedom (McCullagh & Nelder, 1989). For HF > 3 we used standard ANOVA after the appropriate transformation. Data were analysed using R statistical software (Ihaka & Gentleman, 1996, version 1.71). Model selection was made by comparing the Aikake Information Criterion (AIC) of the initial model with the AIC after excluding the line effect from the model. An F-test was used to determine the significance of the line effect. If significant line effects were observed for a behavioural element, the effects of the individual lines were compared using F-tests.

Sex ratio
Separate series of matings were performed to compare the sex ratio produced by ACI males (DB23/9-NL99, DBK-NL00, GBW-NL00, KBH-NL00, NEUVIC-F01, RENNES-F01 and VOSB-NL00) with that of natural arrhenotokous males (CCAP-E00 and DC-E00) when mating to virgin sexual females from a reference line (DC-E00). Matings were observed as described in the courtship behaviour assay. The experiments were carried out from late April to May 2003.

When a successful mating was observed, as defined by genital contact for over 30 s, the female was allowed to oviposit on a patch of approximately 130 D. subobscura larvae for 48 h. After the first oviposition period, the female was transferred to a new patch of approximately 130 D. subobscura larvae for 48 h after which the female and male where stored at −80 °C for further genetic analysis. The larvae were incubated at 25 °C and 65% RH, L : D = 16 : 8 and allowed to pupate. After pupation, the pupae were washed out of the medium and transferred to a new vial. Numbers of emerging males and females were recorded for a period of 4 weeks, during which most parasitoids eclosed. The noneclosed pupae were then opened and the sex of the parasitoids was recorded.

Sex ratio data are usually binomially distributed and are, therefore, best analysed using GLMs (Wilson & Hardy, 2002). We fitted a generalized linear mixed model to the data, using SAS/STAT software (SAS Institute, Cary, NC, USA) and a specific macro (GLIMMIX) as described in Littell et al. (1996). The proportion of males in a clutch was modelled as a binomial random variable, and regression models were coupled to this probability using the logit link function. Matings involving the same combination of lines were treated as repeated measures. Model selection occurred by first fitting an elaborate model. This was then simplified on the basis of likelihood ratio tests excluding factors for which the confidence intervals (CIs) overlapped the value zero. The initial model included brood size, arrhenotokous/thelytokous, first/second clutch and age of female and male as covariates, female and male as random effects, and allowed for a correlation between sex ratios in first and second clutches of the same mating pair.

Sperm utilization in infected females
To test whether infected females use the sperm of uninfected males to produce hybrid offspring, infected females were crossed to uninfected males (KBH-NL00 × DC-E00, BBH-NL00 × Moll1-E00, HOD-NL00 × Moll1-E00). The parents and F1 offspring were frozen (~80 °C) for DNA analysis. Using amplified fragment length polymorphism (AFLP; Vos et al., 1995) markers, the F1 offspring were checked for the presence of paternal alleles. DNA isolation methods and AFLP procedures were as described in Pannebakker et al. (2004a), using the primer combination Mse-CA/Eco-ACA.

Recovery of parental alleles
To determine the existence of genomic incompatibilities between the two modes of reproduction, the recovery rate of alleles specific to each mode was examined in hybrid offspring. A cross was made between an ACI male from KBH-NL00 and a female from the arrhenotokous line DC-E00 under the conditions outlined above. Six female F1 offspring were frozen as virgins, resulting in 72 F2 recombinant males. The parents and F2 offspring were frozen (~80 °C) for DNA analysis. The F2 offspring were checked for the recovery of parental AFLP alleles that were generated as described in Pannebakker et al. (2004a) with the primer combinations Mse-CA/Eco-ACA and Mse-CA/Eco-AGG.

If genomic incompatibilities exist between the two modes of reproduction, they will be detectable by an unequal recovery of parental alleles in the F2 recombinant males (cf. Gadau et al., 1999). When incompatibilities are absent, equal proportions (1/2) of alleles from both reproductive modes are expected in the F2 males, due to meiosis in the hybrid F1 females. We tested for equal recovery of the parental alleles using a binomial test (Zar, 1996; Schork & Remington, 2000).

Results

Courtship and copulation experiments

Courtship behaviour
Courtship behaviour was described for matings between virgin males and females from the uninfected population DC-E00. Typical courtship behaviour is illustrated in Fig. 1, while the sequence used for recording of male courtship behaviour is illustrated in Fig. 2. The sequence of courtship behaviour in L. clavipes resembles that of other Eucoline parasitoids [i.e. L. heterotoma = Pseudewida bocheci (van den Assem, 1969) and Aganaspis pelleranoi (Ovurski & Aluja, 2002)].

Male courtship behaviour differentiation
The courtship behaviour of the ACI males and the males from the arrhenotokous population is summarized in
Table 1. The transition to wingfan that indicates detection of the resident virgin female shows significant differences between the lines (Table 1). This difference, however, is caused solely by the difference in transition to wingfan of the thelytokous line GBW-NL00 with the arrhenotokous and, all the other ACI males ($F_{8,102} = 3.47, P < 0.001$). No differences in further transitions towards copulation were found, although considerable variation exists in the proportion of copulations that result in successful fertilization.

The quantitative behavioural traits show high variability between the different lines. However, no consistent differences were found in male courtship behaviour between males from the arrhenotokous population and ACI males from the thelytokous populations.

**Sex ratio experiments**

The sex ratios (proportion of male offspring) produced by ACI males from the thelytokous lines are significantly higher ($F_{1,8.65} = 23.06, P < 0.05$, Tables 2 and 3) than those produced by the natural arrhenotokous males. No significant differences were found between arrhenotokous males from CCAP-E00 and DC-E00. Within the thelytokous males, however, the variance is higher than among the females (Tables 2 and 3). There is overall only a weakly significant effect of male line within the sex.

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**Fig. 1** Schematic representation of male courtship behaviour in uninfected *Leptopilina clavipes*. (a) Male introduced to mating arena. (b) Following detection of the female, the male starts vigorous wing fanning until genital contact is made. (c) While wing fanning, the male follows the female and makes antennal contact with the female’s dorsal gaster or thorax upon which receptive females remain motionless. (d) Male mounts female until his head is over that of the female after which the female raises her antennae in an upward position. (e) Male starts antennation, consisting of male–female antennal contact (antennal sweeps) and wing fanning bouts. Antennal sweeps are characterized by forward extension of the antennae, which are both then raised and lowered simultaneously in small circular movements. (f) After a number of antennal sweeps, a receptive female moves her antennae from a straight upward to a downward position after which she opens up her abdomen. (g) Upon this signal, the male stops the antennal sweeps and wing fanning and begins copulation. During copulation, the female remains arrested, while the male slowly moves his antennae up and down. (h) Copulation is usually terminated by the female by pushing the male with her hind legs and removing him from her back.

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**Fig. 2** Sequence of male courtship behaviour in *Leptopilina clavipes* used in the behavioural recordings.
Transitions and quantitative data on male *Leptopilina clavipes* courtship behaviour from arrhenotokous (DC-E00) and thelytokous lines (BBH-NL00, DB17/9-NL99, DBK-NL00, GBW-NL00, HOD-NL00, KBH-NL00, NEU-F01, VOS-NL00).

<table>
<thead>
<tr>
<th>Line</th>
<th>N</th>
<th>Antennal sweeps</th>
<th>Copulation attempts</th>
<th>Copulation latency</th>
<th>Fertilization</th>
</tr>
</thead>
<tbody>
<tr>
<td>DC-E00</td>
<td>24</td>
<td>0.96^a</td>
<td>0.94</td>
<td>0.93</td>
<td>0.64</td>
</tr>
<tr>
<td>BBH-NL00</td>
<td>13</td>
<td>0.85^a</td>
<td>0.90</td>
<td>0.89</td>
<td>0.25</td>
</tr>
<tr>
<td>DB17/9-NL99</td>
<td>13</td>
<td>0.85^a</td>
<td>1.00</td>
<td>0.75</td>
<td>0.63</td>
</tr>
<tr>
<td>DBK-NL00</td>
<td>13</td>
<td>0.85^a</td>
<td>1.00</td>
<td>0.50</td>
<td>1.00</td>
</tr>
<tr>
<td>GBW-NL00</td>
<td>11</td>
<td>0.27^b</td>
<td>1.00</td>
<td>0.00</td>
<td>-</td>
</tr>
<tr>
<td>HOD-NL00</td>
<td>11</td>
<td>0.82^a</td>
<td>1.00</td>
<td>0.63</td>
<td>0.40</td>
</tr>
<tr>
<td>KBH-NL00</td>
<td>13</td>
<td>0.85^a</td>
<td>0.90</td>
<td>0.78</td>
<td>0.86</td>
</tr>
<tr>
<td>NEU-F01</td>
<td>11</td>
<td>1.00^a</td>
<td>1.00</td>
<td>0.82</td>
<td>0.67</td>
</tr>
<tr>
<td>VOS-NL00</td>
<td>9</td>
<td>0.89^a</td>
<td>0.71</td>
<td>0.40</td>
<td>-</td>
</tr>
</tbody>
</table>

The mean proportion of KBH-NL00 alleles in the F2 generation was 0.542 (SD = 0.164; Fig. 3) showed no significant deviation from that expected under equal recovery (F = 0.075, n.s.). Hence, there was no evidence for genomic incompatibilities between the two modes of reproduction.

Duration and latencies are given in seconds ± SD. When statistical differences are present, different superscript letters indicate different groups.

### Discussion

**Sexual function of PI *Wolbachia*-infected *L. clavipes***

Sexual function of PL-infected species differs from that of uninfected species. In PI-infected species, males often mate with PI-infected females, whereas in uninfected species, males mostly mate with uninfected females. However, PI-infected males show a higher rate of successful copulation than uninfected males, indicating that PI-infected males are more successful than uninfected males in obtaining sperm from PI-infected females.

**Recovery of parental alleles**

The proportion of parental alleles in the F2 generation was not significantly different from that expected under equal recovery (F = 0.075, n.s.). Hence, there was no evidence for genomic incompatibilities between the two modes of reproduction.

**Sperm use by infected females**

PI-infected females do not use the sperm of PI-infected males. Sperm use by infected females was lower than that of uninfected females (F = 0.26, P < 0.001), indicating that PI-infected females are less likely to fertilize their eggs with PI-infected males.

**Spreads of PI-infected males**

PI-infected males show a higher rate of successful copulation than uninfected males, indicating that PI-infected males are more successful than uninfected males in obtaining sperm from PI-infected females.

**Infected females mated to arrhenotokous males**

Infected females mated readily with arrhenotokous males, but did not produce hybrid offspring. Paternal markers were not incorporated into the genome of the offspring, indicating that PI-infected females do not fertilize their eggs when mated to arrhenotokous males.
arrhenotokous and thelytokous lines is most likely due to small age differences between the tested wasps, as age was not strictly controlled for.

**Sex ratio**
The sex ratios produced by males from thelytokous lines are higher than those produced by arrhenotokous males. Because *L. clavipes* has haplodiploid sex determination (i.e. females develop from fertilized and males from unfertilized eggs), a higher sex ratio implies a lower fertilization success for the thelytokous males. As no difference in sex ratio was found between the two arrhenotokous lines, the effect is presumably due to the difference in reproductive mode rather than to the differences among populations.

For most lines, the sex ratio of the first clutch was lower than the sex ratio of the second clutch. Because females mated only once, the number of sperm in the spermatheca diminishes after ovipositing the first clutch which could have limited the production of female

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Reproductive mode, mean sex ratio of first and second clutch, and mean total broodsize produced by DC-E00 females after mating to males from different populations.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Population</td>
<td>Reproductive mode</td>
</tr>
<tr>
<td>CCAP-E00</td>
<td>A</td>
</tr>
<tr>
<td>DC-E00</td>
<td>A</td>
</tr>
<tr>
<td>DB23/9-NL99</td>
<td>T</td>
</tr>
<tr>
<td>DBK-NL00</td>
<td>T</td>
</tr>
<tr>
<td>GBW-NL00</td>
<td>T</td>
</tr>
<tr>
<td>KBH-NL00</td>
<td>T</td>
</tr>
<tr>
<td>VOSB-NL00</td>
<td>T</td>
</tr>
<tr>
<td>NEUVIC-F01</td>
<td>T</td>
</tr>
<tr>
<td>RENNES-F01</td>
<td>T</td>
</tr>
</tbody>
</table>

A, arrhenotokous; T, thelytokous. Standard error in parentheses. Different letters indicate statistical differences in broodsize at *P* < 0.05.

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Generalized linear mixed model for sex ratios. Note that the parameter estimates of fixed effects are given as logits and need to be transformed to obtain predicted mean sex ratios.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effect</td>
<td>Estimate*</td>
</tr>
<tr>
<td>Fixed</td>
<td></td>
</tr>
<tr>
<td>Arrhenotokous</td>
<td>–0.526</td>
</tr>
<tr>
<td>Thelytokous</td>
<td>1.547</td>
</tr>
<tr>
<td>First clutch</td>
<td>–0.504</td>
</tr>
<tr>
<td>Second clutch</td>
<td>–0.006</td>
</tr>
<tr>
<td>Random</td>
<td></td>
</tr>
<tr>
<td>Female/thelytokous</td>
<td></td>
</tr>
<tr>
<td>Male/thelytokous</td>
<td></td>
</tr>
<tr>
<td>Residual variance</td>
<td></td>
</tr>
</tbody>
</table>

*Back-transformed fixed effect parameter estimates to sex ratio, with brood size covariate fixed at the average over the total dataset: arrhenotokous, first clutch: 0.195; second clutch: 0.287; thelytokous, first clutch: 0.659; second clutch: 0.761.

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Sperm utilization by <em>Wolbachia</em>-infected thelytokous <em>Leptopilina clavipes</em> females when mated with arrhenotokous males</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of females tested</td>
<td>No. of unique loci</td>
</tr>
<tr>
<td>Mother</td>
<td>Father</td>
</tr>
<tr>
<td>BBH-NL00</td>
<td>Moll1-E00</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>KBH-NL00</td>
<td>DC-E00</td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>HOD-NL00</td>
<td>Moll1-E00</td>
</tr>
</tbody>
</table>

offspring in the second clutch (Godfray, 1994). Another possibility is low sperm viability, because of which many sperm could have been rendered nonfunctional between oviposition of the first and second clutch. Only NEUVIC-F01 and DB23/99-NL99 showed a lower second sex ratio.

Measured over all lines, brood size had a negative effect on sex ratio. The opposite effect might be expected when sperm is the limiting factor for sex ratio. However, according to the local mate competition (LMC) theory (Hamilton, 1967) a female should produce just enough males to fertilize her own daughters. If the LMC model is modified to include variation in brood size, it predicts large clutches to be relatively female biased (Werren, 1980), i.e. a negative effect of brood size on sex ratio.

In contrast to the observations in the courtship experiment, males from all the tested thelytokous lines mated and produced offspring, including males from GBW-NL00. The reasons for this difference are unclear, but may relate to differences in air pressure between the experiments. Leptopilina are known to adjust their behaviour to fluctuations in barometric pressure (Roitberg et al., 1993) which we could not control for in our experiments. This is further supported by a futile attempt to start the sex ratio experiment in the winter of 2003. During this trial, courtship was only displayed by the arrhenotokous males (data not shown).

Thus far, L. clavipes is the only species in which a difference in fertilization success was found between males of arrhenotokous lines, and those induced from allopatric thelytokous populations. Males induced from thelytokous Trichogramma deion, T. pretiosus, Apoanagyrus diversicornis and Telonomus nawai populations do not produce a higher sex ratio than their arrhenotokous conspecifics when mated to arrhenotokous females (Stouthamer et al., 1990b; Pijs et al., 1996; Arakaki et al., 2000).

Sperm use

The AFLP analyses indicate that Wolbachia-infected L. clavipes females do not incorporate paternal genes into their offspring after mating to arrhenotokous males. Thus although thelytokous L. clavipes females readily mate with arrhenotokous males, they do not use the sperm to fertilize their eggs. Failed attempts to establish cured thelytokous lines (data not shown) support these results. Failure to establish arrhenotokous lines from PI Wolbachia-infected lines has been reported for many species. For example, Arakaki et al. (2000) found no evidence of sperm use after mating by cured females from the parasitoid Telonomus nawai. In other species sperm transfer does not occur because females are not receptive to courting males [e.g. Apoanagyrus diversicornis (Pijs et al., 1996) and Muscidifurax uniraptor (Gottlieb & Zchori-Fein, 2001)].

The pattern of sexually functional males and dysfunctional females found in L. clavipes is consistent with that observed in other parasitoids where completely infected and uninfected populations occur allopatrically (Pijs et al., 1996; Arakaki et al., 2000). Also in species where only infected populations were studied, matings between infected females and cured males do not result in fertilized eggs (Zchori-Fein et al., 1992, 1994, 1995; De Barro & Hart, 2001; Gottlieb & Zchori-Fein, 2001; Weeks & Breeuwer, 2001). Only infected females from populations where infected and uninfected females co-exist fertilize their eggs after mating with cured thelytokous or arrhenotokous males. Thus far, these mixed populations are only known from several Trichogramma species and cured infected females from these populations proved to be fully functional (Stouthamer et al., 1990a,b, 2001; Stouthamer & Luck, 1993; van Meer, 1999).

The pattern of nonfertilizing females and (partly) functional males is in line with the 'virginity-mutant' hypothesis of Huigens & Stouthamer (2003). According to this scenario, a virginity mutation in L. clavipes females was selected for in the initial stages of the infection. The virginity mutation is then fixed in an all-female population, where male sexual function is no longer actively maintained by selection and is prone to decay due to the random accumulation of mutations (Muller, 1949). However, mutation accumulation predicts that losses of unused functions should accumulate stochastically (Cooper & Lenski, 2000), and different male sexual functions should decay in different thelytokous lines.

We found a reduction in male fertilization capacity in thelytokous lines of L. clavipes from different geographic
origins. There are several explanations possible for this pattern. First, the allopatric modes of reproduction may have been separated long enough for incompatibilities to arise, which can be expressed as a reduction in hybrid (female) offspring. However, in the present study we found no evidence for genomic incompatibilities. The observed pattern of equal recovery of maternal and paternal markers is consistent with other intraspecific crosses in Hymenoptera (Hunt & Page, 1995; Antolin et al., 1996; Laurent et al., 1998), but not with interspecific crosses (Gadau et al., 1999). In addition, the genetic distance between the two modes of reproduction is low when compared with closely related outgroups (Pannebakker et al., 2004c).

A second explanation is that the infected L. clavipes lines originate from the same infection and hence show the same phenotypes. Genetic analysis, however, showed that there are at least two major clonal genotypes present in north-western Europe (Pannebakker et al., 2004c) which were both used in this study (DB/23/9-NL99 & VOS-NL00 vs. DBK-NL00, GBW-NL00, KBH-NL00, NEUVIC-F01 & RENNES-F01). Hence, the observed similarity in phenotypes between the lines is not expected based on their genotypic diversity.

A third explanation for the similar pattern in male fertility among different thelytokous lines is that it is the result of indirect selection due to antagonistic pleiotropy. This may then lead to the loss of sexual function in males due to selection on genetically linked traits that are adaptive in the parthenogenetic strains. When assumed that selection pressures for these traits are similar in different populations, the same sexual trait is expected to decline in different populations (cf. Cooper & Lenski, 2000).

Under the ‘virginity-mutant’ hypothesis of Stouthamer & Huijgens (2003), strong selection on female virginity is expected in populations in the early stages of the infection. The data on L. clavipes show that females from infected populations do not fertilize their eggs after mating, while males from all infected populations examined show a reduced fertility. This pattern could indicate a role for antagonistic pleiotropy between the nonfertilizing mutation in females and the reduction in fertility in males. Linkage analysis in the KBH-NL00 strain identified a single quantitative trait locus (QTL) of large effect for the reduction in male fertility (Pannebakker et al., 2004a), which suggests a key mutation in a single gene.

The role of antagonistic pleiotropy can be determined by comparing QTL involved in reduced male fertility in males induced from other infected populations than KBH-NL00. When the reduced sexual functionality is due to antagonistic pleiotropy, a QTL at the same genomic location is expected in males induced from other infected populations assuming similar selection pressures as predicted by the ‘virginity-mutant’ hypothesis. If the reduction in male fertility is the result of antagonistic pleiotropy, the same gene or pathway should be responsible for nonfertilization in infected females. Examples of genes in other taxa where a single mutation would have such an effect include the gene coding for the sperm-egg attachment mediating protein binding in sea urchins (Palumbi, 1999), and various male-female-sterile (mfs) genes in Drosophila (i.e. mfs(1)6E, mfs(2)350, mfs(3)73A and mfs(3)L). Gelbart et al., 2003) that have different functions in male and female reproduction (Fukunaga, 1980; Lopez et al., 2001). It should be noted however, that linkage disequilibrium between the genes coding for nonfertilization in females and the reduced males fertility in males creates a similar pattern.

In conclusion, we found evidence for the existence of a single reproductive barrier between allopatric athenototous and thelytokous L. clavipes populations. Females from Wolbachia-infected populations do not use sperm after mating to arrhenotokous males. Males induced from infected populations show normal courtship behaviour and are capable of fertilizing arrhenotokous females; however, with a lower fertilization success than arrhenotokous males.

Although the lack of genomic incompatibilities and a low genetic distance indicates the two allopatric modes of reproduction belong to a single species, the reproductive barrier between them may be a first step in the process of speciation. Due to sexual degradation, Wolbachia-infected populations can become ‘locked’ into thelytokous reproduction even if the infection is lost, eventually resulting in speciation between infected and uninfected populations (Werren, 1998; Bordenstein, 2003). In natural infected L. clavipes populations, presumably due to inefficient transmission of the Wolbachia bacteria, males are occasionally produced (Driessen et al., 1990). Although these males can produce viable offspring when mated to arrhenotokous females, (unidirectional) gene flow between the two modes of reproduction is prevented by two additional barriers: a disjunct distribution and differences in phenology (Pannebakker et al., 2004c). To unravel the evolutionary scenario, further detailed research into the ecological differences between the two modes of reproduction is needed, as well as an investigation of the Wolbachia infection history. Moreover, genetic characterization of the nonfertilization trait in infected females, as well as that of the reduced male fertility in different thelytokous lines can help to clarify the genetic mechanism and evolutionary history involved in the loss of sexual function in PI Wolbachia-infected species.

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