#### **ORIGINAL RESEARCH**



# Symbiont-Driven Male Mating Success in the Neotropical *Drosophila* paulistorum Superspecies

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#### **Abstract**

Microbial symbionts are ubiquitous associates of living organisms but their role in mediating reproductive isolation (RI) remains controversial. We addressed this knowledge gap by employing the *Drosophila paulistorum-Wolbachia* model system. Semispecies in the *D. paulistorum* species complex exhibit strong RI between each other and knockdown of obligate mutualistic *Wolbachia* bacteria in female *D. paulistorum* flies triggers loss of assortative mating behavior against males carrying incompatible *Wolbachia* strains. Here we set out to determine whether de novo RI can be introduced by *Wolbachia*-knockdown in *D. paulistorum* males. We show that *Wolbachia*-knockdown *D. paulistorum* males (i) are rejected as mates by wild type females, (ii) express altered sexual pheromone profiles, and (iii) are devoid of the endosymbiont in pheromone producing cells. Our findings suggest that changes in *Wolbachia* titer and tissue tropism can induce de novo premating isolation by directly or indirectly modulating sexual behavior of their native *D. paulistorum* hosts.

Keywords Wolbachia · Drosophila · Reproductive isolation · Premating isolation · Pheromones

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#### Introduction

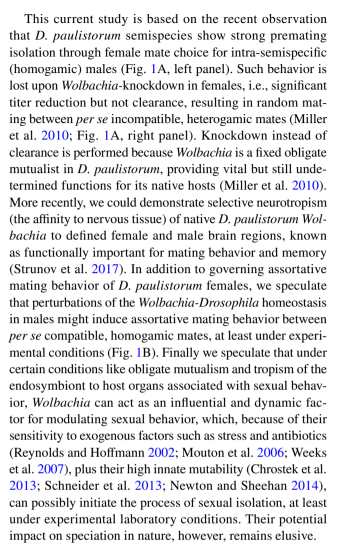
Potential impacts of microbial symbionts as drivers of host speciation have been discussed frequently (Bordenstein et al. 2001; Jaenike et al. 2006; Telschow et al. 2007, 2014; Brucker and Bordenstein 2012; Gebiola et al. 2016), but their broader evolutionary significance fostering speciation remains controversial (Brucker and Bordenstein 2013, 2014; Chandler and Turelli 2014; Turelli et al. 2014; Najarro et al. 2015; Leftwich et al. 2017), and standard models of speciation commonly disregard such impacts (Coyne and Orr 2004; Brucker and Bordenstein 2012; Miller and Schneider 2012). Several theoretical models have nominated the maternally transmitted reproductive tract endosymbiont Wolbachia as a candidate worthy of consideration (Telschow et al. 2007, 2014). The  $\alpha$ -proteobacterium Wolbachia is arguably the most prevalent intracellular invertebrate infection on the planet, infecting as many as 40% of all terrestrial arthropods (Zug and Hammerstein 2012). Wolbachia have attracted attention as participants in arthropod symbioses because they successfully manipulate host biology in manifold ways, ranging from reproductive parasitism like cytoplasmic incompatibility (CI), male killing, feminization and



parthenogenesis to nutritional supplementation, fecundity and protection from pathogens and parasites (Werren et al. 2008; Teixeira et al. 2008; Brownlie et al. 2009; Fast et al. 2011; Moriyama et al. 2015). Although *Wolbachia* are mainly found as facultative endosymbionts in most insect hosts (Martinez et al. 2015 and references therein), they also can evolve fixed obligate associations such as in the parasitoid wasp *Asobara* (Dedeine et al. 2001), bedbugs (Hosokawa et al. 2010), or neotropical fruit flies belonging to the *Drosophila paulistorum* species complex (Miller et al. 2010).

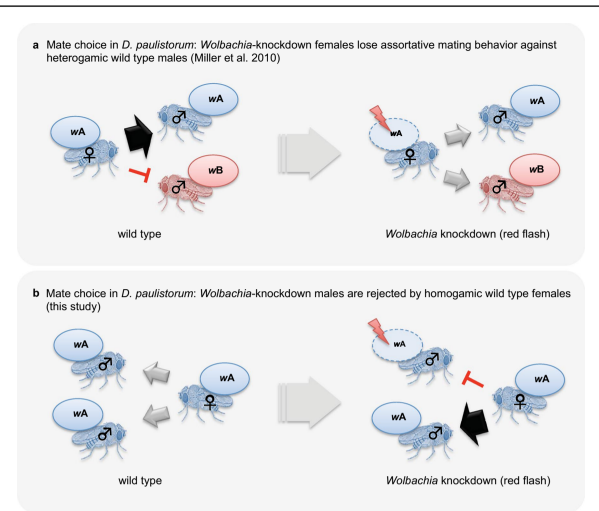
In the latter case, this superspecies is under incipient speciation in the wild and consists of six semispecies (Dobzhansky and Spassky 1959), named Amazonian (AM), Andean Brazilian (AB), Centroamerican (CA) Interior (IN), Orinocan (OR) and Transitional (TR), expressing strong RI at the pre- and postzygotic level in inter-semispecies crosses (reviewed in Ehrman and Powell 1982). As recently found they all carry different loads of obligate mutualistic but distinctive Wolbachia strains (Miller et al. 2010), which can cause strong cytoplasmic incompatibilities (embryonic mortality) plus complete hybrid male sterilities in reciprocal crosses between different semispecies in the laboratory. Because Wolbachia are obligate endosymbionts in this system, no uninfected flies can be generated (Ehrman 1968; Kernaghan and Ehrman 1970; Miller et al. 2010) to test for classic bidirectional CI as feasible in facultative symbioses such as in Culex pipiens mosquitos (reviewed in Werren et al. 2008). The association between the obligate Wolbachia symbiont (earlier designated Mycoplasma-like-organisms, MLOs; reviewed in Ehrman and Powell 1982) and the induction of postmating isolation, however, is based on earlier and recent findings that hybrid lethality and male sterility are partly reversible upon mild antibiotic or heat treatments of the parents before mating (Ehrman 1968; Kernaghan and Ehrman 1970; Miller & Schneider, unpublished).

In two earlier publications, Dobzhansky and collaborators (Dobzhansky and Pavlovsky 1966, 1971) observed the spontaneous evolution of de novo postmating isolation as some lines, which were originally fully compatible with the Orinocan reference strain O11, were no longer compatible with this semispecies or any other semispecies of *D. paulistorum*. This resulted in occurrence of high embryonic F<sub>1</sub> mortality and complete male hybrid sterility, presumably because of drift and isolation in these lines in the laboratory. However, the role of endosymbionts on the formation of premating mechanisms has been less studied in the past (Dobzhansky and Pavlovsky 1966, 1971; Miller et al. 2010). Importantly, in facultative symbioses, where Wolbachia has not reached fixation in their natural hosts as in D. melanogaster or D. yakuba group species, the authors did not detect any Wolbachia effect on premating isolation (Sharon et al. 2010; Arbuthnott et al. 2016; Cooper et al. 2017).



The key components of premating sexual isolation mechanisms are visual, acoustic, and chemical courtship signals. In many insects, sexual pheromones play an important role as olfactory signals influencing mate choice (Blomquist and Bagnères 2010; Chung and Carroll 2015; Dembeck et al. 2015). In D. paulistorum, males from the six semispecies exhibit characteristic sexual pheromone profiles (same compounds but different quantities), including four male-specific long-chained ester compounds, which are recognized by females (Kim et al. 2004; Chao et al. 2010). In contrast to other *Drosophila* species such as *D. mela*nogaster, no female-specific compounds are present in this system. Hence, the authors of previous studies (Kim et al. 2004; Chao et al. 2010) concluded that, based on chemical profiles, D. paulistorum females accept homogamic (compatible), but reject heterogamic (incompatible) males as mates (Chao et al. 2010). Forced mating between heterogamic mates carrying incompatible Wolbachia strains is highly detrimental for both host and symbionts by triggering high levels of embryonic lethality and complete male sterility among F<sub>1</sub> hybrids (Ehrman 1968; Kernaghan and





**Fig. 1** Impact of *Wolbachia* on mate choice in *D. paulistorum.* **a** Mate choice in *D. paulistorum: Wolbachia*-knockdown females lose assortative mating behavior against heterogamic wild type males. Left: wild type females (blue) infected with *Wolbachia* A (wA) express strong assortative mating behavior (red block) against males of a different semispecies (red) carrying a different, incompatible *Wolbachia* (wB). Right: *Wolbachia* kd females (symbolized with red flash) with reduced titers of wA lose assortative mating behavior (grey arrows) and hence mate randomly with males from same (blue) but also from different/incompatible (red) semispecies. Female knockdown phenotype: loss of assortative mating behavior. Data

Ehrman 1970; Miller et al. 2010). Hence strong selection should act on both intimate partners to evolve or maintain such mechanisms for mate recognition.

Here we experimentally investigated whether bacterial symbionts are capable of initiating artificial de novo reproductive isolation (RI) by targeting host-derived olfactory cues. Therefore, we have (i) transiently knocked down obligate *Wolbachia* in *D. paulistorum* males, (ii) assayed their mating success in homogamic choice assays with wild type females, (iii) monitored their sexual pheromone profiles, and (iv) investigated the presence of *Wolbachia* in putative pheromone-producing cells.

taken from (Miller et al. 2010). **b** Mate choice in *D. paulistorum: Wolbachia*-knockdown males are rejected by homogamic wildtype females. Left: wild type males are accepted randomly (grey arrows) for mating by females of the same semispecies (all blue), carrying the same/compatible *Wolbachia* (both *wA*). Right: *Wolbachia* kd males (indicated by red flash) with reduced titers of *wA* are rejected as mate partners (red block) from wild type females of the same semispecies. Thickness of arrows indicates strength of mate choice between partners: thick black arrow=high mate preference, red block=assortative mating, and grey medium arrow=lack of mate choice, i.e., random mating. *kd* knockdown. (Color figure online)

### **Methods**

## Fly strains and generation of *Wolbachia*-knockdown (kd), axenic (et), and penicillin/streptomycin-treated (ps) flies

Reference strains for two semispecies from the *Drosophila paulistorum* superspecies were chosen for this study (Amazonian, A28 and Orinocan, O11; originally described in Burla et al. 1949; Dobzhansky and Pavlovsky 1966). The *Wolbachia* infection status (*w*Pau) of these *D. paulistorum* strains was previously described in Miller et al.



2010. In brief, A28 carries very low densities of the Wolbachia wPauAM strain, and O11 is infected with the high-titer wPauOR strain (Miller et al. 2010). Wolbachia positive controls for Polymerase chain reaction (PCR) assays were D. willistoni P98 and JS6.3, both lab strains carrying native high-titer Wolbachia (wWil; Miller and Riegler 2006). The D. melanogaster H2-strain (wMel; Miller and Riegler 2006) was used as an additional Wolbachia positive control. Wolbachia negative controls for PCR were D. simulans NouméaTC (Poinsot et al. 2000), D. melanogaster w<sup>1118</sup> (BDSC, USA), and D. willistoni Willi3 (14030-0811.2 DSSC, USA). Flies were reared on Formula 4–24 Drosophila instant food (Carolina, USA) at 24–25 °C on a 12 h light–dark cycle.

Wolbachia-knockdown lines A28kd and O11kd were generated by mass-treating wild type flies with 0.2% (w/v) rifampicin (Duchefa, Netherlands) added to Formula 4-24 Drosophila instant food (Carolina, USA) over three sequential generations  $(T_1-T_3)$ . The low dosage of the antibiotic is sublethal and allows for reduction of mutualistic Wolbachia in D. paulistorum (Miller et al. 2010). Offspring from T<sub>3</sub> parents were then transferred onto antibiotic-free food for more than 10 generations under a mass-rearing regime designated as knockdown pools (A28<sup>kd</sup>, O11<sup>kd</sup>; Supporting Information Fig. S1). In parallel, F<sub>2</sub><sup>kd</sup> was used to initiate 8–10 isofemale lines per semispecies (A28<sup>kd-i</sup>, O11<sup>kd-i</sup>; Supporting Information Fig. S1). Biological assays were performed at different generations between  $F_4^{kd}$  and  $F_{13}^{kd}$  and their respective Wolbachia load was monitored by qPCR (see below). Gut flora restoration (gfr) lines were generated by feeding the native gut microbiome to knockdown flies as follows: A28<sup>wt</sup> and O11<sup>wt</sup> virgin females were kept on instant food for 2-3 days to collect feces. Flies were then removed and the corresponding knockdown (kd) flies were transferred onto the feces-containing food vials. After egg deposition adults were removed and emerging flies were consequently used as gfr lines. Axenic ('gut microbe-free flies') were generated by consequently washing freshly collected eggs from A28wt and O11wt with 70% ethanol for 5 min to surface-sterilize them. This treatment prevented larvae ingesting microbes from the outer layer of the eggshell when hatching. Collected eggs were then transferred into food vials and hatching F<sub>1</sub> and F<sub>2</sub> adults (A28<sup>et</sup>, O11<sup>et</sup>) were consequently used for mate choice assays. Penicillin/ streptomycin-treated (ps) flies were generated by supplementing Formula 4-24 Drosophila instant food (Carolina, USA) with a 1:100 dilution of a 100× pen/strep stock solution (10,000 units/ml penicillin, 10,000 µg/ml streptomycin). A28<sup>wt</sup> and O11<sup>wt</sup> were kept on this food for at least 1 week to lay eggs and the hatching F<sub>1</sub> (A28<sup>ps</sup>, O11<sup>ps</sup>) was consequently used for mate choice assays against the wt counterparts. Strains are reported in Table S1.



### Quantification of wPau Wolbachia

DNA was extracted from a pool of ten 3-day old flies using Gentra Puregene chemistry (Qiagen, Germany). Concentrations of DNA were measured on a Nanodrop 2000 spectrophotometer. Consequently, diagnostic Wolbachia-PCR (quantitative real time PCR) was performed using the Wolbachia outer surface protein gene wsp (Yamada et al. 2007), and the Wolbachia-specific 16S rRNA gene (16SW\_RTf 5'-CCTGATCCAGCCATGCCGCAT-3', 5'-16SW\_RTr CGGCTGCTGGCACGGAGTTA-3'). The wsp primer set generates a 69 bp amplicon, and the 16S rRNA set produces a 77 bp fragment. Wolbachia titers obtained with MxPro QPCR v4.10 Software (Agilent Technologies, USA) were normalized against *Drosophila* ribosomal protein RPS17 (RPSmel f 5'-CACTCCCAGGTGCGTGGTAT-3', RPSwil\_r 5'- GGAAACGGCGGGCACGTA-3'). A temperature profile of 95 °C for 3 s, 60 °C for 20 s, and 72 °C for 6 s was used for 45 cycles. All samples were run in duplicates on a Stratagene MxPro4000 cycler. Quantitative PCR with wsp was run to confirm results obtained with the 16S rRNA primer set. Only results for the latter one are presented in the manuscript.

## Measuring sexual isolation via multiple mate choice assays

Mate choice assays between wild type, knockdown and control assays involved double blind direct multiple-mating observations carried out mornings at room temperature in daylight. Virgin flies were aged 2-3 days (females isolated from males), and half were marked via distal wing clips before running the choice assay. These marks were rotated (wing to wing and knockdown to wild type). Such minute abrasions have tested neutral regarding behavioral influences in this superspecies (Dobzhansky and Pavlovsky 1966; Leonard and Ehrman 1983). For each mate choice assay five replicas and 120 mating events (240 individual flies) were scored (12A  $\mathcal{Q}\mathcal{Q}$  + 12B  $\mathcal{Q}\mathcal{Q}$  + 12A  $\mathcal{Q}\mathcal{Q}$  + 12B  $\mathcal{Q}\mathcal{Q}$  differentiated by alternating wing clips; Supporting information Fig. S2a, b). Flies were placed (females first) in mating observation chambers (10 cm in diameter) without anesthetization and the following parameters were scored until all flies copulated in approximately 30-40 min: the time each mating took place (from start of observations; each copulation approximately lasted 15-17 min); its sequence among other copulae which occured; where in the chamber the mating pair was located; the kind of female involved; and the kind of male involved. Recording the location of each copula, even upside down, prevented scoring a copula more than once. Sexual isolation index (SII) was computed according to the following formula (Malogolowkin-Cohen et al. 1965).

SII = (nho-nhe)/N, where *nho* is the number of homogamic matings, *nhe* is the number of heterogamic matings, and N the total number of matings. SIIs range from -1.00(preference for unlikes, heterogamy) through 0 (random mating) to +1.00 (preference for likes, homogamy). This experimental design is not devoid of bias. Indeed, when the number of partners is finite and remating is limited, the choice of the last individuals is conditioned by the choice of the previous ones. This could lead to erroneous signals for assortative mating, e.g. when both males and females from the same population tend to mate earlier than those from the other population (Ehrman and Parsons 1981). However, unbiased rates of assortative mating could be estimated by accounting for the order of mating pairs, using a statistical model described extensively in Supporting Methods. This method provides the maximum-likelihood estimates of the assortative mating coefficient (called eSII, or h in the model and Supporting Methods) and accounts for the possibility of remating in males. Several remating rates were tested and did not significantly influence the estimation of SII (see Fig. S3), hence in practice a remating rate of 0.5 was chosen for the general analysis (Tables S2–S4). To avoid convergence issues, the female remating rate was set to 0.001. Two additional parameters (biases in mating order for males and females) were also estimated by the model (Table S5). However, their biological interpretation is not that straightforward as they may correspond to either a general preference for one population of one sex, or a different mating speed between both populations. The statistical departure of the SII from 0 was estimated by a likelihood ratio test. Simulations showed that this procedure efficiently corrects for such mating order biases, and was even slightly more powerful than homogeneity (Fisher) tests when mating order was unbiased.

The model was implemented in R version 3.1.1 (R Core Team 2016), the code is available in Supporting Methods, along with the raw data. A comparison of the uncorrected SII and the model-estimated SII is provided in Supporting Information Fig. S3 and Tables S2–S4. Hereafter, we will refer only to the estimated SII that will be simply called eSII.

# Analysis of cuticular hydrocarbons (CHCs) *via* gas chromatography-ion trap mass spectrometry (GC-MS)

Male virgin flies were collected, isolated from females, and aged to 3 days. Flies were pooled in glass vials and submerged in 1 ml of HPLC-grade hexane (Carl Roth, Germany). Replicates consisted of 10 males per sample. We did nine replicates for A28<sup>wt</sup> and 10 replicates for each A28<sup>kd</sup>, O11<sup>wt</sup> and O11<sup>kd</sup>. Two µg of octadecane (C18) per sample was added as the internal standard for absolute quantification of CHCs. Extraction of CHCs was performed for 10 min at room temperature under constant agitation, after which

the flies were removed. Extracts were evaporated to about 20-30 µl of hexane under a constant stream of argon, and then transferred to a 150 µl GC-µ-vial (CZT, Germany). 1 µl aliquots were injected into a Varian 450GC gas chromatograph coupled to a Varian 240MS mass spectrometer (Agilent Technologies, Germany). A DB5-MS capillary column (30 m×0.25 mm diameter, film thickness: 0.25 μm, Agilent Technologies, Germany) was used and the GC was programmed from 150 °C to 300 °C at 15 °C/min with a 27 min final isothermal hold. Helium, with a constant flow rate of 1 ml/min, was used as carrier gas. Recording of mass spectra was performed using electron ionization (EI-MS) in external ionization mode and data acquisition plus quantifications were done with MS Workstation v6.9.3 Software (Agilent Technologies, Germany). Consequently, peaks were identified by their mass spectra in comparison to previously published hydrocarbon profile analyses of D. paulistorum (Kim et al. 2004; Chao et al. 2010) and D. melanogaster (Ueyama et al. 2005). Hydrocarbon quantities were calculated from peak areas and then centered log-ratio-transformed according to Aitchinson 1986. To test for differences in chemical profiles between wild type and knockdown individuals, principal component analyses (PCAs) based on eigenanalysis of covariances were performed to reduce numbers of variables. Consequently, resulting PCs were used for discriminant analyses (DAs), to test for among-group differences.

### Fluorescence in situ hybridization (FISH) on D. paulistorum oenocytes

Ten to 15 female and male abdomen per semispecies were dissected in RNase-free 1x phosphate buffered saline (PBS). After removing inner organs, cuticles were fixed in 3.7% formaldehyde in RNase-free PBS for 20 min at room temperature and consequently washed with PBTX (1 × PBS, 0.3% Triton-X 100). After permeabilization with 70% ethanol overnight at 4 °C, samples were hybridized overnight in 10% formamide, saline sodium citrate (SSC) and 0.5 nmol of CAL Fluor Red 590-labeled customized Wolbachia 16-23S rRNA probe (Biosearch Technologies, USA; Schneider et al., in press). Samples were then washed in 10% formamide and SSC, stained with DAPI-SSC (1 µg/ml) and mounted in Roti®-Mount FluorCare (Carl Roth, Germany) on sterilized microscope slides. Cuticles were analyzed on an Olympus FluoView confocal microscope. Beam paths were adjusted to excitation/emission peaks of used fluorophores: 569/591 nm for CAL Fluor Red 590 (Wolbachia), and 350/450 nm for 4',6-diamidin-2-phenylindol (DAPI). Images were processed with Fiji software (http://fiji.sc). Hybridization experiment was repeated twice and a minimum of 15 flies was assayed in each experiment.



#### Results

### Knockdown of obligate Wolbachia in Drosophila paulistorum semispecies is of transient nature

To test whether manipulation of native Wolbachia titers and associated disruption of the host-symbiont homeostasis can trigger de novo reproductive isolation (RI) in D. paulistorum, we generated Wolbachia-knockdown (kd) males of two different semispecies (Amazonian: A28, and Orinocan: O11) by mild rifampicin treatment for three consecutive generations (T1-T3) and in the following generations on regular media without antibiotics (F<sub>1</sub><sup>kd</sup>-F<sub>14</sub><sup>kd</sup>) for restoration. For details see Supporting Information Table S1, Fig. S1. Global Wolbachia titers of these males (A28kd, O11kd) were tested against wild type (wt) controls (A28<sup>wt</sup>, O11<sup>wt</sup>) in quantitative real time PCR (qPCR) assays targeting the Wolbachia 16S ribosomal RNA gene. Assays obtained from F<sub>7</sub><sup>kd</sup> (knockdown generation 7) revealed massive titer reduction to about 5 and 14% of the average native wt Wolbachia titers (Fig. 2a, b).

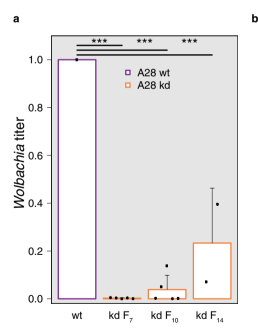
To further assess whether *Wolbachia* titers return to native levels in later generations post kd, we tested symbiont load by  $F_{14}$  and earlier generations. Whereas in O11

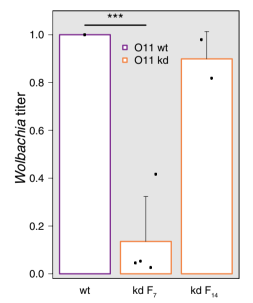
the global symbiont titer was close to wt-levels at  $F_{14}$  (90%; Fig. 2b), A28 flies have gradually reached only 23% of their wt-titer at this time point (Fig. 2a).

## Mate choice assays reveal de novo assortative mating phenotype of wild type (wt) females towards Wolbachia-knockdown males

Among sympatric *D. paulistorum* semispecies, no hybrids are formed inter-semispecifically in nature (Dobzhansky and Spassky 1959). When performing inter-semispecific mate choice assays involving A28<sup>wt</sup> and O11<sup>wt</sup> semispecies, we observed strong RI, revealed by a high estimated Sexual Isolation Index (eSII) (Ehrman 1965; Ehrman and Powell 1982; Supporting Information: Table S2, assays 1, 2). In contrast, intra-semispecific control assays showed no assortative mating in standard interbreeding 'pool' lines (Fig. 3a, b wt×wt (purple) and Supporting Information Table S2, assays 3, 4). Results of statistical testing are summarized in Table S2.

In order to test our hypothesis that manipulation of the native *Wolbachia* titer and the associated disruption of the host-symbiont homeostasis in *D. paulistorum* might trigger de novo premating isolation intra-semispecifically, we assayed mating behavior of *Wolbachia*-kd pool lines towards wt flies of the same semispecies (Fig. 3a, b, orange, and Table S3,





**Fig. 2** Quantitative analysis of *Wolbachia* titers in wild type (wt) and knockdown (kd) *D. paulistorum* males. Bars represent *Wolbachia* titers measured *via* quantitative real time PCR on DNA from whole body extracts of A28 (a), and O11 (b) using *Wolbachia 16S* rRNA normalized against *Drosophila* ribosomal protein 17 (*RPS17*). Each purple bar represents the mean *Wolbachia* titer in wt males (mean from 8 to 10 biological replicates is plotted). Orange bars represent titers in  $F_7/F_{14}$  kd males (post treatment); individual biological repli-

cates are shown as black dots. In addition, for A28<sup>kd</sup>,  $F_{10}$  titers were tested. Kd-titers are relative to wt ones (set to 1). Each sample was run in technical duplicates. Error bars represent the standard deviation of the biological replicates. Asterisks indicate significance of differences between titer levels. *P* values were calculated using *Fisher's* exact test. *wt* wild type,  $F_7$ ,  $F_{10}$ ,  $F_{14}$  fly generations 7, 10 and 14. (Color figure online)



assays 1-6). Assays run with knockdown flies at generation F<sub>4</sub> or F<sub>5</sub> post-treatment, consistently showed high eSII, i.e. mating between likes rather than unlikes. This points towards de novo mate discrimination between wt and kd flies from the same semispecies that significantly differ in their symbiont load (kd  $F_7$  in Fig. 2a, b). However, at generation  $F_{13}$ homogamy was less pronounced among all of them, and eSII did not differ significantly from what is expected under random mating. Importantly, although O11 flies showed rapid increase of Wolbachia levels by kd F<sub>14</sub> (Fig. 2b), only a slower, gradual titer increase was observed in A28 (Fig. 2a). Hence the restoration of the random mating phenotype in O11 might directly correlate with the global restoration of the symbiont level at faster pace, whereas the reversion to random mating of A28 flies might depend on a critical titer threshold of Wolbachia and/or their tropism to sensitive host tissues such as oenocytes (see discussion).

To rule out that drift effects were responsible for observed de novo sexual isolation, we additionally assayed wt and kd lines that were reared as ten isofemale lines per semispecies (A28<sup>wt-i</sup>, O11<sup>wt-i</sup> and A28<sup>kd-i</sup>, O11<sup>kd-i</sup>, respectively). First, two out of ten randomly picked control isofemale lines (wt-i) representing each semispecies were tested against each other (intra) in  $F_{10}$  in order to verify random mating between them (Fig. 3c, d wt-i×wt-i, purple, and Supporting Information Table S2, assays 5–8). The observed low eSIIs indicated the absence of any drift-dependent sexual isolation. To test for emerging sexual isolation between kd-isofemale lines upon drift (as a response to Wolbachia-kd and consequent restoration) we performed intra-semispecific choice assays using independent F<sub>8</sub> isofemale knockdown lines (kd-i). Surprisingly in this case the assays revealed significant isolation between them (pvalues < 10<sup>-4</sup>; Fig. 3c, d, pink, and Supporting Information Table S3, assays 7, 8).

Taken together, control assays between independent wt isofemale lines rule out drift as a potential factor for triggering de novo premating sexual isolation observed between wt and kd lines, whereas independent knockdown lines have possibly drifted apart from each other upon isolation having potentially different levels and/or individual tissue tropisms of *Wolbachia* during restoration at F<sub>8</sub>. Overall, the emergence of high eSIIs in intrastrain assays suggests that even slight perturbations of the native *Wolbachia - D. paulistorum* homeostasis and/or tropism are sufficient to induce de novo RI in this system, even between individual knockdown isofemale lines. This latter and quite intriguing observation, however, will need further investigations, which are beyond the scope of our current analyses (see discussion).

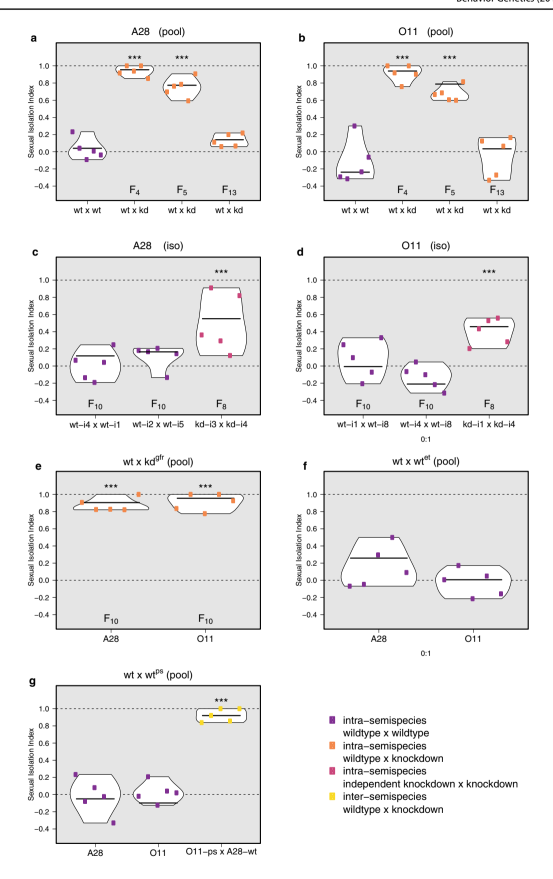
### No evidence for influence of gut microbiota on de novo assortative mating phenotype

Earlier and recent studies, however, have addressed the potential influence of diet and consequently the gut microbiome on host sexual isolation not only in D. pseudoobscura (Dodd et al. 1989), but also in highly inbred lines of D. melanogaster (Sharon et al. 2010; Ringo et al. 2011). Here, sexual isolation can be triggered by manipulating the fly's gut microbiome, based on a certain dietary regime. However, two recent studies have questioned the general role for gut bacteria or diet composition in driving reproductive isolation in D. melanogaster (Najarro et al. 2015; Leftwich et al. 2017). In the light of these still controversial findings obtained from inbred D. melanogaster strains, we have tested D. paulistorum gut flora restored (gfr) lines, where kd lines were fed with wt feces, in parallel with our unfed kd lines (see "Methods" for details). Assays with these lines and wt flies revealed almost complete SII suggesting that gut flora restoration does not affect the de novo assortative mating phenotype (p values  $< 10^{-4}$ , Fig. 3e, and Supporting Information Table S4, assays 1, 2). To further exclude an effect of the gut microbiome associated with the egg smear, we assayed axenic flies, i.e., eggs washed with ethanol (et, Fig. 3f, and Supporting Information Table S4, assays 3, 4), and penicillin/streptomycin-treated flies (ps, Fig. 3g, purple, and Supporting Information Table S4, assays 5, 6). Mating behavior with both sets were compatible with random mating (absence of assortative mating). Hence, in contrast to rifampicin, which acts on Wolbachia levels, neither sterilizing eggs, nor treatments with penicillin/streptomycin had any effect on mate choice behavior in this system. Furthermore, to test whether penicillin/streptomycin affects mate behavior of females we performed inter-semispecific control assay between wildtype A28 and penicillin/streptomycintreated O11 flies that still revealed high sexual isolation (ps, Fig. 3g, yellow, Supporting Information Table S4, assay 7, p value  $< 10^{-4}$ ), confirming that penicillin/streptomycinsensitive gut microbes in females have no effect on mating behavior in this system.

Importantly, we are aware that none of our control assays on its own can exclude the spurious presence of some hidden non-*Wolbachia* microbe affecting mate behavior. However, the combination of three independent control assays, i.e., gut-flora restoration, egg-surface sterilization and penicillin/streptomycin-treatment, strongly supports our hypothesis that endosymbiotic *Wolbachia*, which are resilient to penicillin/streptomycin and egg surface sterilizing treatments (Audsley et al. 2017; Leclerq et al. 2017; Ye et al. 2017), are interfering with mate discrimination of *D. paulistorum* in both sexes and not any other bacteria.

Concordantly, only rifampicin treatment massively reduces Wolbachia load and induces de novo assortative







**◄Fig. 3** Violin plots showing estimated sexual Isolation Index (eSII) in multiple choice assays between D. paulistorum wild type (wt), Wolbachia-knockdown (kd), axenic (et), and penicillin/streptomycintreated (ps) flies. (a, b) Intra-semispecific mate choice assays with pool lines. Wild type (wt) flies (control, in purple) or knockdown (kd) flies (in orange) were assayed at generations F4, F5, F13, against wt flies, for A28 and O11. (c, d) Intra-semispecific mate choice assays with isofemale lines. wt vs. wt (purple), and kd vs. kd (pink) mating choice assays were performed between different isofemale lines, respectively at generation F<sub>10</sub> and F<sub>8</sub>. e Intra-semispecific assays between wt and gut-flora-restored flies (gfr) performed at generation F<sub>10</sub> for A28 and O11. f Intra-semispecific assays between wt and ethanol-washed wt flies (et) performed at generation F<sub>2</sub> for A28 and O11. g Intra-semispecific assays between wt and penicillin/streptomycin-treated flies (ps), performed at generation F<sub>3</sub> for the A28 and O11. The last assay (yellow) is an additional inter-semispecific assay between wt and ps flies. For all assays, the model-estimated SII was used on either individual replicates (dots), or considering the 5 replicates (horizontal bars). Asterisks denote assays for which the likelihood ratio test (LRT) is highly significant (p values  $< 10^{-4}$ ), the null model being random mating (SII = 0). All the p values are reported in Tables S2–S4. (Color figure online)

mating behavior of wildtype females against knockdown males from the same semispecies (this study) that also selectively triggers loss of assortative mating behavior in knockdown females against males from a different semispecies (Fig. 3g, yellow; Miller et al. 2010).

## Sexual pheromone profiles are altered in *D.* paulistorum knockdown males

Based on the observed de novo assortative mating phenotype, we tested whether the pheromone composition of Wolbachia-kd males was altered in comparison with wt males. Males from all six D. paulistorum semispecies exhibit characteristic pheromone profiles (all share the same 15 major compounds but in varying quantities), including four malespecific long-chain esters, which are recognized by females as a semispecific blend because of the differences in their relative concentrations (Supporting Information Fig. S4; Kim et al. 2004; Chao et al. 2010). In contrast to other Drosophila species such as D. melanogaster, no female-specific compounds are known in *D. paulistorum* (Chao et al. 2010) and mate decisions appear—to our current knowledge—to be exclusively made by females (Ehrman 1965; Ehrman and Parsons 1981). Here we extended hydrocarbon profile analyses to a total of 27 compounds found in all tested semispecies by adding two new, unknown compounds (Supporting Information Table S6, compounds 9, 13) and by resolving the original C33, C35, and C37 peak clusters (C=chain length of the compound; 33, 35, and 37 carbon atoms).

In full agreement with earlier reports (Kim et al. 2004; Chao et al. 2010), discriminant analyses of sex pheromone profiles from A28<sup>wt</sup> and O11<sup>wt</sup> males revealed major differences among the semispecies (Supporting Information Fig. S5). In order to test for differences in male pheromone

profiles upon Wolbachia-kd, we compared profiles between F<sub>7</sub> kd males and wt males for each semispecies. Based on the 27 quantified components (Supporting Information Table S6), six principal components were extracted capturing 84.9% of the total variance. The discriminant analysis revealed statistically significant differences between both semi-species A28<sup>wt</sup> and O11<sup>wt</sup> males (Wilks' Lambda = 0.178,  $X^2 = 24.2$ , df = 6, p < 0.001) as well as between A28<sup>wt</sup> and A28<sup>kd</sup> males (Wilks' Lambda = 0.059,  $X^2 = 39.6$ , df = 6, p < 0.001) and also between  $O11^{wt}$  and  $O11^{kd}$  (Wilks' 446 Lambda = 0.117,  $X^2 = 34.3$ , df = 6, p < 0.001, Supporting Information Fig. S5). To further analyze differences in pheromone profiles between wt and kd males, we calculated the relative quantities of the four malespecific compounds in D. paulistorum male profiles (11-docosenyl-acetate, DA, 19-triacontenyl-acetate, TA, di-unsaturated acetate C<sub>32</sub>H<sub>60</sub>O<sub>2</sub>, DU, and methyl-(Z)-tetradecanoate, MD). As shown in Fig. 4, we observed most drastic changes in quantities of these compounds in A28<sup>kd</sup> males, where all four male-specific compounds were decreased between 9and 23-fold when compared to wt levels (Fig. 4a, p < 0.0001for all four compounds). Although not as drastic as A28<sup>kd</sup>, in O11<sup>kd</sup> males at least one male-specific compound also showed significant changes: compared to O11wt males, MD was reduced fivefold (Fig. 4b, p = 0.0399).

### Wolbachia colonize oenocytes in D. paulistorum

In insects, pheromone components are mainly synthesized in specialized cells located under the fly's cuticle, the oenocytes (reviewed in Blomquist and Bagnères 2010; Chung and Carroll 2015; Dembeck et al. 2015). We hypothesized that these specialized cells could serve a primary somatic target for Wolbachia to manipulate host pheromone profiles in D. paulistorum. Since male pheromone profiles were altered upon Wolbachia kd, we tested the infection status of male oenocytes in both D. paulistorum wt and kd individuals. As shown in Fig. 5, oenocytes from O11<sup>wt</sup> females (Fig. 5a, a') and males (Fig. 5b, b') harbor Wolbachia. We could also detect the symbiont in oenocytes of A28<sup>wt</sup> males (not shown). Post Wolbachia-kd oenocytes, however, are cleared from the symbiont (F<sub>4</sub> O11<sup>kd</sup>; males in Fig. 5c; females in c'). Since our mate choice assays indicated reversion to wt mating behavior, i.e., loss of de novo assortative mating, around F<sub>13/14</sub> post Wolbachia-kd, we tested whether this phenotype correlates with a potential recolonization of Wolbachia to oenocytes around this generation post kd. As shown in Fig. 5d, oenocytes of F<sub>13</sub> O11<sup>kd</sup> males are, at least partially, recolonized by the endosymbiont. Oenocytes of other Drosophila hosts like the wMel-infected D. melanogaster strain Harwich-2, (males in Fig. 5e, females in e') in which Wolbachia have evolved facultative symbiotic interactions and do not affect mate behavior, are devoid of the



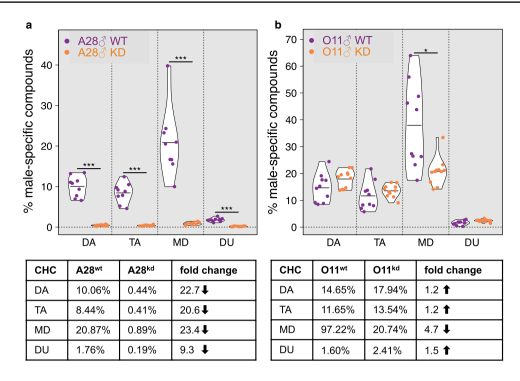


Fig. 4 Quantitative changes in *D. paulistorum* male-specific pheromone compounds upon *Wolbachia*-knockdown. Violin plots show percentage of the four male-specific ester compounds of the total pheromone profiles (27 compounds) of A28 ( $\bf a$ ) and O11 ( $\bf b$ ). Individual replicates for wild type controls (wt) are shown in purple, and in orange for  $F_7$  knockdown males (kd). Horizontal bars correspond to

the mean. Tables below violin plots list corresponding percent change of compounds plus n-fold increase ( $\uparrow$ ) or decrease ( $\downarrow$ ) between wt and kd. Asterisks indicate statistical significance based on two-tailed p values from *Student's t* tests. *DA* 11-docosenyl-acetate, *TA* 19-triacontenyl-acetate, *MD* methyl-(Z)-tetradecanoate, *DU* di-unsaturated acetate C<sub>32</sub>H<sub>60</sub>O<sub>2</sub>. (Color figure online)

symbiont. This is in contrast to *D. paulistorum*, where *w*Pau *Wolbachia* are obligate mutualists affecting host behavior (Miller et al. 2010 and this study). Oenocytes derived from the *Wolbachia*-uninfected *D. willistoni* strain Willi3, a sister species of *D. paulistorum*, were used as negative controls (Fig. 5f). These data suggest that the oenocyte-tropism of *w*Pau is a phenotypic specificity for the *Wolbachia-D. paulistorum* system.

### Discussion

## Wolbachia-knockdown provokes a de novo assortative mating phenotype

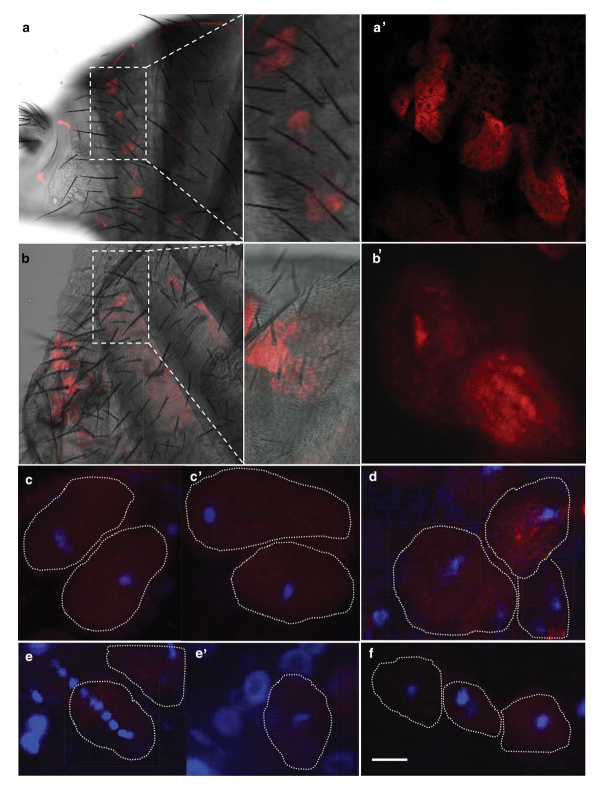
D. paulistorum semispecies are very sensitive to standard antibiotic treatments (Ehrman 1968; Kernaghan and Ehrman 1970) since in this symbiosis, *Wolbachia* is a fixed obligate mutualist providing vital, but still undetermined functions for its native hosts (Miller et al. 2010). Similar *Wolbachia*-dependencies were found in the parasitoid wasp *Asobara tabida* (Dedeine et al. 2001; Pannebakker et al. 2007), bedbugs of the genus *Cimex* (Hosokawa et al. 2010; Nikoh et al.

2014; Moriyama et al. 2015), and in the rice water weevil *Lissorhoptrus oryzophilus* (Chen et al. 2012). Furthermore, even partial depletion of fixed mutualistic *Wolbachia* from filarial nematodes affects host fitness and fecundity but also triggers overexpression of host nuclear and mitochondrial genes in order to partially compensate the loss of some gene functions provided by *Wolbachia* (Pfarr et al. 2008; Strübing et al. 2010).

Here we have assayed for behavioral consequences of symbiont knockdowns in males of two different D. paulistorum semispecies, i.e., Wolbachia low-titer Amazonian (A28) and high titer Orinocan (O11) strains. Our data strongly suggest that in both systems partial knockdown of the endosymbiont in males significantly affects their attractiveness for homogamic wt-females, which seems to correlate with global titer levels. However, restoration to random mating in later generations post treatment (Fig. 3a, b) seems to correlate with the gradual quantitative reconstitution of global symbiont titers that are fast in O11 (Fig. 2b), but not in A28 (Fig. 2a). In our assays, A28 flies expressing random mating at F<sub>13</sub> still show significantly lower global Wolbachia levels than wt flies. We speculate that reversion to random mating of A28 flies observed at later generations post-treatment might not depend on the global restoration of the symbiont



Behavior Genetics (2019) 49:83-98



**Fig. 5** Fluorescence *in situ* hybridization (FISH) on *Drosophila* oenocytes. (**a, a**') *Wolbachia* are present in *D. paulistorum* O11<sup>wt</sup> females, and (**b, b**') in O11<sup>wt</sup> males. **c** Symbionts are lacking from oenocytes of O11  $F_4^{kd}$  males and from  $F_4^{kd}$  females (**c**'). **d** In  $F_{13}$  post kd, oenocytes seem recolonized by *Wolbachia*. (**e, e**') Oenocytes from *Wolbachia*-infected *D. melanogaster* Harwich strain (H2) tested also

negative. **f** *Wolbachia*-uninfected *D. willistoni* (Willi3) was used as negative control. *Wolbachia* are shown in red (*16-23S* rRNA-specific probe); *Drosophila* nuclei are stained in blue (DAPI). Grey dashed lines mark borders of oenocyte cells. *Wol Wolbachia, kd* knockdown. Scale bar is 10 µm. (Color figure online)



titer in this semispecies to wt, but possibly because of a critical titer threshold and/or the symbiont's tropism to sensitive tissues and organs associated with mating behavior. In order to test this hypothesis, however, detailed temporal and spatial *in situ* quantifications of the symbiont in different host tissues during the restoration phase are required, which were beyond the scope of our current study.

As shown by this and earlier studies, under laboratory conditions different D. paulistorum semispecies express strong assortative mating behavior against each other, which is sensitive to rifampicin and tetracycline (Miller et al. 2010), but insensitive to penicillin and streptomycin that do not target the Wolbachia endosymbiont (this study). It seems feasible although not tested that strict female mate choice might prevent mating between incompatible members of different D. paulistorum semispecies also in nature. If true, this behavior has most likely coevolved in conjunction with the obligate endosymbiont to avoid detrimental reproductive phenotypes like hybrid mortality and male sterility in crosses between inter-semispecies (Ehrman 1968; Kim and Ehrman 1998). Here we have evaluated the potential of Wolbachia to trigger assortative mating behavior in D. paulistorum towards per se compatible mates under laboratory conditions. Our results suggest that kd-males appear sexually less attractive for their wt-female counterparts in mate choice assays upon manipulation of the native wPau-Wolbachia titer since they were expressing high SIIs in all assays. Such high SIIs were previously found only between members of different *D. paulistorum* semispecies (Ehrman 1965; Malogolowkin-Cohen et al. 1965) or between *Dros*ophila species that are sexually isolated from each other (reviewed in Spieth and Ringo 1983; Martin and Hosken 2003; Anderson and Kim 2005; Castrezana and Markow 2008). Our data also indicate that de novo sexual isolation in D. paulistorum under our experimental conditions is transient as eSIIs reverted to random mating around generation F<sub>13</sub> post kd (only 45–57% homogamy) together with the observed re-colonization of male oenocytes. However, experimental reduction of native D. paulistorum-Wolbachia titers was sufficient to alter males so that they were subsequently rejected by wt-females in the two semispecies tested, at least transiently. Interestingly we observed significant assortative mating between independently established kd-isofemale lines of the same semispecies, a finding most likely caused by drift. In such, heterogeneity of kd isolines could be a result of different temporal and spatial dynamics of Wolbachia recolonization of behavioral important tissues in the individual lines post knockdown. To test this drift hypothesis for kd lines, however, detailed quantitative and qualitative analyses of global symbiont titer levels, but also their in situ tropism and densities will be necessary from multiple staged flies and tissues of the same generation. In our current study, extensive analyses like these were not possible due to limitations of fly material at this time point.

### Host pheromonal signatures may be a target for *Wolbachia* to signal the infection state and to impact mate choice

In many insects, sexual pheromones serve as recognition cues for mate choice between and within species and are hence important players in reproductive isolation (Coyne et al. 1994; Savarit et al. 1998; Ferveur 2005). Moreover, such cues might even be prone to manipulation by pathogenic bacteria, as recently shown in *D. melanogaster* (Keesey et al. 2017).

In our study, the observed alteration of pheromone blends in kd-males may explain why females prefer wt-mates carrying the intact native profile. This finding is corroborated by studies showing that in lekking sandflies (*Lutzomya*), female mate choice is influenced by the amount of pheromones released by males before potential mating (Jones and Hamilton 1998; Jones et al. 2000). Another study demonstrated the influence of Wolbachia on host odor-linked mate preference in the terrestrial isopod Armadillidium vulgare (Richard 2017). In A28<sup>kd</sup> and O11<sup>kd</sup> males, relative quantities of pheromone compounds were affected differently, suggesting that the quantitative change in at least one of the male-specific components, but possibly also in conjunction with some other non-sex-specific CHCs, might be sufficient to trigger rejection of kd-males by wt-females in choice experiments. Importantly, our findings are in line with a recent study suggesting that Wolbachia can influence pupal communication between females and males in *Drosophila melanogaster* by modulating CI levels (Pontier and Schweissguth 2015, 2017; but also see; Jacquet et al. 2017). However, as shown earlier, facultative Wolbachia symbionts of D. melanogaster adults do not affect mate choice behavior in this system at all, but possibly other gut microbes (Sharon et al. 2010; Ringo et al. 2011; Arbuthnott et al. 2016). Importantly, a very recent study has severely questioned the potential impact of the gut microbiome on reproductive isolation since the microbiome of D. melanogaster is not fixed but reported as flexible and environmental determined (Leftwich et al. 2017). In the mutualistic D. paulistorum-Wolbachia system, however, where the endosymbiont is fixed by serving vital but still undetermined functions to their native host (Miller et al. 2010) we suggest that these obligate Wolbachia have also a direct or indirect impact on adult cuticular hydrocarbon (CHC) profiles, where male-specific and sex-unspecific compounds change quantitatively upon Wolbachia-kd in adult males.

The presence of *Wolbachia* in oenocytes of O11<sup>wt</sup> males implies a direct or indirect interaction of the symbiont with



host pheromone production and mate choice behavior as a consequence. The potential effects of the endosymbiont in oenocytes on female pheromone blending, however, await elucidation. If the presence of Wolbachia in male oenocytes is essential to express the semispecies-specific pheromone profile, loss of the symbiont from male oenocytes ( $F_4$  post kd) might explain why pheromone profiles are altered and knockdown males are consequently rejected as mate partners from homogamic wt-females. Based on our mate choice assays, the de novo assortative mating phenotype behavior is transient and results in reversion to a wt-like situation around F<sub>13</sub>/F<sub>14</sub> post knockdown. Assuming a direct link between Wolbachia tropism in the oenocytes and expression of the male pheromone profile, we expected a recolonization of oenocytes in parallel with reversion to random mating phenotype. We could confirm such recolonization, and thus potentially explain reversion to random mating, in oenocytes of F<sub>13</sub> O11<sup>kd</sup> males. Our finding is particularly interesting in the light of the contrasting situation in *D. melanogaster*, where Wolbachia do not play a role in mate choice (Sharon et al. 2010; Ringo et al. 2011; Arbuthnott et al. 2016). In our experimental setup, we did not detect Wolbachia in D. melanogaster oenocytes, which most likely explains why the symbiont does not affect mate choice and pheromone expression in this model system (Sharon et al. 2010). Finally, in contrast with the strict neurotropism of obligate Wolbachia to defined brain regions of D. paulistorum, (Strunov et al. 2017) native Wolbachia are randomly dispersed at low densities in *D. melanogaster* brains (Albertson et al. 2009). However, it remains to be elucidated how Wolbachia manipulate male pheromone expression and female mate choice in D. paulistorum.

To conclude, our combined data strongly imply that artificial reduction of the obligate Wolbachia endosymbiont of D. paulistorum males significantly reduces mating success with homogamic wildtype females belonging to the same semispecies. Although these and earlier findings in D. paulistorum lead to interpretations based on associations rather than causation and their functional and molecular bases are still undetermined, we propose the following model for the potential impact of the D. paulistorum endosymbiont on RI and host speciation in this system. In contrast to most facultative Wolbachia-insect associations, where the symbiont is not fixed and does not serve vital host functions, D. paulistorum semispecies have evolved vital mutualistic associations with their endosymbiont (Miller et al. 2010) as well as strict tissue tropisms to reproductive host organs, such as the primordial germline cells of embryos and adult gonads (Miller et al. 2010), but also to defined larval and adult brain regions associated with sexual behavior (Strunov et al. 2017) and pheromone producing oenocytes (this study). As implicated from the results of this study even partial depletions of the mutualist from their primary somatic host targets, such as oenocytes, might directly or indirectly alter pheromone signatures of males, which are no longer accepted as mates from stress-free wt females.

It appears likely that spontaneous symbiont knockdowns might also happen in the wild by stochastic exposure of the host to natural antibiotics or heat stress. This de novo assortative phenotype expressed by females against aberrant homogamic males can significantly disrupt gene flow within populations via premating isolation, at least transiently. As shown earlier, however, Wolbachia-kd males are accepted by kd females, which randomly mate even with heterogamic males under lab conditions (Miller et al. 2010). Under this scenario two sexually isolated reproductive groups would emerge and coexist next to each other. Since population sizes of Wolbachia have dropped significantly upon such external stresses, it seems quite likely that genetic drift (Chrostek et al. 2013; Schneider et al. 2013; Newton et al. 2014) can cause disruptive diversification of earlier-compatible Wolbachia variants that consequently trigger high cytoplasmic incompatibilities plus complete male sterility at postmating levels.

Hence it seems plausible that in Dobzhansky's earlier studies (Dobzhansky and Pavlovsky 1966, 1971, 1975) where spontaneous emergence of strong post mating incompatibilities between long term isolated sub lines of the same *D. paulistorum* semispecies were observed, he had actually detected the final outcome of this symbiotic stress and drift effect, observed in our current experimental study. Planned studies, however, should elucidate the mechanistic basis of *Wolbachia*-influence on male pheromone production and whether similar scenarios observed under lab conditions also take place in nature.

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#### Compliance with ethical standards

Conflict of interest Daniela I. Schneider, Lee Ehrman, Tobias Engl, Martin Kaltenpoth, Aurélie Hua-Van, Arnaud Le Rouzic and Wolfgang J. Miller declare that they have no conflict of interest.

**Ethical approval** For this type of study formal consent is not required. This article does not contain any studies with human participants or animals performed by any of the authors.



Informed consent Not applicable.

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#### References

- Aitchinson J (1986) The statistical analysis of compositional data. Monographs on statistics and applied probability. Chapman and Hall Ltd., London
- Albertson R, Casper-Lindley C, Cao J, Tram U, Sullivan W (2009) Symmetric and asymmetric mitotic segregation patterns influence Wolbachia distribution in host somatic tissue. J Cell Sci 122:4570–4583
- Anderson WW, Kim YK (2005) Sexual isolation between sympatric and allopatric populations of *Drosophila pseudoobscura* and *D. persimilis*. Behav Genet 35:305–312
- Arbuthnott D, Levin TC, Promislow DE (2016) The impacts of *Wolbachia* and the microbiome on mate choice in *Drosophila* melanogaster. J Evol Biol 29:461–468
- Audsley MD, Ye YH, McGraw EA (2017) The microbiome composition of *Aedes aegypti* is not critical for *Wolbachia*-mediated inhibition of dengue virus. PLoS Negl Trop Dis 11(3):e0005426
- Blomquist GJ, Bagnères AG (2010) Insect hydrocarbons: biology, biochemistry, and chemical ecology. Cambridge University Press, Cambridge
- Bordenstein SR, O'Hara FP, Werren JH (2001) *Wolbachia*-induced incompatibility precedes other hybrid incompatibilities in *Nasonia*. Nature 409:707–710
- Brownlie JC, Cass BN, Riegler M (2009) Evidence for metabolic provisioning by a common invertebrate endosymbiont, *Wolbachia pipientis*, during periods of nutritional stress. *PLoS Pathog*. (4):e1000368
- Brucker RM, Bordenstein SR (2012) Speciation by symbiosis. Trends Ecol Evol 27:443–451
- Brucker RM, Bordenstein SR (2013) The hologenomic basis of speciation: gut bacteria cause hybrid lethality in the genus *Nasonia*. Science 341:667–669
- Brucker RM, Bordenstein SR (2014) Response to Comment on "The hologenomic basis of speciation: gut bacteria cause hybrid lethality in the genus *Nasonia*". Science 345:1011
- Burla H, Brito da Cunha A, Cordeiro AR, Dobzhansky T, Malogolowkin C, Pavan C (1949) The *willistoni* group of sibling species of *Drosophila*. Evolution 3:300–314
- Castrezana SJ, Markow TA (2008) Sexual isolation and mating propensity among allopatric *Drosophila mettleri* populations. Behav Genet 38:437–445
- Chandler JA, Turelli M (2014) Comment on "The hologenomic basis of speciation: gut bacteria cause hybrid lethality in the genus *Nasonia*". Science 345:1011
- Chao TH, Ehrman L, Permaul A, Vincent R, Sattaur L, Brandt D (2010) Male-specific cuticular compounds of the six *Drosophila* paulistorum semispecies: structural identification and mating effect. J Chem Ecol 36:933–942
- Chen SJ, Lu F, Cheng JA, Jiang MX, Way MO (2012) Identification and biological role of the endosymbionts *Wolbachia* in rice water weevil (*Coleoptera: Curculionidae*). Environ Entomol 41:469–477

- Chrostek E, Marialva MS, Esteves SS, Weinert LA, Martinez J, Jiggins F, Teixeira M, L (2013) *Wolbachia* variants induce differential protection to viruses in *Drosophila melanogaster*: a phenotypic and phylogenomic analysis. PLoS Genet 9(12):e1003896
- Chung H, Carroll SB (2015) Wax, sex and the origin of species: dual roles of insect cuticular hydrocarbons in adaptation and mating. Bioessays 37:822–830
- Cooper BS, Ginsberg PS, Turelli M, Matute DR (2017) *Wolbachia* in the Drosophila yakuba complex: pervasive frequency variation and weak cytoplasmic incompatibility, but no apparent effect on reproductive isolation. Genetics 205:333–351
- Coyne JA, Orr HA (2004) Speciation. Sinauer Associates, Sunderland
- Coyne JA, Crittenden AP, Mah K (1994) Genetics of a pheromonal difference contributing to reproductive isolation in *Drosophila*. Science 265:1461–1464
- Dedeine F, Vavre F, Fleury F, Loppin B, Hochberg ME, Bouletreau M (2001) Removing symbiotic *Wolbachia* bacteria specifically inhibits oogenesis in a parasitic wasp. Proc Natl Acad Sci USA 98:6247–6252
- Dembeck LM, Böröczky K, Huang W, Schal C, Anholt RR, Mackay TF (2015) Genetic architecture of natural variation in cuticular hydrocarbon composition in *Drosophila melanogaster*. Elife. https://doi.org/10.7554/eLife.09861
- Dobzhansky T, Pavlovsky O (1966) Spontaneous origin of an incipient species in the *Drosophila paulistorum* complex. Proc Natl Acad Sci USA 55:727–733
- Dobzhansky T, Pavlovsky O (1971) Experimentally created incipient species of *Drosophila*. Nature 230:289–292
- Dobzhansky T, Pavlovsky O (1975) Unstable intermediates between Orinocan and Interior semispecies of *Drosophila paulistorum*. Evolution 29:242–248
- Dobzhansky T, Spassky B (1959) Drosophila paulistorum, a cluster of species in statu nascendi. Proc Natl Acad Sci USA 45:419–428
- Dodd DMB (1989) Reproductive isolation as a consequence of adaptive divergence in *Drosophila pseudoobscura*. Evolution 43:1308–1311
- Ehrman L (1965) Direct observation of sexual isolation between allopatric and between sympatric strains of the different *Drosophila paulistorum* races. Evolution 19:459–464
- Ehrman L (1968) Antibiotics and infections hybrid sterility in *Drosophila paulistorum*. Mol Gen Genet 103:218–222
- Ehrman L, Parsons PA (1981) Behavior genetics and evolution. McGraw-Hill, New York
- Ehrman L, Powell JR (1982) The *Drosophila willistoni* species group.
  In: Ashburner M, Carson HL, Thompson JN (eds) The genetics and biology of *Drosophila*. Academic Press, New York, pp 192–225
- Fast EM, Toomey ME, Panaram K, Desjardins D, Kolaczyk ED, Frydman HM (2011) Wolbachia enhance Drosophila stem cell proliferation and target the germline stem cell niche. Science 334:990–992
- Ferveur JF (2005) Cuticular hydrocarbons: their evolution and roles in *Drosophila* pheromonal communication. Behav Genet 35:279–295
- Gebiola M, Kelly SE, Hammerstein P, Giorgini M, Hunter MS (2016) "Darwin's corollary" and cytoplasmic incompatibility induced by *Cardinium* may contribute to speciation in Encarsia wasps (Hymenoptera: Aphelinidae). Evolution 70:2447–2458
- Hosokawa T, Koga R, Kikuchi Y, Meng XY, Fukatsu T (2010) Wolbachia as a bacteriocyte-associated nutritional mutualist. Proc Natl Acad Sci USA 107:769–774
- Jacquet A, Horard B, Loppin B (2017) Does pupal communication influence Wolbachia-mediated cytoplasmic incompatibility? Curr Biol 27:R55–R56



- Jaenike J, Dyer KA, Cornish C, Minhas MS (2006) Asymmetrical reinforcement and Wolbachia infection in Drosophila. Plos Biol 4(10):e325
- Jones TM, Hamilton JGC (1998) A role for pheromones in mate choice in a lekking sandfly. Animal Behav 56:891–898
- Jones TM, Balmford A, Quinnell RJ (2000) Adaptive female choice for middle-aged mates in a lekking sandfly. Proc R Soc B 267:681-686
- Keesey IW, Koerte S, Khallaf MA, Retzke T, Guillou A, Grosse-Wilde E, Buchon N, Knaden M, Hansson BS (2017) Pathogenic bacteria enhance dispersal through alteration of *Drosophila* social communication. Nat Commun 8:265
- Kernaghan RP, Ehrman L (1970) Antimycoplasmal antibiotics and hybrid sterility in *Drosophila paulistorum*. Science 169:63–64
- Kim YK, Ehrman L (1998) Developmental isolation and subsequent adult behavior of *Drosophila paulistorum*. IV. Courtship. Behav Genet 28:57–65
- Kim YK, Phillips DR, Chao T, Ehrman L (2004) Developmental isolation and subsequent adult behavior of *Drosophila paulistorum*. VI. Quantitative variation in cuticular hydrocarbons. Behav Genet 34:385–394
- Leclerq S, Mian FM, Stanisz AM, Bindels LB, Cambier E, Ben-Amram H, Koren O, Forsythe P, Bienenstock J (2017) Low-dose penicillin in early life induces long-term changes in murine gut microbiota, brain cytokines and behavior. Nat Commun 8:15062
- Leftwich PT, Clarke NVE, Hutchings MI, Chapman T (2017) Gut microbiomes and reproductive isolation in *Drosophila*. Proc Natl Acad Sci USA 114:12767–12772
- Leonard J, Ehrman L (1983) Does the rare male advantage result from faulty experimental design? Genetics 104:713–716
- Malogolowkin-Cohen C, Simmons AS, Levene H (1965) A study of sexual isolation between certain strains of *Drosophila paulisto*rum. Evolution 19:95–103
- Martin OY, Hosken DJ (2003) The evolution of reproductive isolation through sexual conflict. Nature 423:979–982
- Martinez J, Ok S, Smith S, Snoeck K, Day JP, Jiggins FM (2015) Should symbionts be nice or selfish? Antiviral effects of Wol-bachia are costly but reproductive parasitism is not. PLoS Pathog 11(7):e1005021
- Miller WJ, Riegler M (2006) Evolutionary dynamics of wAu-like Wolbachia variants in Neotropical Drosophila species. Appl Environ Microbiol 72:826–835
- Miller WJ, Schneider D (2012) Endosymbiotic microbes as adaptive manipulators of arthropod behaviour and natural driving sources of host speciation. In: Hughes D, Brodeur J J, and Thomas F (eds) Host manipulation by parasites. University Press, Oxford, pp 119–139
- Miller WJ, Ehrman L, Schneider D (2010) Infectious speciation revisited: impact of symbiont-depletion on female fitness and mating behavior of *Drosophila paulistorum*. PLoS Pathog 6(12):e1001214
- Moriyama M, Nikoh N, Hosokawa T, Fukatsu T (2015) Riboflavin provisioning underlies *Wolbachia*'s fitness contribution to its insect host. mBio 6(6):e01732–e01715
- Mouton L, Henri H, Bouletreau M, Vavre F (2006) Effect of temperature on Wolbachia density and impact on cytoplasmic incompatibility. Parasitology 132(Pt):49–56
- Najarro MA, Sumethasorn M, Lamoureux A, Turner TL (2015) Choosing mates based on the diet of your ancestors: replication of nongenetic assortative mating in *Drosophila melanogaster*. PeerJ 3:e1173
- Newton IL, Sheehan, and K. B (2014) Passage of *Wolbachia pipientis* through mutant *Drosophila melanogaster* induces phenotypic and genomic changes. Appl Environ Microbiol 81:1032–1037

- Nikoh N, Hosokawa T, Moriyama M, Oshima K, Hattori M, Fukatsu T (2014) Evolutionary origin of insect-*Wolbachia* nutritional mutualism. Proc Natl Acad Sci USA 111:10257–10262
- Pannebakker BA, Loppin B, Elemans CP, Humblot L, Vavre F (2007)
  Parasitic inhibition of cell death facilitates symbiosis. Proc Natl
  Acad Sci USA 104:213–215
- Pfarr KM, Heider U, Schmetz C, Buttner DW, Hoerauf A (2008) The mitochondrial heat shock protein 60 (HSP 60) is up-regulated in *Onchocerca volvulus* after the depletion of *Wolbachia*. Parasitology 135:529–538
- Poinsot D, Montchamp-Moreau C, Merçot H (2000) Wolbachia segregation rate in *Drosophila simulans* naturally bi-infected cytoplasmic lineages. Heredity 85:191–198
- Pontier SM, Schweissguth F (2015) A *Wolbachia*-sensitive communication between male and female pupae controls gamete compatibility in *Drosophila*. Curr Biol 25:2339–2348
- Pontier SM, Schweissguth F (2017) Response to "Does pupal communication influence Wolbachia-mediated cytoplasmic incompatibility? Curr Biol 27(2):R55–R56
- Reynolds KT, Hoffmann AA (2002) Male age, host effects and the weak expression or non-expression of cytoplasmic incompatibility in *Drosophila* strains infected by maternally transmitted *Wolbachia*. Genet Res 80:79–87
- Richard F-J (2017) Symbiotic bacteria influence the odor and mating preference of their hosts. Front Ecol Evol. https://doi.org/10.3389/ fevo.2017.00143
- Ringo J, Sharon G, Segal D (2011) Bacteria-induced sexual isolation in *Drosophila*. Fly 5:310–315
- Savarit F, Sureau G, Cobb M, Ferveur JF (1998) Genetic elimination of known pheromones reveals the fundamental chemical bases of mating and isolation in *Drosophila*. Proc Natl Acad Sci USA 96:9015–9020
- Schneider DI, Riegler M, Arthofer W, Merçot H, Stauffer C, Miller WJ (2013) Uncovering *Wolbachia* diversity upon artificial host transfer. PLoS ONE 8(12):e82402
- Schneider DI, Parker AG, Abd-Alla AM, Miller WJ (2018) High-sensitivity detection of cryptic *Wolbachia* in the African tsetse fly (*Glossina* spp.). BMC Microbiol, in press
- Sharon G, Segal D, Ringo JM, Hefetz A, Zilber-Rosenberg I, Rosenberg E (2010) Commensal bacteria play a role in mating preference of *Drosophila melanogaster*. Proc Natl Acad Sci USA 107:20051–20056
- Spieth HT, Ringo JM (1983) Mating behavior and sexual isolation in *Drosophila*. In: Ashburner M, Carson HL, Thompson JN (eds) The genetics and biology of *Drosophila*. Academic Press, London, pp 345–384
- Strübing U, Lucius R, Hoerauf A, Pfarr KM (2010) Mitochondrial genes for heme-dependent respiratory chain complexes are upregulated after depletion of *Wolbachia* from filarial nematodes. Int J Parasitol 40:1193–1202
- Strunov A, Schneider DI, Alberston R, Miller WJ (2017) Restricted distribution and lateralization of mutualistic *Wolbachia* in the *Drosophila* brain. Cell Microbiol. https://doi.org/10.1111/cmi.12639
- Teixeira L, Ferreira Å, Ashburner M (2008) The bacterial symbiont *Wolbachia* induces resistance to RNA viral infections in *Drosophila melanogaster* PLoS Biol 6:2753–2763
- Telschow A, Flor M, Kobayashi Y, Hammerstein P, Werren JH (2007) Wolbachia-induced unidirectional cytoplasmic incompatibility and speciation: mainland-island model. PLoS ONE 2(8):e701
- Telschow A, Hilgenboecker K, Hammerstein P, Werren JH (2014) Dobzhansky-Muller and *Wolbachia*-induced incompatibilities in a diploid genetic system. PLoS ONE 9(4):e95488
- Turelli M, Lipkowitz JR, Brandvain Y (2014) On the Coyne and Orrigin of species: effects of intrinsic postzygotic isolation, ecological differentiation, X chromosome size, and sympatry on *Drosophila* speciation. Evolution 68:1176–1187



98 Behavior Genetics (2019) 49:83–98

Ueyama M, Chertemps T, Labeur C, Wicker-Thomas C (2005) Mutations in the *desat1* gene reduces the production of courtship stimulatory pheromones through a marked effect on fatty acids in *Drosophila melanogaster*. Insect Biochem Mol Biol 35:911–920

- Weeks AR, Turelli M, Harcombe WR, Reynolds KT, Hoffmann AA (2007) From parasite to mutualist: rapid evolution of *Wolbachia* in natural populations of *Drosophila*. PLoS Biol 5(5):e114
- Werren JH, Baldo L, Clark ME (2008) *Wolbachia*: master manipulators of invertebrate biology. Nat Rev Microbiol 6:741–751
- Yamada R, Floate K, Riegler M, O'Neill S (2007) Male development time influences the strength of *Wolbachia*-induced cytoplasmic
- incompatibility expression in Drosophila melanogaster. Genetics 177:801–808
- Ye YH, Seleznev A, Flores HA, Woolfit M, McGraw EA (2017) Gut microbiota in *Drosophila melanogaster* interacts with *Wolbachia* but does not contribute to *Wolbachia*-mediated antiviral protection. J Invertebr Pathol 143:18–25
- Zug R, Hammerstein P (2012) Still a host of hosts for Wolbachia: analysis of recent data suggests that 40% of terrestrial arthropod species are infected. PLoS ONE 7(6): e38544

