

Decoupling of Host–Symbiont–Phage Coadaptations Following Transfer Between Insect Species

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ABSTRACT

Transferring endosymbiotic bacteria between different host species can perturb the coordinated regulation of the host and bacterial genomes. Here we use the most common maternally transmitted bacteria, *Wolbachia pipientis*, to test the consequences of host genetic background on infection densities and the processes underlying those changes in the parasitoid wasp genus *Nasonia*. Introgressing the genome of *Nasonia giraulti* into the infected cytoplasm of *N. vitripennis* causes a two-order-of-magnitude increase in bacterial loads in adults and a proliferation of the infection to somatic tissues. The host effect on *W. pipientis* distribution and densities is associated with a twofold decrease in densities of the temperate phage WO-B. Returning the bacteria from the new host species back to the resident host species restores the bacteria and phage to their native densities. To our knowledge, this is the first study to report a host–microbe genetic interaction that affects the densities of both *W. pipientis* and bacteriophage WO-B. The consequences of the increased bacterial density include a reduction in fecundity, an increase in levels of cytoplasmic incompatibility (CI), and unexpectedly, male-to-female transfer of the bacteria to uninfected females and an increased acceptance of densely infected females to interspecific mates. While paternal inheritance of the *W. pipientis* was not observed, the high incidence of male-to-female transfer in the introgressed background raises the possibility that paternal transmission could be more likely in hybrids where paternal leakage of other cytoplasmic elements is also known to occur. Taken together, these results establish a major change in *W. pipientis* densities and tissue tropism between closely related species and support a model in which phage WO, *Wolbachia*, and arthropods form a tripartite symbiotic association in which all three are integral to understanding the biology of this widespread endosymbiosis.

ALL metazoans are populated by symbiotic bacteria, some of which live in intimate association with their hosts as maternally transmitted infections. *Wolbachia pipientis* is one of the most prevalent species of bacterial endosymbionts in the animal world that exemplifies this lifestyle. It is maternally transmitted from host ovaries to developing eggs in filarial nematodes and arthropods. In filarial nematodes, *W. pipientis* are obligate mutualists (TAYLOR *et al.* 2005) and assist fertility (HOERAUF *et al.* 1999) and larval development (SMITH and RAJAN 2000). Proinflammatory responses in filarial-infected vertebrates indicate that the endosymbiont rather than the nematode is the significant

contributor to the acute pathologies of human river blindness (SAINT ANDRE *et al.* 2002) and elephantiasis (TAYLOR *et al.* 2000). In arthropods, *W. pipientis* are reproductive parasites (STOUTHAMER *et al.* 1999; WERREN *et al.* 2008) and, less often, mutualists (PANNEBAKKER *et al.* 2007; HOSOKAWA *et al.* 2010), and occur in ~66% of all insect species worldwide (HILGENBOECKER *et al.* 2008). Thus, millions of animal species are infected by *W. pipientis* and over 1 billion people in >80 countries are at risk of filarial/*W. pipientis* diseases (OTTESEN *et al.* 2008). As one of the great pandemics in the animal world, *W. pipientis* offer a preeminent model to explore the varied outcomes of infection between animal cells and obligate intracellular bacteria.

Transfer of symbionts between closely related species is a useful tool to determine the role of host genes in regulating a symbiosis. Over time, maternally inherited bacteria may coevolve with their hosts to finely tune their densities, tissue tropism, and interactions with host fitness. Many studies have found that relocating *W. pipientis* to a new host species causes changes in reproductive parasitism, bacterial titers, and tissue distri-

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bution not observed in the donor (BOYLE *et al.* 1993; BORDENSTEIN and WERREN 1998; MCGRAW *et al.* 2002; RIEGLER *et al.* 2004; RUANG-AREERATE and KITTAYAPONG 2006).

In addition to host genetic background, temperate phage WO of *W. pipientis* is hypothesized to regulate infection densities in *Nasonia vitripennis* wasps owing to its lifecycle as a lytic and lysogenic phage (BORDENSTEIN *et al.* 2006; KENT and BORDENSTEIN 2010). In that investigation, phage WO was observed in ~12% of the *W. pipientis* cells in late pupal testes. These bacteria showed cellular defects consistent with phage-induced mortality, including degraded DNA, a collapsed inner membrane typical of holin enzymes, and lysed membranes associated with phage. Introduction of a new host background could alter lytic development and result in changes in *W. pipientis* infection densities and tissue tropism from the donor species.

The *Nasonia* genus is composed of four closely related parasitoid wasp species that are naturally infected with different *W. pipientis* strains (RAYCHOUDHURY *et al.* 2009). *N. vitripennis* and *N. giraulti* laboratory strains and field isolates are coinfecting by each of the two major insect-*W. pipientis* subdivisions, A and B (WERREN *et al.* 1995; RAYCHOUDHURY *et al.* 2009). These infections have been segregated in the lab into strains with single A or B infections (PERROT-MINNOT *et al.* 1996; BORDENSTEIN and WERREN 2007). Crosses between the different A infections, and between the single A and B infections within species, are bidirectionally incompatible due to cytoplasmic incompatibility (CI) (BORDENSTEIN and WERREN 2007). In addition to *Drosophila*, the *Nasonia* genus is one of a few tractable genetic systems for understanding host-*W. pipientis* interactions because it has many features suited for genetic and functional studies. They have a short generation time, simple husbandry, fertile species hybridizations, a wide array of molecular markers (WERREN and LOEHLIN 2009), and three fully sequenced genomes (WERREN *et al.* 2010).

Here, we describe a comprehensive set of experiments to test the consequences of host genetic background on *W. pipientis* densities, the processes underlying the changes in density, and the effects of increased bacterial densities on cytoplasmic incompatibility, fitness, behavior, and paternal transmission. To accomplish this, we introgressed the *N. giraulti* genome into a *N. vitripennis* A-infected cytoplasm. Results showed a stable increase in *W. pipientis* density, a reduction in phage WO density, and a vast expansion of the infection into somatic tissues. The ~100-fold increase in *W. pipientis* in the new genetic background resulted in male-to-female transfer of *W. pipientis* and a behavioral mating pathology. Return of the bacteria/bacteriophage combination from the introgression line back to the resident *N. vitripennis* host restores the native microbial densities. The host genetic effect on microbial densities was not observed with the supergroup-B infected *N. vitripennis* strain that lacks a

lytic phage, suggesting the increase in bacterial densities and reduction in phage WO densities could be specific to the supergroup A strain that harbors the virion-producing phage.

MATERIALS AND METHODS

Strains: *N. vitripennis* strains 12.1 (A-infected), 4.9 (B-infected), and 13.2 (uninfected) were derived from the double AB-infected R511 line after a prolonged period of diapause (PERROT-MINNOT *et al.* 1996). Introgression strains for this study were derived by repeatedly backcrossing females of 12.1, 4.9, and 13.2 to uninfected *N. giraulti* RV2R males (uninfected) for nine, six, and six generations, respectively. These strains are essentially composed of *N. giraulti* nuclear genomes and *N. vitripennis* cytoplasm; introgressed strains are IntG12.1 with an A-infected cytoplasm, IntG4.9 with a B-infected cytoplasm, and IntG13.2 with an uninfected cytoplasm. Strains 12.1 and IntG12.1 were subsequently treated with 30 mg/ml tetracycline for three generations to generate *W. pipientis*-free versions of these lines and are denoted 12.1T and IntG12.1T. The IntG12.1 line was backcrossed to *N. vitripennis* 12.1 males for seven generations to reintroduce the *N. vitripennis* genome and is denoted ReInt12.1. All strains were incubated at 25° at constant light during the experiments. To set up crosses, *Nasonia* wasps were collected as black pupae, sorted into separate virgin male and female stocks, and mated upon eclosion as adults. Copulations were observed for all crosses.

Fluorescent imaging: Females were presented with one *Sarcophaga bullata* host pupa embedded halfway in foam plugs to localize egg laying in the host for 24 hr. Embryos to be used in immunofluorescent imaging were then collected and rinsed with heptane into a final 1:1 heptane-methanol mixture and then shaken for ~5 min. Embryos, which became dechorionated and fixed, settled to the bottom of the methanol layer and were removed with a pipette, transferred to a microcentrifuge tube, and washed three times with methanol for 5 min each. Fixed embryos were then hydrated with TBST (50 mM Tris, 150 mM NaCl, 0.1% Tween, 0.05% NaN₃, pH 7.5) through a series of 1:3, 1:1, 3:1 TBST:methanol mixture for 5 min each, followed by TBST and blocked in TBST-BSA (with 1% bovine serum albumin) for 10 min.

Seminal vesicles and malpighian tubules were dissected from 2- to 3-day posteclosion males in TBST and fixed in 3.7% formaldehyde in TBST for 30 min, followed by three 5-min washes in TBST, and blocked in TBST-BSA (with 1% bovine serum albumin) for 10 min.

W. pipientis were labeled using a mouse monoclonal anti-human hsp60 antibody (Sigma), which also labels *W. pipientis* (HOERAUF *et al.* 2000). Tissues were incubated with 1:250 primary antibody for 1 hr at room temperature in TBST-BSA and 2 mg/ml RNaseA followed by three 5-min washes in TBST. This was followed by 1 hr at room temperature in 1:500 Alexa Fluor 488 anti-mouse antibody (Invitrogen) followed by three 5-min washes in TBST. DNA was then stained with either 5 µg/ml propidium iodide (embryos) for 20 min, or DAPI (malpighian tubules) for 10 min followed by a brief wash in TBST before mounting in ProLong Gold antifade mounting media (Invitrogen). Confocal images of embryos were obtained using a Leica SP confocal microscope. Seminal vesicles and malpighian tubules were imaged with a Zeiss Axio-Imager Z1 microscope.

Fluorescent *in situ* hybridization (FISH) of 30-µm *Nasonia* cross-sections was employed to visually show differences in bacterial densities and tissue tropism in the high-density

(IntG12.1) and low-density (12.1) strains. Reproductive tissues were dissected within 1 hr after mating directly in 4% PFA on PLUS GOLD adhesion slides. All tissues were fixed for 1 hr and dehydrated using a graded ethanol series. Two *W. pipientis*-specific 16S oligonucleotide probes, W1 and W2 (HEDDI *et al.* 1999), labeled with Texas Red at the 5' end were used at 25 ng/ml. Hybridization was performed at 37° in a dark moisture chamber in 0.5 ml hybridization buffer (50% formamide, 5× SSC, 200 g/liter dextran sulfate, 250 mg/liter poly(A), 250 mg/liter salmon sperm DNA, 250 mg/liter tRNA, 0.1 M dithiothreitol (DTT), and 0.5× Denhardt's solution). After an overnight incubation, the slides were washed twice in 1× SSC–10 mM DTT and twice in 0.5× SSC–10 mM DTT at 55°. The slides were rinsed in deionized water, mounted with Vectashield antifade mounting medium containing 1.5 µg/ml DAPI (4',6'-diamidino-2-phenylindole) and viewed on a Zeiss axiovert compound microscope fitted with epifluorescent optics and photographed with a Cannon G10 fitted with a Zeiss Soligor adapter tube. Confocal images were taken using a Zeiss Axioplan 2 confocal microscope fitted with a HeNe1 laser (543-nm excitation) and DIC.

Quantitative analysis of *W. pipientis* and phage WO densities: Genomic DNA was extracted using the Puregene tissue kit (Qiagen, Valencia, CA) from single adult males and females. Real-time quantitative polymerase chain reaction (qPCR) was performed in either an iCycler system (Bio-Rad) or CFX96 Real-Time system (Bio-Rad). Reaction volumes of 25 µl contained 12.5 µl of BioRad SYBR Green Supermix, 10.5 µl sterile water, 0.5 µl of each 10 µM forward and reverse primer, and 2 µl target DNA in single wells of a 96-well plate (Bio-Rad). Selective amplification was performed on a small portion of the *Nasonia S6 Kinase (S6K, 133 bp)*, *W. pipientis groEL (groEL, 97 bp)*, and bacteriophage WO-B (*ORF7, 125 bp*) genes, as previously described (BORDENSTEIN *et al.* 2006). A melting curve analysis was performed following each PCR to check for primer dimers and nonspecific amplification. Standard curves for each gene were constructed with a log10 dilution series of PCR products cloned into plasmid vectors using the TOPO TA cloning kit (Invitrogen, Carlsbad, CA). qPCR assays were performed in triplicate on all DNA, and data analyses were performed on the average of the three replicates.

Fecundity and survival: The effect of *W. pipientis* on fecundity was examined with virgin females that were *W. pipientis* infected (IntG12.1), antibioticly treated (IntG12.1T), or uninfected (IntG13.2). Virgin females were provided three hosts in a 12 × 75-mm vial for 48 hr, at which time females were removed and transferred to a single host for 8 hr. Limited ovipositioning time prevents developing wasps from becoming resource limited (BORDENSTEIN and WERREN 2000). Adult offspring were scored from 80 females during two studies (October 2007 and June 2008) using uninfected, control (IntG13.2) females and antibioticly treated (IntG12.1T) females, respectively. In the haplodiploid *Nasonia* system, unmated females exclusively produce haploid males. Adult and diapause male F₁ offspring were included in offspring production per female.

The effect of *W. pipientis* on survival under constant environmental conditions without food was also evaluated with infected (IntG12.1) and uninfected, control (IntG13.2) males and females. Upon eclosure, individual insects were transferred to vials and held without food. Deaths were counted at 0, 24, 48, 65, 72, 89, 96, and 120 hr.

Mate discrimination: The role of *W. pipientis* in host mate discrimination was assessed by scoring the percentage of successful single-pair copulations observed within a 10-min time span in glass vials. No choice interspecific tests were set up between infected tester *N. vitripennis* males (12.1) and

infected (IntG12.1), antibioticly treated (IntG12.1T) and control, uninfected (IntG13.2) females harboring the *N. giraulti* genome. Behavioral observations were conducted with 1-day-old individuals. In total, 157 males were mated with uninfected females and 176 were mated with infected females over two replicate experiments in September 2007 and May 2008.

Bidirectional cytoplasmic incompatibility: Single-pair crosses between introgression and control strains were set up to determine whether gender differences in bacterial loads cause bidirectional CI. Following single-pair copulations, females were hosted on three hosts for 48 hr and transferred to one host for eight hr for egg laying. CI was scored on the basis of the number of adult F₁ male and female offspring per female.

Additionally, low-density (12.1) males used in the qPCR experiments were also employed in a study of unidirectional CI to correlate *W. pipientis* and phage gene copy numbers to levels of CI. Following single-pair copulations with these males, 12.1T tetracycline-cured females were hosted on three hosts for 48 hr and transferred to one host for 8 hr for egg laying. Adult offspring were scored upon death for number of female offspring as a proxy for unidirectional CI.

Male-to-female transfer: Females were collected immediately after mating to determine the frequency of *W. pipientis* transmission from an infected male to an uninfected female on the basis of polymerase chain reaction (PCR). DNA from uninfected females was extracted following the methods described below and assessed for transmission of *W. pipientis* from infected males at time point 0 and/or every 24 hr after mating for 5 days. Primers and PCR conditions are also described below.

Inheritance of paternally transmitted *W. pipientis* to offspring was tested in two replicate experiments that were carried out in September 2009 (*N* = 48 families) and October 2009 (*N* = 40 families). Upon mating with infected males, uninfected mothers in replicate 1 were hosted on two hosts in glass vials for 4 days and then rehosted on two fresh hosts for an additional 5 days in new vials. Females were collected after 9 days and batches of 10 adult offspring from each hosting were collected and frozen at –80° upon emergence. In experimental replicate 2, females were hosted on two hosts for 48 hr and batches of 10 offspring from four life stages (eggs, larvae, yellow-red pupae, and adults) were collected and frozen at –80°. DNA from mothers and offspring was later isolated and PCR tested for *W. pipientis* transmission. DNA was extracted using a Genra Puregene tissue kit (Qiagen). PCR was conducted with primers from the *W. pipientis*-specific 16S rRNA gene (Lo *et al.* 2002) and bacteriophage WO-specific primers from the gp17 gene from WOCauB1 (FUJII *et al.* 2004): gp17F (5'CCAGATCAATTAGCATATCTTGCT) and gp17R2 (5'TCTTGGCCGATAAAAACTAAC). PCR conditions for gp17 are as follows: 95° for 2 min, followed by 35 cycles of 94° for 1 min, 53° for 1 min, 72° for 1 min 30 sec, and 72° for 5 min.

Preliminary PCR data indicated that high-density, introgressed males (IntG12.1) transmit *W. pipientis* to uninfected females immediately after mating but the infection disappears in the mated female after 24 hr. A comparison study with deceased females was then set up to test the female's role in eliminating *W. pipientis* over time. Female heads were decapitated immediately after mating and the remaining body was held in glass vials. Decapitated females were collected at 0 hr and every 24 hr after mating for 20 days and frozen at –80°. PCR was then used to determine whether the *W. pipientis* signal disappears in dead females at the same rate as it did in live females.

Statistics: Pairwise comparisons of adult family size were conducted with the nonparametric Mann–Whitney *U*-tests (MWU) at $\alpha = 0.05$ (MiniTab v.12.23, State College, PA), as

some distributions were not normal. Fisher's exact test at $\alpha = 0.05$ was calculated (JMP v. 5.0, SAS Institute) to determine significant deviations from the null hypothesis in discrete datasets. Correlation coefficients were calculated using the nonparametric Spearman's ρ (JMP v. 5.0). A Mantel Cox log-rank test was used to compare survival distributions.

RESULTS

Host variation in infection of the somatic tissues:

Genome replacement by genetic introgression of the *N. giraulti* genome into a *N. vitripennis* A-infected cytoplasm for nine generations in April 2006 led to a stable and heritable increase in *W. pipientis* densities in the introgression line (IntG12.1) compared to the resident A-infected *N. vitripennis* line (12.1). Extensive proliferation of the infection in *N. giraulti* IntG12.1 was evident by the titer increase at the posterior end of the embryo and the spread of the bacteria toward the anterior end (Figure 1A). The proliferation of the infection throughout the embryo in the *N. giraulti* IntG12.1 line placed *W. pipientis* infections in cells fated to become somatic cells. Consequently, bacterial proliferation into adult tissues not typically infected by the resident *N. vitripennis* 12.1 infection occurred in all tissues of the *N. giraulti* IntG12.1 line examined, including malpighian tubules (Figure 1B), accessory glands, and seminal vesicles (Figure 1C).

Whole female *Nasonia* tissue sections hybridized with *W. pipientis*-specific 16S rRNA fluorescent probes W1 and W2 (HEDDI *et al.* 1999) showed somatic tissue infection throughout the entire *N. giraulti* IntG12.1 wasp (Figure 2C). Strong *W. pipientis* signals were also observed in the abdomen, flight muscle tissues in the thorax, and throughout the head (Figure 2, C and D). In contrast, the resident host *N. vitripennis* 12.1 showed localization only to the abdomen (Figure 2B), and the control unstained *N. vitripennis* 12.1 yielded little autofluorescence of the probes (Figure 2A).

In the male gonads of the new *N. giraulti* IntG12.1 genetic background, fluorescent imaging of the W1 and W2 probes showed that the bacteria proliferate in the testes, seminal vesicle, and accessory gland (Figure 3A). One notable observation is an unusual signal aggregation or "clumping" of *W. pipientis* at the base of the testes (Figure 3A, white arrows), suggesting *W. pipientis* is shed from the sperm by the individualization complexes during spermatogenesis, possibly into waste bags, at high densities (CLARK and KARR 2002; RIPARBELLI *et al.* 2007). There is a dense fluorescence in the testes that is not present in the accessory gland and seminal vesicles. It could be due to an artifact of autofluorescence or an extremely dense infection in the testes. Control experiments with the testes from cured and low-density *N. vitripennis* 12.1 males did not show the dense signal in the testes (supporting information, Figure S1). The presence of *W. pipientis* in the seminal vesicle and accessory gland suggests it could be transmitted pater-

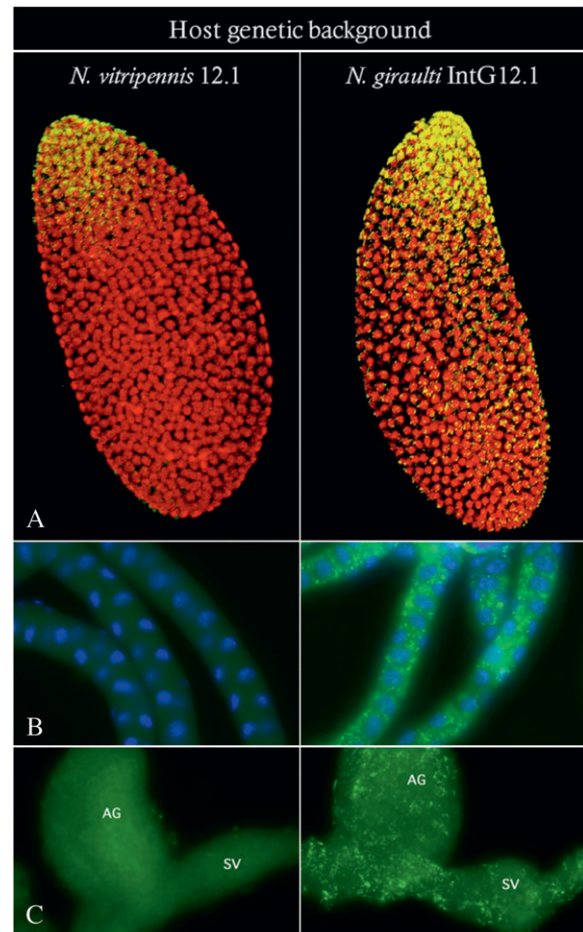


FIGURE 1.—Immunofluorescence staining of A-group *W. pipientis* in *N. vitripennis* and *N. giraulti*. Images show *W. pipientis* stained in yellow or green and host DNA stained in red or blue. Density differences and tissue tropism are shown in (A) embryos; posterior end is at the top, (B) malpighian tubules, and (C) ag, accessory glands and sv, seminal vesicles.

nally through the seminal fluid (see section on male-to-female transfer).

Host variation in microbial densities and expressivity of CI: To examine *W. pipientis* and phage WO-B titer differences between the two host species, we used qPCR of single-copy genes of the *N. vitripennis* and *N. giraulti* genomes (*S6K*), *W. pipientis* genomes (*groEL*), and phage WO-B (*ORF7*). Consistent with the fluorescent imaging, *W. pipientis* densities were significantly higher in *N. giraulti* IntG12.1 adults than in *N. vitripennis* 12.1 adults, for both males (MWU, $P < 0.0001$) and females (MWU, $P < 0.0001$) (Figure 4). The mean \pm standard error for *W. pipientis* density in *N. giraulti* IntG12.1 (males, 8.98 ± 0.82 ; females, 6.08 ± 0.16) was 128-fold and 36-fold higher than the average *W. pipientis* density in the resident *N. vitripennis* 12.1 (males, 0.07 ± 0.01 ; females, 0.17 ± 0.01). These increases were associated with significant decreases in phage WO-B densities (phage-to-bacteria ratios) for males and fe-

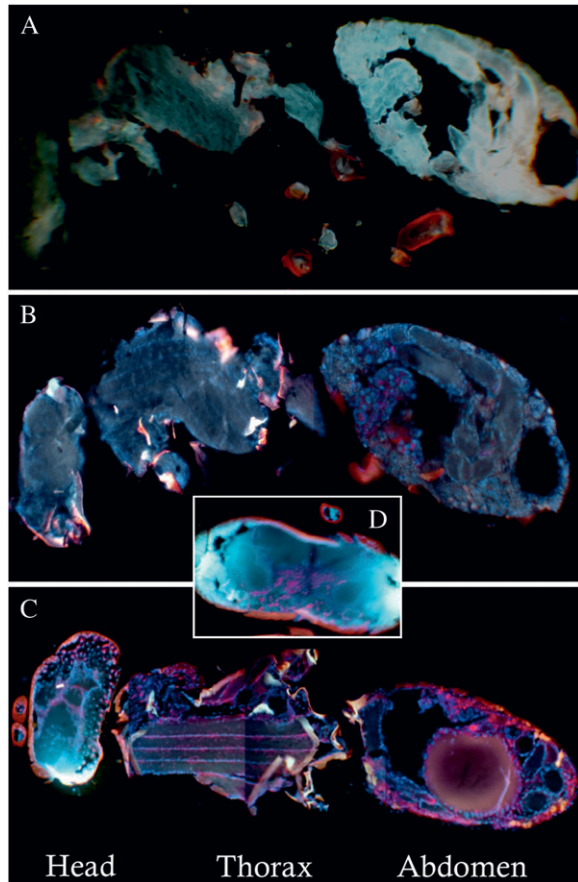


FIGURE 2.—Fluorescent *in situ* hybridizations of *W. pipientis* in adults of *N. vitripennis* and *N. giraulti*. Comparisons of 30- μ m cross-sections from adult females of (A) unstained *N. vitripennis* 12.1 (control for autofluorescence), (B) stained *N. vitripennis* 12.1, (C) stained *N. giraulti* IntG12.1, and (D) a stained *N. giraulti* IntG12.1 head. Strong *W. pipientis* signals (red) are observed in IntG12.1 in the head, the flight mussel tissues in the thorax, and the abdomen. Images B–D are counterstained with DAPI (blue) and are presented as composite images from the same individual to enhance resolution of the *W. pipientis*.

males, respectively (Figure 4). There were 2.0-fold and 1.7-fold reductions in phage WO-B densities in the densely infected *N. giraulti* IntG12.1 relative to *N. vitripennis* 12.1 for males (MWU, $P < 0.0001$) and females (MWU, $P < 0.0001$), respectively. Furthermore, there were significant inverse associations between phage and *W. pipientis* densities within species in *N. vitripennis* 12.1 males ($\rho = -0.6932$, $P < 0.0001$, Figure 4) and *N. giraulti* IntG12.1 males ($\rho = -0.8455$, $P < 0.0001$, Figure 4), but not females. For instance, the *N. vitripennis* male with the highest WO-B density (45.36 *ORF7* gene copies per *W. pipientis groEL* gene copies) had the lowest *W. pipientis* density, and the *N. vitripennis* male with the lowest WO-B density (9.30) had the highest *W. pipientis* density. Similarly for *N. giraulti*, the male with the highest WO-B density (21.43 *ORF7* gene copies per *W. pipientis groEL* gene copies) had the second lowest *W. pipientis* density, and the male with

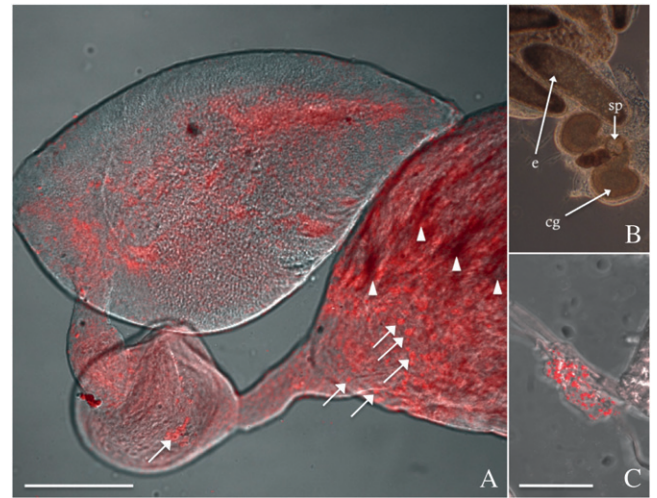


FIGURE 3.—Fluorescent *in situ* hybridizations of *W. pipientis* in reproductive tissues of *N. giraulti*. *W. pipientis* (red) is shown in (A) the testes of an infected IntG12.1 male and (C) the spermatheca of an uninfected female mated to a IntG12.1 male. The ovaries of an uninfected female are also shown in B, e, mature egg at the base of an ovariole; cg, colleterial glands; sp, spermatheca. Arrows indicate *W. pipientis* aggregation within the testes (A) and may indicate the cellular unit transmitted to the spermatheca of uninfected females (C). Bars, 50 μ m.

the lowest WO-B density (6.86) had the second highest *W. pipientis* density.

A replicate experiment of the host variation in *W. pipientis* densities was performed with additional *Nasonia* strains infected with different bacterial strains (Figure 5). First, the study recapitulated the major density differences described above. Female 12.1 and IntG12.1 infected with the supergroup A strain showed a 112-fold difference (MWU, $P = 0.02$). Second, the increased bacterial densities appear specific to the A infection as the introgression of the *N. giraulti* genome into the *N. vitripennis* cytoplasm carrying the supergroup B infection did not yield different bacterial loads (4.9 vs. IntG4.9, MWU, $P = 0.73$). The strain-specific increase was further evident by quantifications of the divergent A infection in its resident *N. giraulti* genome (strain 16.2), in which it exhibited low densities (16.2 vs. 12.1, MWU, $P = 0.86$). Finally, replacement of the IntG12.1 *N. giraulti* genome back to a *N. vitripennis* genome resulted in the return of low *W. pipientis* densities (ReInt12.1 vs. 12.1, 0.51 vs. 0.70; MWU, $P = 0.14$) and high phage densities (ReInt12.1 vs. 12.1, 6.1 vs. 5.6; MWU, $P = 1.0$), indicating the native bacterial and phage densities were restored in the resident *N. vitripennis* background (Figure 5).

Fecundity: To determine whether the extensive proliferation of *W. pipientis* densities and tissue tropism in the new *N. giraulti* host influences fitness, we compared the fecundity of IntG12.1 infected, virgin females (haploid male offspring only), IntG12.1T antibioticly cured virgin females, and IntG13.2 uninfected, control

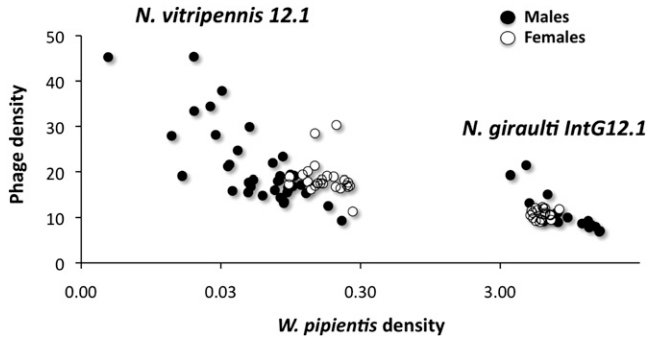


FIGURE 4.—Inverse relationships between *W. pipientis* and phage WO-B densities. Bacteriophage WO-B *vs.* *W. pipientis* gene copy number in the resident *N. vitripennis* and novel *N. giraulti* host genetic backgrounds. Points on the chart denote bacterial and bacteriophage densities in individual males and females.

virgin females that went through the same introgression scheme as the infected strain. A Mann–Whitney *U*-test revealed a marginally nonsignificant decrease ($P = 0.07$) in comparison to the IntG13.2 uninfected, control females and a significant decrease ($P = 0.0007$) in mean fecundity per infected female in comparison with IntG12.1T tetracycline-treated females (Figure 6). The difference in the significance of the two datasets was apparently due to the fecundity of infected females (10.83 *vs.* 6.03, MWU, $P = 0.0002$), as uninfected females yielded similar offspring numbers (12.93 *vs.* 11.36, MWU, $P = 0.17$).

Starvation resistance: Increases in *W. pipientis* densities and infection of somatic tissues in the *N. giraulti* genetic background may also affect the host's duration of life, similar to the rapidly proliferating *wMel* Popcorn strain in *Drosophila melanogaster* that causes early mortality (MIN and BENZER 1997). We measured survival of individual IntG12.1 infected and IntG13.2 uninfected males and females under constant environmental conditions without food (Figure 7). Survival of starved wasps was slightly extended in infected males and significantly extended in infected females (Mantel Cox log-rank test, $\chi^2 = 19.3$, d.f. = 1, $P = 0.00001$). The time points that were significantly different occurred at 65 and 72 hr (Fisher's exact test, $P = 0.0002$ and $P = 0.0001$, respectively). Further, males survived longer than females, for both infected (Mantel Cox log-rank test, $\chi^2 = 1282$, d.f. = 1, $P < 0.00001$) and uninfected wasps (Mantel Cox log-rank test, $\chi^2 = 55.1$, d.f. = 1, $P < 0.00001$).

Host mating behavior: We hypothesized that increases in *W. pipientis* densities in the *N. giraulti* background, especially into the brain or sensory organs of the head region (Figure 2, C and D), could affect mate discrimination. We therefore measured the frequency of interspecific matings between *N. vitripennis* 12.1 males and IntG12.1 infected, IntG12.1T tetracycline-treated, or IntG13.2 uninfected, control females in the

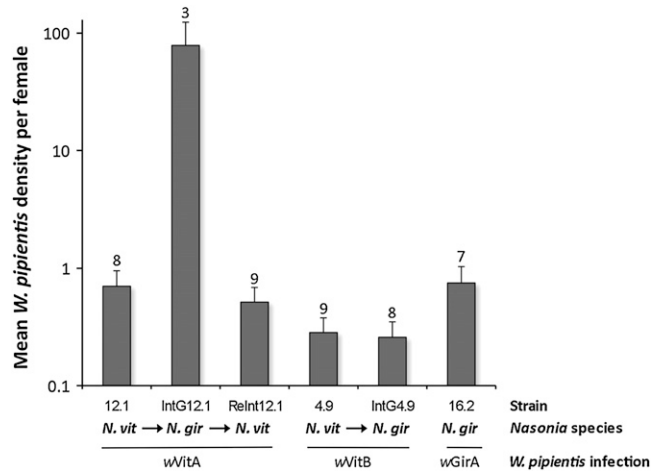


FIGURE 5.—Increase in *W. pipientis* densities is specific to the bacterial genotype. Columns denote the mean *W. pipientis* densities \pm standard error in the native *N. vitripennis* host (A-infected 12.1 and B-infected 4.9), the native *N. giraulti* host (A-infected 16.2), and the introgressed *N. giraulti* (A-infected IntG12.1 and B-infected IntG4.9) and reintroduced *N. vitripennis* (A-infected RelInt12.1) genetic backgrounds. Numbers above columns denote female sample size.

introgressed *N. giraulti* genetic background. Figure 8A shows a highly significant increase (χ^2 , $P < 0.0001$) in interspecific mating percentage between IntG12.1 infected females (66% acceptance, $N = 118$, IntG12.1) *vs.* the IntG13.2 uninfected, control females (33% acceptance, $N = 87$, IntG13.2).

The same increase in interspecific mating was also evident between IntG12.1 infected females (60% acceptance, $N = 58$, IntG12.1) and IntG12.1T antibiotic-treated females (40% acceptance, $N = 70$, IntG12.1T) (χ^2 , $P = 0.02$). Overall, the presence of high *W. pipientis* densities in females resulted in less discrimination and therefore more interspecific copulations. Similar effects were not observed in low-density 12.1 *N. vitripennis* females who showed strong interspecific mate discrimination irrespective of whether they were infected (4.9% acceptance, $N = 82$) or antibiotic-treated (7.9% acceptance, $N = 38$) (χ^2 , $P = 0.51$) when paired with *N. giraulti* males.

CI: CI is a postmating incompatibility between infected males and uninfected females or females harboring a different infection from the male. The bacterial density model of CI posits that variation in incompatibility levels positively correlates with bacterial densities in the host (BREEUWER and WERREN 1993). While investigations of the bacterial density model of CI traditionally test for a positive correlation between *W. pipientis* titers and the expressivity of CI, as is observed in crosses between *N. vitripennis* 12.1 infected males and uninfected females in this study ($\rho = -0.38$, $P = 0.021$, Figure 9), an extension of the model described in the original article was that differences in bacterial densities between males and females

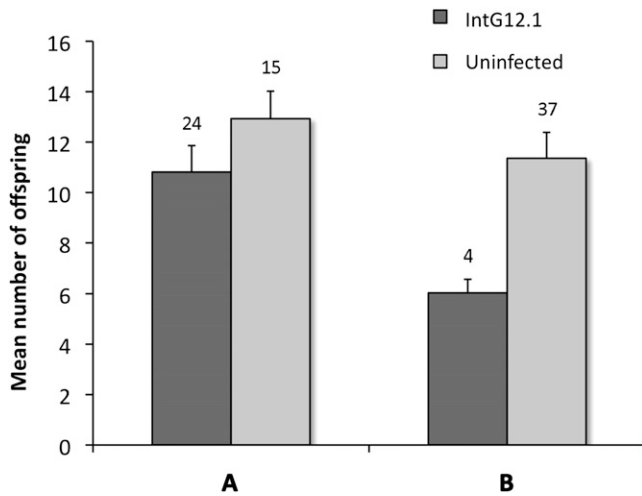


FIGURE 6.—Fecundity cost in infected *N. giraulti* IntG12.1 females. High densities of *W. pipientis* in *N. giraulti* IntG12.1 results in reduced fecundity relative to IntG13.2 uninfected, control females (A) and IntG12.1T tetracycline-cured females (B). Bars represent means \pm standard errors. Numbers above columns denote female sample size.

could lead to different incompatibility types that cause bidirectional incompatibility. For instance, males with high-density infections in testes could be incompatible with females that have low densities in their eggs, or conceivably eggs with significantly higher bacterial densities than the testes may be predisposed to a cell cycle asynchrony in the relative timing of the male and female pronuclei. Here we take advantage of the high- and low-density *Nasonia* lines in this study to explicitly test this prediction.

In the haplodiploid *Nasonia*, *W. pipientis*-induced CI results in a reduction in the number of diploid females produced since haploid males are produced without fertilization. Therefore, the predicted trend for the bacterial density model is that incompatible crosses yield low numbers of female offspring, *i.e.*, high CI inducers. Results indicate that differences in bacterial density do not cause major effects on bidirectional CI. Crosses between low-density 12.1 *N. vitripennis* females and either IntG12.1 high-density or IntG13.2 control, uninfected *N. giraulti* males (Figure 10A) showed no statistical difference in mean number of male offspring (9.56 *vs.* 9.45, MWU, $P = 0.86$), female offspring (29.69 *vs.* 23.27, MWU, $P = 0.33$), and total family size (39.25 *vs.* 32.72, MWU, $P = 0.23$). However as expected for unidirectional CI, the IntG12.1 high-density males were completely incompatible with uninfected females with a complete reduction in female offspring (0 *vs.* 23.27, $P = 0.0001$) and correlated increase in the number of males (24.22 *vs.* 9.45, MWU, $P = 0.002$) in comparison to the control cross with uninfected males.

Crosses in the reciprocal direction between high-density IntG12.1 *N. giraulti* females and either low-density or control, uninfected *N. vitripennis* males

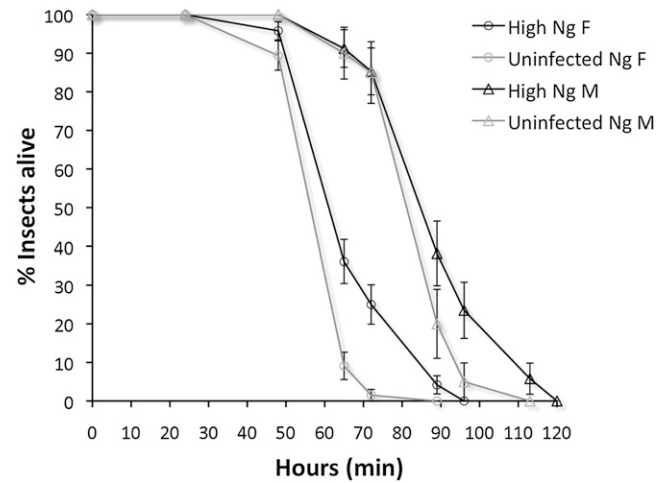


FIGURE 7.—Survival curves of infected and uninfected *N. giraulti* introgression males and females. Survival under starvation conditions was higher in infected *N. giraulti* females relative to IntG13.2 uninfected, control females. The fraction of live individuals was counted at 0, 24, 48, 65, 72, 89, 96, and 120 hr. Each point denotes mean \pm standard error. Labeling denotes male (M) and female (F).

(Figure 10B) showed no statistical difference in the number of males (0.96 *vs.* 0.87, MWU, $P = 0.71$) and a slight, significant difference in the number of female offspring (7.50 *vs.* 9.73, MWU, $P = 0.02$), which caused a difference in total family size (8.46 *vs.* 10.60, MWU, $P = 0.01$). These results specify a marginal increase in CI between a high-density IntG12.1 *N. giraulti* female and a low-density 12.1 *N. vitripennis* male. As expected in the control incompatible cross, unidirectional CI between the low-density 12.1 males and uninfected IntG13.2 females caused a strong but incomplete reduction in female offspring (0.80 *vs.* 9.73, MWU, $P < 0.0001$) and increase in males (3.30 *vs.* 0.87, MWU, $P = 0.0002$), which is consistent with previous experiments on these lines (BORDENSTEIN *et al.* 2003).

Male-to-female transfer of *W. pipientis*: Finally, to test for transfer of the infection from an infected male with unusually high bacterial densities to an uninfected female, two sets of intraspecific crosses were set up using the IntG12.1 and 12.1 infected males with their antibioticly treated female counterparts. In the resident, low-density 12.1 *N. vitripennis* males ($N = 40$), no transfer of *W. pipientis* DNA was observed on the basis of the absence of PCR-amplifiable DNA in uninfected females immediately after mating. However, the high-density IntG12.1 *N. giraulti* males showed clear evidence of transfer of *W. pipientis* DNA to uninfected females. Immediately after mating, 14 out of 20 uninfected females yielded a PCR product for *W. pipientis* (16S rRNA gene) and 15 out of 20 yielded a product for bacteriophage WO (gp17). Subsequent PCRs from 2 through 5 days after mating indicated no detectable *W. pipientis* DNA, suggesting that most transferred *W. pipientis* or *W. pipientis* DNA persists for <24 hr after

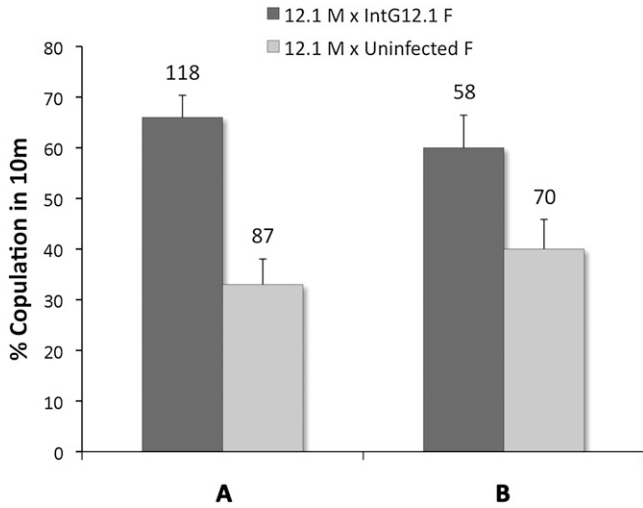


FIGURE 8.—Interspecific female mate discrimination is reduced in IntG12.1 infected females. Mate discrimination is reduced in infected *N. giraulti* females in comparison to IntG13.2 uninfected, control females (A) and IntG12.1T antibioticly treated females (B). Labeling denotes male (M) and female (F). Bars represent means \pm standard errors, and sample sizes are shown above the bars.

mating (Figure 11). Fluorescent *in situ* hybridizations with the *W. pipientis* 16S rRNA-targeting probes in freshly mated females confirmed transfer of *W. pipientis* cells in the female’s spermatheca (4 out of 9 samples, Figure 3C). No *W. pipientis* fluorescence was observed in six, negative uninfected control spermathecae. Further, no *W. pipientis* or bacteriophage WO PCR product was detected in DNA from 10 pooled offspring of uninfected females mated to high-density IntG12.1 *N. giraulti* males ($N = 55$ families, 0/550 offspring, 95% confidence interval from 0 to 0.67%), indicating that on a laboratory scale, paternal transmission to offspring is rare or does not occur.

Detection of the transferred *W. pipientis* cells in the spermathecae of IntG12.1T uninfected females and in the testes of IntG12.1 infected males showed an aggregation of bacterial cells clumped together (Figure 3, A and C). Control crosses between tetracycline-

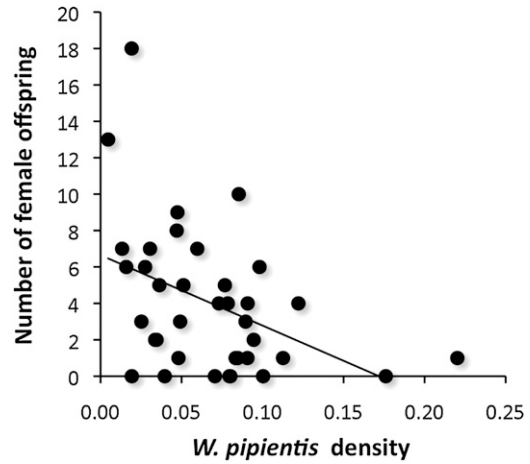


FIGURE 9.—*W. pipientis* densities correlate with CI levels. Points denote the *W. pipientis* density and total number of female offspring from single-pair matings between *N. vitripennis* 12.1 males and uninfected females. Line represents the least squares regression.

cured IntG12.1T *N. giraulti* males and females ($N = 6$) did not show this signal, ruling out the alternative that the aggregate is due to nonspecific binding of the probe. To preliminarily discern the role of a live female in eliminating paternally transmitted *W. pipientis*, crosses were set up similar to above, but the uninfected females were decapitated immediately after mating. Comparisons of these dead females at 24 hr time points ($N = 5$ females for each time point) showed that dead females recapitulated the high frequency of paternally transmitted *W. pipientis* in the first 24 hr, but *W. pipientis* and phage DNA were still detectable in decapitated, IntG12.1T uninfected females even after 20 days (*W. pipientis*, 2/5 individuals; bacteriophage, 5/5 individuals). To test whether *W. pipientis* DNA normally remains in infected wasps postmortem, we examined dead *N. vitripennis* females from infected 12.1 wasps between 0 and 150 days after death. All template DNA across the time points yielded PCR amplicons from *W. pipientis*, phage WO-B, and *Nasonia* (Figure 12).

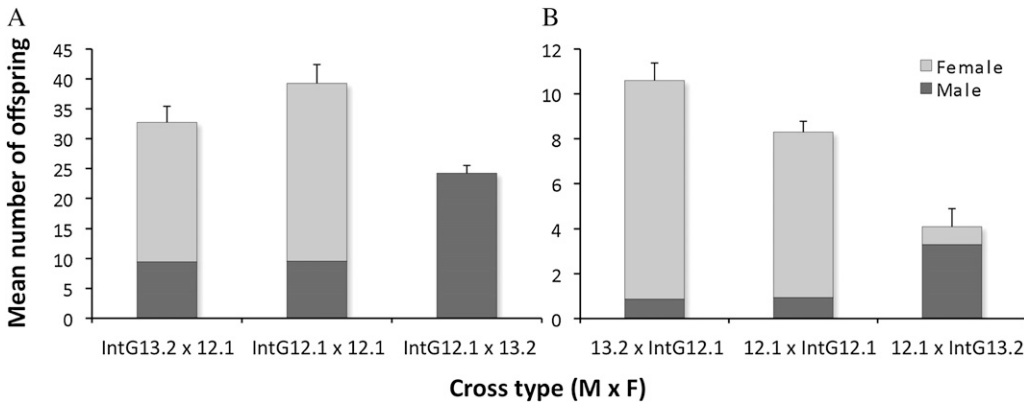


FIGURE 10.—The bacterial density model of CI does not explain bidirectional CI. Experimental and control crosses are shown for testing incompatibility between (A) high-density males and low-density females and (B) low-density males and high-density females. Data are represented as the average number of male and female offspring produced per cross \pm the standard error for total offspring.

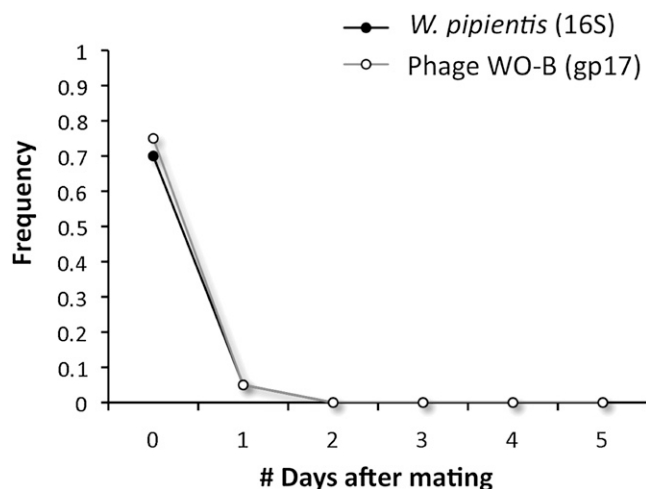


FIGURE 11.—Male-to-female transfer of *W. pipientis* upon transfer to new host species. *W. pipientis* transferred from infected, IntG12.1 males to uninfected females is transiently detectable in young females. Points denote mean frequency of PCR product detected in uninfected females over 5 days in 20 female replicates per day.

DISCUSSION

Transfer of symbionts to new host species can decouple host–symbiont coadaptations within the resident species background. Within *N. vitripennis*, the supergroup A strain of *W. pipientis* has a relatively low level of infection. Upon introgression of the *N. giraulti* host genome into the A-infected *N. vitripennis* cytoplasm, the bacterial concentrations approach seemingly virulent numbers, inundating tissues throughout the entire body cavity. The new strain is therefore an excellent candidate to study *W. pipientis*–host interactions with a strong host genetic basis. The influence of *W. pipientis* infections in different host species varies widely in both nature and intensity (BREEUWER and WERREN 1993; BORDENSTEIN and WERREN 1998; GRENIER *et al.* 1998; POINSOT *et al.* 1998; FUJII *et al.* 2001). For some strains, the type of reproductive parasitism expressed can alternate depending on the host species (SASAKI and ISHIKAWA 2000; JAENIKE 2007). For others, host genetic background has been shown to influence *W. pipientis* density and the degree of CI (BOYLE *et al.* 1993; BORDENSTEIN and WERREN 1998; SINKINS *et al.* 2005; ZABALOU *et al.* 2008), as well as to redefine the types of tissues able to support *W. pipientis* replication (MIN and BENZER 1997; DOBSON *et al.* 1999; CHENG *et al.* 2000; FRYDMAN *et al.* 2006).

Compared to the low-density 12.1 *N. vitripennis*, the IntG12.1 *N. giraulti* host bears a relative bacterial load that is two orders of magnitude higher and a relative phage/bacteria ratio that is twofold lower (Figure 4). The perturbations in microbial densities associated with the *N. giraulti* host were not observed with the *N. vitripennis* B infection, which does not harbor a phage with lytic activity (BORDENSTEIN *et al.* 2006). Thus, the

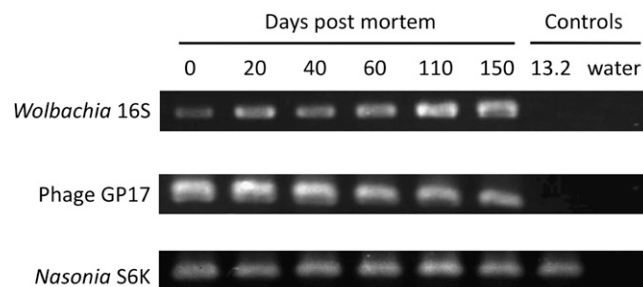


FIGURE 12.—Amplifiable DNA from *W. pipientis*, phage WO-B, and *Nasonia* in dead, infected females. PCR products from infected 12.1 *N. vitripennis* females spanning 0–150 days postmortem. Negative controls are derived from the uninfected 13.2 *N. vitripennis* and double distilled water.

differences in density and somatic tissue infection are due to a specific interaction between the A infection/phage combination of *N. vitripennis* and the *Nasonia* nuclear genomes. It is also possible that mitochondria (mt) are involved in the interaction since the *N. vitripennis* cytoplasm with both *W. pipientis* and mt is placed into a *N. giraulti* genetic background. The *Nasonia* mtDNA have one of the highest rates of substitution among all animal mitochondria (OLIVEIRA *et al.* 2008). Breakdown between cytonuclear interactions could conceivably alter the homeostasis of the endosymbiosis. For example, the mitochondrial heat shock protein 60 is upregulated in *Onchocerca volvulus* filarial nematodes following *W. pipientis* removal (PFARR *et al.* 2008). Reversal of the introgression from *N. giraulti* back into *N. vitripennis* recapitulates the low *W. pipientis* densities and high phage densities native to this species. The genetic pathways involved in this tripartite interaction between invertebrates, bacteria, and phage remain an important, future direction of study.

Inverse association between phage WO-B and *W. pipientis*: We established that *W. pipientis* densities within infected males of each species correlate inversely with phage WO-B densities. We also showed that the dramatic increase of *W. pipientis* densities in the new host species was associated with a significant reduction in relative phage densities. These data corroborate our previous findings on how the phage and Wolbachia densities are intertwined in *N. vitripennis* (BORDENSTEIN *et al.* 2006). There are at least three possible explanations for why the phage densities change after movement to a new host species.

First, a tritrophic interaction between the phage, *W. pipientis*, and insect nuclear genome may govern a reduction in phage densities in the novel host species, which ultimately results in higher relative *W. pipientis* densities. This reticulated control of the symbiosis could, in theory, arise by selection on arthropods to utilize the phage as a natural therapy against *W. pipientis* infection. This hypothesis is based on two arguments: (i) in the absence of symbiont-induced fitness advantages, the phage and arthropod have a common selective

pressure in killing the bacteria—the phage uses it for replication and the host arthropod could suppress fitness costs or expressivity of cytoplasmic incompatibility; (ii) furthermore, ankyrin repeats are found within both the *W. pipientis* and phage genomes (MASUI *et al.* 2000; WU *et al.* 2004; ITURBE-ORMAETXE *et al.* 2005; KLASSON *et al.* 2008; KLASSON *et al.* 2009), indicating the potential occurrence of protein–protein interactions between the phage and insect genome. While these ankyrin repeats have been suggested to be causative factors in the induction of reproductive parasitism, they may alternatively facilitate a direct interaction between *Nasonia* and the phage that mediates the regulation of lytic activity and a reduction in *W. pipientis* densities. Similarly, host level selection could also explain its absence from the genome of mutualistic *W. pipientis* in filarial nematodes (FOSTER *et al.* 2005). Selection at the host level to maintain the beneficial symbiosis could select against a lytic phage of a mutualistic infection, which is required for host oogenesis and larval development (BORDENSTEIN *et al.* 2006). Strict vertical transmission, genome reduction, and absence of coinfections of *W. pipientis* in filarial nematodes are also adequate reasons for the absence of bacteriophage in this species.

A second explanation for the inverse association between phage WO-B and *W. pipientis* is that the *W. pipientis* cells are stably replicating in the new *N. giraulti* background, leading to a suppression of the bacterial interactors that elicit lytic phage development. Perhaps the new host is more permissive for *W. pipientis* growth. The basic rationale here is that if phage WO-B responds to bacterial stress responses by entering lytic development, then by extension, the *W. pipientis* in *N. vitripennis* could be more stressed than the same infection in *N. giraulti*. Many temperate phages “sense” stressed bacteria, thereby inducing prophage excision, replicating as virions, and triggering bacterial lysis. There may be many ways by which a sensing feedback mechanism leads to phage induction, and one of the most common is the SOS response. Curiously, key SOS genes are present in the *Wolbachia* genome, including *recA*, *uvrA*, *uvrB*, *uvrD*, and a *cI*-like repressor; *lexA*, however, is missing.

Third and finally, stochastic changes in phage densities during the introgression rather than host genotype could also explain the change in phage densities. However, the IntG12.1 strain has now been maintained in the laboratory since April 2006, indicating a stable host genome interaction that is not subject to random fluctuations in bacterial density. In addition, it is the second time we have generated the strain and observed an increase to complete CI, which is indicative of higher densities, although the densities were not measured in that experiment (BORDENSTEIN and WERREN 1998).

Fitness: The dramatic 100-fold increase in infection densities in the naive host species is likely to modify the traditional costs and benefits associated with the symbiosis. Previous studies indicate fecundity effects of *W.*

pipientis vary widely. Some studies suggest there is no effect of *W. pipientis* on fecundity under laboratory conditions in *D. melanogaster* (CLARK and KARR 2002; REYNOLDS and HOFFMANN 2002) or in *N. vitripennis* (BORDENSTEIN and WERREN 2000), while others have found harmful effects (HOFFMANN *et al.* 1990; NIGRO 1991; STOUTHAMER and LUCK 1993; CALVITI *et al.* 2010). In our study, a decrease in fecundity was found in the high-density IntG12.1 *N. giraulti* females. This negative fecundity effect was not observed within *N. vitripennis* where *W. pipientis* densities are lower and restricted to the host reproductive tissues. Thus, the high-density infection in the introgression line may have intensified effects on fecundity in *Nasonia* that were not seen in comparisons with lower densities (BORDENSTEIN and WERREN 2000).

In contrast to fecundity, survival of starved wasps was slightly extended in *Nasonia* harboring high *W. pipientis* densities. This finding was unexpected since we anticipated the high densities in somatic tissues would yield a decrease in survival. Previous studies showed that *W. pipientis* did not influence adult starvation resistance in the quiescent *wMel* infection of *D. melanogaster* (HARCOMBE and HOFFMANN 2004). However, the rapidly proliferating *wMel* Popcorn strain in laboratory-reared *D. melanogaster* causes early mortality (MIN and BENZER 1997). Possible reasons for the increase in survival include (i) an increase in immunity to other bacterial or viral parasites, as evident from recent findings of RNA viral resistance mediated by *W. pipientis* (HEDGES *et al.* 2008; TEIXEIRA *et al.* 2008); (ii) a life history tradeoff in which decreased fecundity in infected females may lead to an increased energy investment in adult survival; in this study, fecundity was significantly reduced, but not measured over the lifetime of the female; and (iii) mitochondrial haplotype differences between the experimental lines; IntG12.1 and IntG13.2 are derived from *N. vitripennis* lines 12.1 and 13.2 in a segregation experiment conducted in 1996 (PERROT-MINNOT *et al.* 1996). Further work is necessary to determine the importance of these mechanisms.

Other recent studies suggest the fitness interactions between host and *W. pipientis* are more complex than previously recognized. For instance, loss of insulin-like ligands produced in the *Drosophila* brain greatly extended lifespan in the presence of *W. pipientis*, demonstrating a specific interaction between the insulin/IGF signaling (IIS) pathway and the bacteria in lifespan regulation (GRONKE *et al.* 2010). Coevolution between host and *W. pipientis* results in an intricate merger of signaling pathways, which can result in highly specific phenotypic effects unique to each association.

Bidirectional cytoplasmic incompatibility: The positive association between *W. pipientis* densities and incompatibility levels is characterized as the “bacterial density model” of CI (BREEUWER and WERREN 1993), and numerous studies have confirmed positive correla-

tions between CI levels and bacterial densities within eggs, sperm cysts, whole adults, and reproductive tissues (POINSOT *et al.* 1998; PERROT-MINNOT and WERREN 1999; NODA *et al.* 2001; CLARK and KARR 2002; VENETI *et al.* 2003). Extrinsic factors that affect *W. pipientis* density can modify the nature and degree of CI including host genetic background (BOYLE *et al.* 1993; BORDENSTEIN and WERREN 1998; SINKINS *et al.* 2005) and environmental stresses such as host age (HOFFMANN *et al.* 1986; SINKINS *et al.* 1995; FEDER *et al.* 1999; CLARK *et al.* 2002), heat treatment (FEDER *et al.* 1999), mating history (KARR *et al.* 1998), larval crowding (SINKINS *et al.* 1995), as well as lytic phage proposed as part of the “phage density model” (BORDENSTEIN *et al.* 2006).

Here we used high- and low-density *Nasonia* lines to explicitly test a second prediction of the bacterial density model of CI: bidirectional CI may result from gender differences in bacterial densities. We showed that the large differences in bacterial loads between IntG12.1 and 12.1 do not lead to strong bidirectional CI. Slight incompatibility observed between low-density males and high-density females could possibly be due to the dense infection in the egg, which causes a change to the embryonic factors that rescue the paternal genome modification caused by CI. Previous studies demonstrated that rescue is either achieved through correction of cell cycle defects in the male pronucleus or through a compensatory slowing of the female pronucleus cell cycle to restore the cell cycle synchrony with the altered male pronucleus (TRAM and SULLIVAN 2002; SERBUS *et al.* 2008). Nonetheless, results generally indicate that male and female differences in bacterial densities do not single handedly explain the significant differences in compatibility (POINSOT *et al.* 1998; PERROT-MINNOT and WERREN 1999; NODA *et al.* 2001; CLARK and KARR 2002; VENETI *et al.* 2003; BORDENSTEIN and WERREN 2007). Thus, these results show that bidirectional incompatibility is more likely due to rapid genetic changes between strains and/or hosts.

Male-to-female transfer of *W. pipientis*: The exceptionally high level of infection in the new genetic background leads to colonization of the male seminal vesicles and accessory glands. This change presumably leads to the high rate of male-to-female transfer observed in crosses between infected males and uninfected females (70%). Both PCR-amplifiable DNA and 16S rDNA probes provided evidence for the sexual transmission. Although the infection did not remain within the living female for >24 hr or transfer to offspring, the presence of *W. pipientis* within an uninfected female could potentiate a rare opportunity for paternal transmission, which is not likely to be captured in small-scale laboratory experiments. Strikingly, dead females who had been decapitated for up to 20 days yielded transferred DNA, suggesting there may be some direct or indirect physiological process that eliminates the *W. pipientis* within 24 hr after transfer. Alternatively, the dead

female could offer a more permissive environment for *W. pipientis* to inhabit. Experiments are underway to determine the viability of *W. pipientis* in dead wasps.

Recently there have been a number of animal and plant systems in which paternal leakage of mitochondrial genomes has been demonstrated in hybrids (WHITE *et al.* 2008). Presumably, there is a hybrid breakdown of mechanisms that recognize and remove paternal cytoplasmic elements such as mitochondria. Therefore, while paternal transfer of *W. pipientis* is absent or rare in *D. melanogaster* (HOFFMANN *et al.* 1998), *D. simulans* (HOFFMANN and TURELLI 1988), and *Aedes albopictus* (DOBSON *et al.* 2002) within species, our study suggests more attention to paternal transfer in hybrid zones or hybrids in the lab because the IntG12.1 is a hybrid composed of a *N. vitripennis* cytoplasm and *N. giraulti* genome.

Host behavioral pathology: Finally, previous studies showed that *W. pipientis* can alter host behaviors. One study found *W. pipientis*-induced effects on precopulatory behavior where inbred, uninfected female spider mites preferentially mated with uninfected males to avoid *W. pipientis*-induced CI (VALA *et al.* 2004). A change in ovipositioning behavior was also reported in that study. Mating behavior influences have also been described in isopods (MOREAU *et al.* 2001) where *W. pipientis* infections distort sex ratios toward females through the feminization of males. Here, males showed preference for genetic females as opposed to copulations with feminized males.

Interspecific mate discrimination between *N. vitripennis* and *N. giraulti* is well characterized and likely plays a role in reinforcing reproductive isolation in sympatric populations because of F₁ CI and severe F₂ hybrid breakdown between the species (BREEUWER and WERREN 1990; BORDENSTEIN and WERREN 1998; DRAPEAU and WERREN 1999). Our studies in the laboratory showed that IntG12.1 infected females have reduced interspecific mate discrimination (*i.e.*, more copulations). Similar mating behavior effects in low-density 12.1 *N. vitripennis* females were not observed. Our results are the opposite of those of a previous study in which the removal of *W. pipientis* from choosy *D. melanogaster* strains resulted in a reduction of discrimination (KOUKOU *et al.* 2006), indicating that *W. pipientis*-induced effects (*i.e.*, on pheromones and/or behavior) enhance a preexisting genetic bias in mate discrimination in both males and females.

A simple explanation for the increase in interspecific mating frequency in our study is that the infection in the head regions of *N. giraulti* females causes a behavioral pathology in which females suffer from a reduced competence to discriminate against interspecific males. This conclusion is based on two observations. First, neurological infections of *W. pipientis* in *Nasonia* are abnormal, potentiating a pathology related to behavior. Second, wasps that are infection-free show high levels of discrimination typical of *N. giraulti* females infected with their native *W. pipientis* strain (BORDENSTEIN and WERREN 1998).

Concluding remarks: The arthropod–*W. pipientis* association is an ancient symbiosis, yet new symbiotic combinations between different *W. pipientis* strains and host species constantly evolve as the infection promiscuously transfers to new arthropod species. A conflict or compromise between the endosymbiont and host must ensue, which determines the outcome of the symbiosis. Here we have shown that the two-order magnitude increase in relative *W. pipientis* densities in a new host species leads to reduced fecundity, higher levels of CI, and unexpectedly, male-mediated transfer of the bacteria to females and a behavioral pathology. Results from this study also raise the idea that the host may have direct or indirect effects on biological entities one trophic level down from *W. pipientis*, *i.e.*, the bacteriophage WO, which could alter densities of the endosymbiont. The host genetic mechanisms within *Nasonia* that permit/suppress a hyperinfection to take a permanent hold, and the question of whether the molecular interactions with the host involve both the bacteria and phage, remain an important topic of investigation.

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Supporting Information

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Decoupling of Host–Symbiont–Phage Coadaptations Following Transfer Between Insect Species

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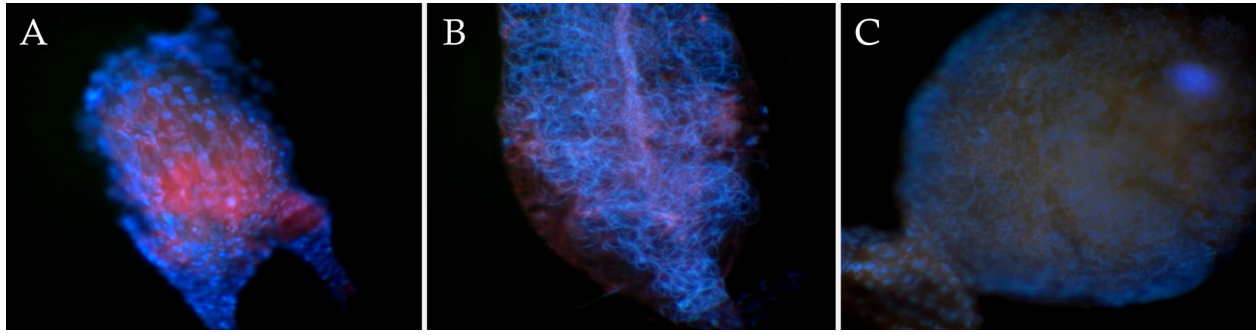


FIGURE S1.—Testes from infected and uninfected *Nasonia*. *W. pipientis* probes W1 and W2 (red) and nucleic acid (DAPI blue) shown in the testes of the high density *N. giraulti* IntG12.1, low density *N. vitripennis* 12.1, and antibiotically-cured *N. giraulti* IntG12.1T.