Fitness costs of phoretic nematodes in the burying beetle, *Nicrophorus vespilloides*

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Abstract

*Nicrophorus vespilloides* is a social beetle that rears its offspring on decomposing carrion. Wild beetles are frequently associated with two types of macrobial symbionts, mites, and nematodes. Although these organisms are believed to be phoretic commensals that harmlessly use beetles as a means of transfer between carcasses, the role of these symbionts on *N. vespilloides* fitness is poorly understood. Here, we show that nematodes have significant negative effects on beetle fitness across a range of worm densities and also quantify the density-dependent transmission of worms between mating individuals and from parents to offspring. Using field-caught beetles, we provide the first report of a new nematode symbiont in *N. vespilloides*, most closely related to *Rhabditoides regina*, and show that worm densities are highly variable across individuals isolated from nature but do not differ between males and females. Next, by inoculating mating females with increasing densities of nematodes, we show that worm infections significantly reduce brood size, larval survival, and larval mass, and also eliminate the trade-off between brood size and larval mass. Finally, we show that nematodes are efficiently transmitted between mating individuals and from mothers to larvae, directly and indirectly via the carcass, and that worms persist through pupation. These results show that the phoretic nematode *R. regina* can be highly parasitic to burying beetles but can nevertheless persist because of efficient mechanisms of intersexual and intergenerational transmission. Phoretic species are exceptionally common and may cause significant harm to their hosts, even though they rely on these larger species for transmission to new resources. However, this harm may be inevitable and unavoidable if transmission of phoretic symbionts requires nematode proliferation. It will be important to determine the generality of our results for other phoretic associates of animals. It will equally be important to assess the fitness effects of phoretic species under changing resource conditions and in the field where diverse interspecific interactions may exacerbate or reduce the negative effects of phoresy.

Keywords

commensalism, nematode, *Nicrophorus vespilloides*, phoresy
1 | INTRODUCTION

Animals that persist on ephemeral and spatially dispersed resources have evolved diverse mechanisms to detect and exploit these resources (Janzen, 1977; Shivik & Clark, 1997; Stavert, Drayton, Beggs, & Gaskett, 2014). Carion feeders, like blowflies and burying beetles, can use olfactory cues to detect minute concentrations of the volatile products of animal decomposition and can orient their search flights accordingly (Ashworth & Wall, 1994; Kalinová, Podskalská, Růžička, & Hoskovec, 2009). However, some animals are incapable of moving across large distances themselves. Instead, these species hitch a ride on the bodies of other more mobile species and are consequently transported from resource to resource (Bartlow, Villa, Thompson, & Bush, 2016; Guerra, Romero, Costa, Lofego, & Benson, 2012). Thus rather than developing mechanisms to detect resources, they have evolved mechanisms to ensure reliable and durable associations with the species that carry them (Krishnan, Muralidharan, Sharma, & Borges, 2010; von Beeren & Tishechkin, 2017). This strategy, known as phoresy, is common in many species of insects, mites, and nematodes and is a form of symbiosis that is typically believed to be harmless to the host (Houck, 2009; Signe White, Moran, & de Roode, 2017). The rationale for this belief is that because phoretic species are wholly dependent on their hosts for their migration, species that cause too much harm and thereby reduce their transport between breeding resources face the risk of local extinction (Signe White et al., 2017). However, just as parasites and pathogens can evolve levels of virulence that balance harm and fitness to host with the need to be transmitted between hosts, so too may phoretic species become parasitic, as long as this harm facilitates their transmission between hosts (Alizon, 2008; Houck & Houck, 1994; Little, Chadwick, & Watt, 2008; Munoz, Perfectti, Martin-Alganza, & Camacho, 1998). Few studies have quantified the direct harm of phoretic species to their hosts while also estimating their persistence and transmission between host individuals and across generations. Our aim in this paper was to address these questions in the context of the burying beetle, Nicrophorus vespilloides, and its phoretic nematodes.

Nicrophorus burying beetles are subsocial insects that breed on small vertebrate carrion (Scott, 1998). After locating a small vertebrate carcass using volatile cues produced from microbial decomposition (Kalinová et al., 2009), a mated female lays eggs in the surrounding soil after which she (or a mated pair) prepares the carcass for the arrival of the hatched larvae (Milne & Milne, 1976). The carcass is buried underground, stripped of fur or feathers, the gut is removed, and then, it is coated in antimicrobial oral and anal secretions (Arce, Smiseth, & Rozen, 2013; Cotter & Kilner, 2010; Duarte et al., 2016; Jacobs et al., 2014; Trumbo, 2017). When larvae migrate to the carcass, parents remain to feed them via regurgitation (Eggert, Reinking, & Müller, 1998; Scott, 1998), which both provides a meal and also transmits the endogenous microbiome to the developing larvae (Duarte, Welch, Swannack, Wagner, & Kilner, 2017; Shukla, Vogel, Heckel, Vilcinskas, & Kaltenploth, 2017; Vogel et al., 2017; Wang & Rozen, 2017). But beetles are not alone in their consumption of the carcass. Nicrophorus adults trapped in the field are conspicuously associated with high densities of mites and nematodes that are attached to their carapace or reside internally (Gasperin, Duarte, & Kilner, 2015; Richter, 1993). Many species of mites have established phoretic associations with burying beetles, and their well-studied effects on beetles range from harmful to beneficial, depending on the context and the study (Gasperin & Kilner, 2015; Nehring, Müller, & Steinmetz, 2017; Wilson & Knollenberg, 1987). By contrast, only one species of phoretic nematode has been described in Nicrophorus and its effects on beetles are unknown (Richter, 1993).

Richter (1993) described the carrion-feeding nematode Rhabditis stammeri isolated from N. vespilloides. He showed that worms were present in the gut and genitalia and could be transmitted between mating individuals. However, although Nicrophorus researchers regularly comment on the presence of nematodes in laboratory and field populations, there is no direct evidence that these worms are actually R. stammeri, nor is there any understanding of their natural abundance in field-caught insects. More importantly, we lack an experimental understanding of the fitness consequences of these nematodes for beetles. As part of our efforts to understand the evolution and ecology of phoretic associates of N. vespilloides, we provide a detailed study of the identity and effects of a novel nematode associate of N. vespilloides, most closely related to Rhabditoides regina, that we cultured and quantified from field-caught beetles. In brief, we find that these nematodes are extremely numerous in wild beetles and significantly reduce N. vespilloides fitness. In addition, worms are efficiently transmitted in high densities between mating adults and from infected mothers to their offspring, which then persist through beetle development and are retained into adulthood. We discuss these results in the context of the evolution of interspecific interactions in N. vespilloides and the evolution of host harm in phoretic species.

2 | METHODS

2.1 | General procedures

All experimental beetles were taken from an outbred laboratory population derived from wild-caught N. vespilloides individuals trapped in Warmond, near Leiden in The Netherlands, between May and June 2016. Beetles were maintained in the laboratory at 20°C with a 15:9 hr light:dark cycle. All adults were fed fresh chicken liver twice a week. To maintain the laboratory population and to establish experimental broods, an unrelated male and female were placed together overnight without food in one small plastic containers filled with 1–2 cm of autoclaved soil for mating. The following morning, mated females were provided with a freshly thawed mouse carcass in a new larger container for egg laying. Broods were reared until larvae dispersed from the carcass, approximately 7 days post-hatching (Monteith, Andrews, & Smiseth, 2012). Dispersed larvae were placed together into a new container of sterile soil until eclosion, at which point they were removed to new individual containers.
To generate nematode-free adults, eggs were collected from broods within 12–24 hr of laying and surface-sterilized with an antimicrobial solution of hen egg white lysozyme (1 mg/ml), streptomycin (500 µg/ml), and ampicillin (100 µg/ml) (Jacobs et al., 2014). These were then transferred onto 1% water agar plates to hatch, after which they were placed onto a freshly thawed mouse carcass that had been opened using a sterile scalpel. To prevent nematode transmission from parents to newly hatched larvae, first-generation nematode-free larvae were reared without parental care. Once these nematode-free individuals had eclosed as adults, they were crossed as above, and maintained thereafter on autoclaved soil.

2.2 Nematode quantification from field-caught and laboratory beetles

Nematodes were collected and counted from the guts and cuticles of field-caught and laboratory-reared beetles. Individual adult beetles were vortexed for 3 min in 700 µl of sterile phosphate buffer saline (PBS, pH = 7.2) to collect nematodes from the cuticle. To quantify nematodes from the beetle gut, we removed individual beetle guts with fine forceps and suspended these in 700 µl of sterile PBS. 10 µl of each suspension (cuticle or gut sample) was then transferred onto an hemocytometer and examined at 10× magnification for counting. Three independent 10 µl aliquots were counted from each sample to generate a mean estimate/sample.

2.3 Nematode maintenance and identification

Experimental nematodes were isolated directly from the cuticles of field-caught beetles. Species identification is explained below. To maintain laboratory populations, newly collected nematodes were transferred onto Petri plates containing Nematode Growth Medium (NGM contains 1.7% agar/l; 50 mM NaCl; 0.25% peptone; 1 mM CaCl₂; 5 µg/ml cholesterol; 25 mM KH₂PO₄; 1 mM MgSO₄; Stiernagle, 2006) and fed with an E. coli strain originally isolated from a mouse carcass and held at 20°C. Nematodes were transferred to fresh plates containing E. coli at an initial density of ~10⁶ cells/plate every 2 days.

To determine species identity, nematode samples reared on NGM plates were collected and suspended in sterile PBS (100 mM, pH 7.2), after which they were surface-sterilized in a wash solution containing a 1:2 ratio of 5 N NaOH and a 5% solution of sodium hypochlorite (Stiernagle, 2006). Washed nematodes were re-suspended in 1 ml PBS and then centrifuged at 13,000×g for 10 min. Nematode pellets were re-suspended in 0.7 ml PBS and stored at −20°C. For DNA extraction, nematode samples were thawed and homogenized with a sterile micropestle and vortexed for 2 min. Samples were then lysed in SDS at 60°C for 30 min. following the method of Donn, Griffiths, Neilson, and Daniell (2008). DNA was extracted using phenol–chloroform and quantified using a Thermo NanoDrop ND-1000 spectrophotometer. The 18S rRNA gene fragment (~900 bp) was amplified using primer pairs Nem_18S_F (CGGCAATRGCTCATACAACAGC) and Nem_18S_R (GGGCCGTATCTGATCC) (Floyd, Rogers, Lambshead, & Smith, 2005). For PCR amplification, 2 µl of template containing 2–10 ng DNA was used directly in a 20 µl reaction mixture using *Pyrococcus furiosus (Pfu)* DNA Polymerase. PCR was performed in a thermal cycler (Bio-RAD T100™) with thermal cycling of 95°C for 5 min, followed by 35 cycles of 95°C for 30 s, 54°C for 30 s, 72°C for 30 s, and a final extension at 72°C for 5 min. PCR products (fragment length of ~900 bp) were gel purified (Illumina™ GFX™ PCR DNA and Gel Band Purification Kit) and sequenced commercially via MaxyGen. The resulting 18S rRNA gene sequence was classified to species using a nucleotide BLAST against the NCBI database.

2.4 Fitness effects and transmission of nematode infections

To determine the fitness effects of nematodes on beetles, broods were established with worm-inoculated mated females (at least 20 broods/inoculation density). All broods were established with virgin females that had eclosed at least 7 days prior to mating. Before inoculation, nematodes were surface-sterilized to remove any surface-associated bacteria and then suspended in sterile PBS. Worm densities were quantified prior to inoculation by direct counts using an hemocytometer. Experimental worm-free beetles were inoculated with either ~10, 10², 10³ or 10⁴ nematodes per beetle by pipetting worm solutions under their elytra and on their mouth and anus. Two days later, inoculated females were paired with an unrelated worm-free male and allowed to mate overnight in small plastic containers. The next morning, males were removed and the mated females were provided with a freshly thawed mouse carcass and allowed to rear their broods until the point of larval dispersal. We tightly controlled carcass mass to ensure uniformity across our treatment groups given the known association between carcass mass and beetle reproductive fitness. Carcass mass ranged from 20 to 24.11 g with an overall mean (±SE) of 22.31 ± 0.9 g. There were no significant differences in carcass mass between our treatment groups (ANOVA: F4,113 = 0.169, p = 0.954). When the beetle larvae dispersed from the carcass, we measured brood size, total brood mass, and mean larval mass for each brood, as well as the number and fraction of eclosing adults.

To quantify nematode transmission between *Nicrophorus* individuals, we measured the number of worms transmitted between experimentally inoculated beetles and worm-free recipients. Transmission was examined between mating adults and from mothers to offspring.

Nematode transmission was quantified bidirectionally between males and females (i.e., female donors to male recipients and male donors to female recipients). Individual adults were first inoculated with different worm densities, as outlined above, and then maintained for 2 days in small boxes containing 1–2 cm of sterile soil. Next, these individuals were transferred to a new box with sterile soil and paired with a nematode-free individual of the opposite sex for mating. Two days later, both individuals were sampled to determine nematode densities.

To estimate worm transmission from parental females to offspring, females were inoculated with different worm densities,
allowed to mate with a worm-free male, and then provided a fresh carcass for breeding. When beetle offspring eclosed, they were sampled to estimate nematode densities.

2.5 | Statistical analyses

There were 17 of 20 successful broods in the “no nematodes” treatment, 26 of 32 in the “10 nematodes” treatment, 25 of 32 in the “100 nematodes” treatment, and 19/32 in the “10,000 nematodes” treatment. A Shapiro–Wilk test based on successful broods was used to test for normality in experiments examining the effects of nematode infection on larval fitness (All <0.05). We used ANCOVA to test for significant effects on (a) brood size, (b) total brood mass, (c) mean larval mass, (d) number of eclosed adults, and (e) fraction of eclosed larvae, while controlling for carcass size. The relationship between brood size and mean larval mass within each nematode treatment was examined using linear regression. We used a generalized linear model (GLM) to test for interactions between nematode infection and the trade-off between brood size and mean larval mass. Differences in nematode transmission were estimated using tests. All analyses were performed using SPSS version 24 (IBM SPSS, Inc., Chicago, IL, USA).

3 | RESULTS

3.1 | Nematode identification and infection densities in wild beetles

The partial nematode 18S rRNA gene sequence (~900 bp) was BLASTed against the NCBI database and showed 95% identity to Rhhabditoideas regina strain DF5012 (AF082997) and was clearly distinct from Rhabditis stammeri, the nematode species already described from N. vespilloides. With the resolution we have from this sequence, it remains uncertain whether our isolate is truly R. regina or an as-yet-undescribed species; further sequencing will be required in a later study to more fully resolve its taxonomy. For ease of presentation, we hereafter tentatively refer to our isolate as R. regina.

3.2 | Effect of different starting nematode densities on larval fitness

Nematode infections are highly costly to beetles. We observed significant treatment effects associated with different nematode densities on all fitness parameters after controlling for carcass mass (Table 1). In addition, we observed a significant negative linear relationship between the number of inoculated nematodes and mean brood size ($r^2 = 0.84$, $p = 0.03$) and mean larval mass ($r^2 = 0.9$, $p = 0.02$) (Figure 2). Brood size declined nearly threefold in broods...
with the highest nematode densities (Mean ± SE: 11.21 ± 2.15 larvae/brood) compared to nematode-free broods (Mean ± SE: 27.25 ± 3.03 larvae/brood), while mean larval mass declined roughly 15% from 0.181 ± 0.006 g in broods without nematode infections to 0.156 ± 0.007 g in broods where females were inoculated with 10,000 worms.

In addition to these direct negative effects of nematode infection, we observed a significant interaction between worm density and the trade-off between brood size and average larval mass ($F = 9.332, df = 1, p = 0.003$, Figure 3). Most interestingly, whereas there was a significant trade-off between brood size and average larval mass in worm-free beetles ($r^2 = 0.50, p = 0.001$), there was no association in broods produced by females infected with worms at any of the treatment densities (10 nematodes: $r^2 = 0.04, p = 0.321$; 100 nematodes: $r^2 = 0.001, p = 0.896$; 1,000 nematodes: $r^2 = 0.072, p = 0.185$ and 10,000 nematodes: $r^2 = 0.0004, p = 0.935$).

### 3.3 Transmission between sexes and from mothers to offspring

We inoculated male or female beetles (donors) with different nematode densities and then measured worm transmission to opposite-sex recipients during mating. As shown in Figure 4a,b, we observed intersexual transmission of nematodes in both directions, although this varied with worm density and the sex of the donor. For both donor sexes, when the initial density of nematodes was 10, there was neither transfer nor retention of worms. Transmission occurred at all other initial worm densities.

At inoculation densities of 100, worm numbers declined slightly in both male and female donors; however, transmission occurred effectively and there were no significant differences in the final worm densities of donors or recipients ($t = -2.05, p = 0.095$). Worm densities were retained at initial values when donor females were inoculated with 1,000 worms but were significantly reduced when the initial inoculum was 10,000 worms ($t = -23.2, p < 0.001$). Final worm densities in males and females (recipients and donors, respectively) did not differ at either inoculum density. In addition, there were no differences in final worm densities between beetles inoculated with 1,000 or 10,000 worms, suggesting an estimated carrying capacity of roughly 2,000 worms per adult beetle (mean ± SE: 1770 ± 238.3).

For male donors, both transmission and retention were reduced compared to female donors. At initial densities of 1,000 or 10,000, we observed significant reductions in donor and recipient worm densities overall (1,000: $t = -56.64, p < 0.001$; 10,000: $t = -83.5, p < 0.001$). There were no differences in final worm densities at all inoculum sizes >10, suggesting a carrying capacity of approximately 125 worms/adult beetle (mean ± SE: 123 ± 41.6). Although both sexes can transfer nematodes to the opposite sex during mating, female transfer and retention are approximately 10× higher in females than in males ($t = 6.3, p < 0.001$).

Next, to measure transmission from mothers to offspring, we inoculated mated females with different densities of worms and then allowed them to rear broods, after which we quantified the number of nematodes on eclosing pupae (Figure 4c). For all inoculum sizes other than 10,000, we observed significant transmission from mothers to larvae (one-sample t test: 10: $t = 2.78, p = 0.039$; 100: $t = 2.43, p = 0.036$; 1,000: $t = 3.02, p = 0.01$). In addition, worm densities on eclosing larvae were significantly or marginally greater than maternal inoculum densities (one-sample t test: 10: $t = 2.6, p = 0.047$; 100: $t = 2.0, p = 0.072$; 1,000: $t = -2.394, p = 0.032$). Finally, we observed no significant differences in the final densities of worms on eclosing larvae from the 10, 100, and 1,000 treatments (one-way ANOVA: $F = 0.79, p = 0.463$, with a mean of approximately 500 nematodes per eclosed individual (mean ± SE: 500 ± 250).
Expectedly, we found negligible transmission when mothers were initially inoculated with 10,000 nematodes, a likely artifact attributed to the extremely high rate of larval mortality in this treatment group (brood success ~7%).

4 | DISCUSSION

Because phoretic species have limited dispersal capacity on their own, they rely on the greater motility of other species to coordinate their longer-distance transport across the environment. Such transport is beneficial and typically obligatory for the phoront, leading to the belief that phoretic species should not harm their hosts, or risk compromising their transmission. However, this expectation is not always realized, and the effects of apparently phoretic species can range from mutualism to parasitism. For example, female bark beetles that carry mites produced larger and heavier offspring, suggesting phoront-specific benefits (Mazza, Cini, Cervo, & Longo, 2011). By contrast, mites carried by the red palm weevil significantly reduce beetle longevity, indicating severe costs (Hodgkin, Elgar, & Symonds, 2010). The factors that determine these different outcomes are varied and context-dependent, and show likely parallels to the diverse factors that influence the evolution of parasite virulence. As with parasites, the virulence of phoretic species may increase if this correlates with increased transmission. It could also increase in cases where the phoront interacts with multiple host species, thereby reducing reliance on any single host, or where there is competition between different genotypes of a single phoront species. Phoront virulence may also vary across different stages of host development, depending on the coupling between the developmental/dispersal stage of the phoretic species and that of its host. To establish the baseline against which to examine these issues, our aim here was to quantify the fitness effects and transmission between adults and across generations of phoretic nematodes of *N. vespilloides*.

![FIGURE 4](image)

Nematode transmission between mated pairs and between generations. (a) Transmission from females to males; (b) transmission from males to females; (c) transmission from mothers to offspring.
the carcass itself (possibly due to consumption of bacteria on the carcass) and also conspicuous worm nictation upon disturbance. Nictation is a behavior commonly seen in phoretic nematodes that is thought to facilitate dispersal. It involves standing upright and waving in all directions, thereby attracting potential hosts (Brown, D’Anna, & Sommer, 2011). In addition, the strong sex bias in nematode densities both in the field (Figure 1) and in the laboratory (Figure 4) is consistent with the idea that worms are maximizing dispersal potential by preferentially associating with the sex most likely to colonize a breeding resource (a carcass). Similar biases have been observed in other phoretic nematodes (Krishnan et al., 2010; Scheffer et al., 2013) and mites (Campbell & Luong, 2016; Frohnofer et al., 2013; Gilburn, Stewart, & Edward, 2009). Our field collections reveal that this species is maintained in high, although variable, densities in male and female wild beetles (Figure 1) while further studies in the benign conditions of the laboratory (YW unpublished) have shown that they are also stably maintained within laboratory populations of burying beetles at even higher densities. Although Nicrophorus nematodes were believed to have no or marginal effects on beetle fitness, our results indicate that this is not the case. Worm infections cause significant harm to beetles, and the extent of this harm scales with worm density for both brood size and mean larval mass (Figures 2 and 3, Table 1), two central measures of adult and larval fitness, respectively. In addition, nematodes are transferred at high rates between adults and from parents to offspring (Figure 4), suggesting that despite host harm, transmission potential is maintained.

We find strong density-dependent effects of nematodes on N. vespilloides. However, even though we see a significant negative relationship between worm numbers and, for example, brood size and mean larval mass, much of the maximum cost observed at the highest worm density (10,000) is already observed at the lowest inoculum size we used (10 worms). In other words, of the ~50% decline in brood size in beetles inoculated with 10,000 worms, around 80% of this decline is already apparent in beetles inoculated with only 10 worms. This result is consistent with the idea that worms are proliferating extensively on the carcass where they can then go on to infect larvae, which seems to occur whether the initial number of colonizing worms is high or low. This result also explains why nematode transmission from parents to offspring has no lower threshold (Figure 4c), in contrast to the threshold of ~100 worms needed for transmission between breeding adults (Figure 4a,b). Equally, while the densities of worms on larvae tend to exceed the inoculum density on females (consistent with proliferation), this is not the case for intersexual transmission, suggesting that nematode reproduction does not occur in this context and that worm transmission between adults suffers from stochastic loss if worms are initially rare.

Less clear are the factors that are responsible for the harm worms cause to reduce beetle fitness. Nematodes can reduce fitness in several ways. Before establishing broods, infected females may either forgo egg laying or reduce the number of eggs they lay to reduce the costs of rearing a brood while interacting with nematodes. Although we do not see any influence of nematode inoculation density on the failure to lay, we did not estimate egg numbers directly so are unable to assess the effects of nematodes on female reproductive investment. This will undoubtedly be of interest in future studies. Nematodes could potentially cause indirect harm to beetle larvae by competing with them for space or resources, or possibly, by physically interfering with larvae while they consume the carcass. Phoretic nematodes are bacterivores, so direct resource competition with beetle larvae seems unlikely, unless some part of beetle nutrition is also microbial (directly or indirectly) (Wang & Rozen, 2017). Competition for physical space may occur if nematode densities are sufficiently high to prevent larval feeding or access to parts of the carcass. This type of interference could also explain the absence of a trade-off between brood size and larval mass (Figure 3), since much of this effect is driven by the reduction in larval size in smaller broods (<~10 larvae/brood). Direct harm could possibly arise at different stages of development. Eggs could be pierced, something observed by Nicrophorus phoretic mites, Poecilocochus carabi (Beninger, 1993), or otherwise damaged by nematodes. Larvae could also be directly harmed by worms during their growth (Dillman et al., 2012). It is notable that worms are not only transported on the surface of beetles, but are also recovered from within the digestive and reproductive tracts (Figure 1), indicating an ability to invade host tissue (Barbercheck, 2005; Sudhaus, 2008). Internalized worms may obtain nutrients from the larvae or otherwise hinder their growth and development (Sudhaus, 2008). At present, this remains unknown because we lack intermediate samples of larvae themselves and instead have focused on characterizing the transmission route of worms from mature adults through to newly eclosing adults. It will be interesting in future work to sample worm densities in developing larvae to better understand how and when worms inflict their damage. In addition, it will be worthwhile to determine the extent of transmission via dispersing adults on the same carcass. This will be especially important for males who disperse from the carcass before larvae are fully matured, as this second source of nematode transmission may serve to reduce the density of worms on individual larvae.

Although our results make clear that R. regina is common in field-caught beetles and can persist through a complete beetle life cycle, there are important limitations to our study. Most notably, our fitness experiments were carried out in the laboratory in the absence of other species that could either mitigate or exacerbate the harm caused by nematodes. Nicrophorus beetles carry other phoretic species: many different species of mites (Gasperin & Kilner, 2015), possibly other species of nematodes (Khan, 1993; Koneru et al., 2016) and potentially diverse genotypes of individual species that compete with one another for transmission, thereby affecting virulence (Herre, 1995). While there is no evidence of simultaneous carriage of different nematode species, this has been observed in other insect:nematode associations and is possible here, too (Koneru et al., 2016). On the other hand, mites and nematodes always co-occur in Nicrophorus. Wilson and Knollenberg (1987) found that the effects of mites varied from harmful at high densities to neutral or even beneficial at lower densities. They also found that mites reduce the burden of nematodes on eclosing
adults by up to sixfold from ~18,000/individual to ~3,000/individual (Wilson & Knollenberg, 1987). This reduction could have different causes, from direct consumption of nematodes to other types of interference competition. Regardless, their experiments make clear the importance of examining the effects of phoretic species in the context of the entire community. This includes mites and nematodes, but also should include the microbial species that live on and within the beetles and carcass, and also the microbes that are carried within the nematodes, especially because these bacteria may be directly associated with nematode entomopathogenicity (Dillman et al., 2012; Jiménez-Cortés et al., 2016).

Nematodes have a broad continuum of effects on their host species (Kiontke & Sudhaus, 2006), and transitions between levels of harm appear to be widespread and context-dependent even among closely related hosts (Perlman & Jaenike, 2003). Although phoresy is often assumed to be a commensal interaction that benefits worms without harming their hosts, the results of our study suggest that this assumption is incorrect—at least in our model system. We show that the phoretic worms of *N. vespilloides* are detrimental to beetles under laboratory conditions even though they rely on beetles for transmission to new resources. However, this harm may be inevitable and unavoidable if worm transmission requires proliferation. It will be important to determine whether this is similarly true for other phoretic associates of animals. It will also be important to examine interactions between nematodes and *Nicrophorus* in the field, where associations with other phoretic species may modify the effects of nematodes on their hosts.

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AUTHORS’ CONTRIBUTIONS

YW and DR conceived the ideas and designed methodology; YW collected the data; YW and DR analyzed the data and co-wrote the manuscript. Both authors contributed critically to the drafts and gave final approval for publication.

DATA ACCESSIBILITY

Data available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.5q8028q and are also available directly from the senior author.

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REFERENCES

Alizon, S. (2008). Transmission-recovery trade-offs to study parasite evolution. *American Naturalist*, 172, 113–121. https://doi.org/10.1086/589892

Arce, A. N., Smiseth, P. T., & Rozen, D. E. (2013). Antimicrobial secretions and social immunity in larval burying beetles. *Nicrophorus vespilloides*. *Animal Behavior*, 86, 741–745. https://doi.org/10.1016/j.anbehav.2013.07.008

Ashworth, J. R., & Wall, R. (1994). Responses of the sheep blowflies *Lucilia sericata* and *L. cuprina* to odor and the development of semiochemical baits. *Med Vet Entom*, 8, 303–309.

Barbercheck, M. E. (2005). *Insect-parasitic nematodes for the management of soil-dwelling insects*. Entomol Notes, Dep Entomol Pennsylvania State Univ.Available at: http://www.ento.psu.edu/extension/factsheets/nematode.htm (verified 15 March 2010)

Bartlow, A. W., Villa, S. M., Thompson, M. W., & Bush, S. E. (2016). Walk or ride? Phoretic behaviour of amblycercan and ischnoceran lice. *International Journal for Parasitology*, 46, 221–227. https://doi.org/10.1016/j.ijpara.2016.01.003

Beninger, C. (1993). Egg predation by *Poeciloclirrus carabi* (Mesostigmata, Parasitidae) and its effect on reproduction of *Nicrophorus vespilloides* (Coleoptera, Sphiliidae). *Environmental Entomology*, 22, 766–769.

Brown, F. D., D’Anna, I., & Sommier, R. J. (2011). Host-finding behaviour in the nematode *Pristionchus pacificus*. *Proceedings of the Royal Society B-Biological Sciences*, 278, 3260–3269. https://doi.org/10.1098/rspb.2011.0129

Campbell, E. O., & Luong, L. T. (2016). Mite group choices- sex-and size-biased infection in *Drosophila hydei*. *Parasitology*, 143, 787–793. https://doi.org/10.1017/S0031182016000305

Cotter, S. C., & Kilner, R. M. (2010). Sexual division of antibacterial resource defence in breeding burying beetles, *Nicrophorus vespilloides*. *Journal of Animal Ecology*, 79, 35–43.

De Gasperin, O., Duarte, A., & Kilner, R. M. (2015). Interspecific interactions explain variation in the duration of paternal care in the burying beetle. *Animal Behavior*, 109, 199–207. https://doi.org/10.1016/j.anbehav.2015.08.014

De Gasperin, O., & Kilner, R. M. (2015). Friend or foe: Inter-specific interactions and conflicts of interest within the family. *Ecological Entomology*, 40, 787–795. https://doi.org/10.1111/een.12259

Dillman, A. R., Chaston, J. M., Adams, B. J., Ciche, T. A., Goodrich-Blair, H., Stock, S. P., & Sternberg, P. W. (2012). An entomopathogenic nematode by any other name. *PloS Path*, 8, 8–11. https://doi.org/10.1371/journal.ppat.1002527

Donn, S., Griffiths, B. S., Neilson, R., & Daniell, T. J. (2008). DNA extraction from soil nematodes for multi-sample community studies. *Applied Soil Ecology*, 38, 20–26. https://doi.org/10.1016/j.apsoil.2007.08.006

Duarte, A., Cotter, S. C., Reavey, C. E., Ward, R. J. S., De Gasperin, O., & Kilner, R. M. (2016). Social immunity of the family: Parental contributions to a public good modulated by brood size. *Evolutionary Ecology*, 30, 123–135. https://doi.org/10.1007/s10682-015-9806-3

Duarte, A., Welch, M., Swannack, C., Wagner, J., & Kilner, R. M. (2017). Strategies for managing rival bacterial communities: Lessons from burying beetles. *Journal of Animal Ecology*, 87, 414–427. https://doi.org/10.1111/1365-2662.12725

Eggers, A.-K., Reinking, M., & Muller, J. K. (1998). Parental care improves offspring survival and growth in burying beetles. *Animal Behavior*, 55, 97–107. https://doi.org/10.1006/anbe.1997.0588

Floyd, R. M., Rogers, A. D., Lambshde, P. J. D., & Smith, C. R. (2005). Nematode-specific PCR primers for the 185 small subunit rRNA gene. *Molecular Ecology Notes*, 5, 611–612. https://doi.org/10.1111/j.1471-8286.2005.01009.x

Fronhofer, E. A., Sperr, E. B., Kreis, A., Ayasse, M., Poethke, H. J., & Tscharpa, M. (2013). Picky hitch-hikers: Vector choice
(Coleoptera: Histeridae: Haeteriinae) with an exceptional mechanism of phoresy. BMC Zool, 2, 3.

Wang, Y., & Rozen, D. E. (2017). Gut microbiota colonization and transmission in the burying beetle Nicrophorus vespilloides throughout development. Applied and Environment Microbiology, 83, e03250–e3316. https://doi.org/10.1128/AEM.03250-16

Wang, Y., & Rozen, D. (2017). Gut microbiota in the burying beetle, Nicrophorus vespilloides, provide colonization resistance against larval bacterial pathogens. Ecology and Evolution, 8, 1646–1654.

Wilson, D. S., & Knollenberg, W. G. (1987). Adaptive indirect effects: The fitness of burying beetles with and without their phoretic mites. Evolutionary Ecology, 1, 139–159. https://doi.org/10.1007/BF02067397

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