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Citation for published version:
Rynkiewicz, EC, Clerc, M, Babayan, SA & Pedersen, AB 2019, 'Variation in local and systemic pro-inflammatory immune markers of wild wood mice after anthelminthic treatment', Integrative and comparative biology. https://doi.org/10.1093/icb/icz136

Digital Object Identifier (DOI):
10.1093/icb/icz136

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Integrative and comparative biology

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SYMPOSIUM

Variation in Local and Systemic Pro-Inflammatory Immune Markers of Wild Wood Mice after Anthelmintic Treatment

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From the symposium “The scale of sickness: how immune variation across space and species affects infectious disease dynamics” presented at the annual meeting of the Society for Integrative and Comparative Biology, January 3–7, 2019 at Tampa, Florida.

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Synopsis

The immune system represents a host’s main defense against infection to parasites and pathogens. In the wild, a host’s response to immune challenges can vary due to physiological condition, demography (age, sex), and coinfection by other parasites or pathogens. These sources of variation, which are intrinsic to natural populations, can significantly impact the strength and type of immune responses elicited after parasite exposure and infection. Importantly, but often neglected, a host’s immune response can also vary within the individual, across tissues and between local and systemic scales. Consequently, how a host responds at each scale may impact its susceptibility to concurrent and subsequent infections. Here we analyzed how characteristics of hosts and their parasite infections drive variation in the pro-inflammatory immune response in wild wood mice (Apodemus sylvaticus) at both the local and systemic scale by experimentally manipulating within-host parasite communities through anthelmintic drug treatment. We measured concentrations of the pro-inflammatory cytokine tumor necrosis factor alpha (TNF-α) produced in vitro in response to a panel of toll-like receptor agonists at the local (mesenteric lymph nodes [MLNs]) and systemic (spleen) scales of individuals naturally infected with two gastrointestinal parasites, the nematode Heligmosomoides polygyrus and the protozoan Eimeria hungaryensis. Anthelmintic-treated mice had a 20-fold lower worm burden compared to control mice, as well as a four-fold higher intensity of the non-drug targeted parasite E. hungaryensis. Anthelmintic treatment differentially impacted levels of TNF-α expression in males and females at the systemic and local scales, with treated males producing higher, and treated females lower, levels of TNF-α, compared to control mice. Also, TNF-α was affected by host age, at the local scale, with MLN cells of young, treated mice producing higher levels of TNF-α than those of old, treated mice. Using complementary, but distinct, measures of inflammation measured across within-host scales allowed us to better assess the wood mouse immune response to changes in parasite infection dynamics after anthelmintic treatment. This same approach could be used to understand helminth infections and responses to parasite control measures in other systems in order to gain a broader view of how variation impacts the immune response.

Introduction

Within-host parasite communities can be extremely diverse, with one individual being host to many parasite species simultaneously or sequentially over the course of its lifetime. This is especially true for individuals living in the wild, which face frequent and repeated challenges from both microparasites (e.g., viruses, bacteria, and protozoa) and macroparasites (e.g., helminths and ectoparasites). The composition of these within-host parasite communities can have significant impacts on individual host health, parasite transmission dynamics, and the epidemiology of each parasite species (Poulin 1996; Pedersen and Fenton 2007; Lafferty et al. 2008; Telfer et al. 2008;
Johnson et al. 2015; Rynkiewicz et al. 2015). However, while we often consider the host individual as the appropriate unit for investigating within-host processes, for parasites, a host represents a diverse ecosystem with many different habitats, each of which can be infected by different parasites (Griffiths et al. 2014; Restif and Graham 2015; Rynkiewicz et al. 2015). Therefore, a parasite’s preferred niche can be functionally determined by the properties of the tissues it infects and by the type of immune response it elicits (Graham et al. 2007; Pedersen and Fenton 2007; Lello and Hussell 2008). Correspondingly, the host immune system has evolved different mechanisms to respond to infection by a diverse set of parasites (Hajnicka et al. 2005; Baucom and de Roode 2011; Maizels et al. 2012; Parker et al. 2017). The host’s response to parasite infection can be measured at multiple sites or scales, including a local response at the site of infection which may focus on tissue repair or controlling initial parasite spread and damage, and a systemic response across the whole organism where circulating immune factors in blood and the spleen coordinate a more general response to infection. While local immune responses impact host susceptibility and infection success of parasites that infect the same location, systemic responses impact susceptibility to parasites coinfesting other tissues within the host (Mwatha et al. 1998; Munoz et al. 2005; Semenyya et al. 2012; Zavala et al. 2018). Therefore, investigating parasite–host and parasite–parasite interactions at multiple within-host scales will improve our ability to measure and predict parasite spread as well as quantify variation among hosts in how they respond to infection. Coinfection is very common in natural host populations, and in turn can lead to within-host parasite interactions (Poulin 1996, 2001; Abu-Raddad et al. 2006; Seabloom et al. 2015). These can be direct via competition for habitat, or indirect through competition for a nutritional resource or via the host’s immune system (Pedersen and Fenton 2007; Graham 2008; Griffiths et al. 2014; Budischak et al. 2015; Griffiths et al. 2015). An analysis of human coinfection networks showed that coinfecting parasites most commonly interact though shared host resources and a common infection site (Griffiths et al. 2014). However, there was also evidence of immune-mediated parasite interactions, where one parasite can negatively or positively impact the infection success or intensity of another through a shared immune response. When parasites interact via the host immune system, it is important to consider at which within-host scales the interaction is occurring (e.g., are these parasites infecting the same or different tissues) to assess their impact on host health or parasite transmission. For example, parasites that elicit a strong systemic immune response can potentially impact host susceptibility to other coinfesting parasite species, even if they infect different within-host habitats. This type of interaction has been shown in wild African Buffalo, where reducing gastrointestinal nematode burdens using anthelminthic treatment significantly increased individual hosts’ odds of survival after bovine tuberculosis infection, due to altered systemic inflammation (Ezenwa and Jolles 2015). Consequently, the scale at which we measure metrics of the immune response will influence our conclusions about how individuals respond to infection, and how the variation in these responses may impact parasite coinfection and transmission dynamics.

How a host responds to infection can also vary depending on demographic/physiological characteristics such as age, sex, or reproductive condition, which can impact resource allocation to local and systemic immune responses (French et al. 2007; Pedersen and Greives 2008; Vandegrift et al. 2008; Buehler et al. 2009). Studies in Soay sheep and tree swallows, for example, have shown significant changes in a host’s immune response with increasing host age, with older hosts in most cases having higher levels of inflammatory markers (Palacios et al. 2007; Nussey et al. 2012; Cevenini et al. 2013; Babayan et al. 2018b). Furthermore, there are key immunological differences between the sexes, with males often being less immune responsive than females. These differences are often attributed to steroid hormones, such as testosterone, whose immunosuppressive effects can also vary across seasons (Demas and Nelson 1998; Lehmer et al. 2010; Pap et al. 2010). There are also immunological differences within the sexes, dependent on life-history stage. For example, pregnant females often have lower levels of inflammatory markers than non-reproductively active females (Martin et al. 2008; Babayan et al. 2018). These cases suggest the physiological trade-offs between sexually-selected traits and the immune response are an evolutionarily adaptive mechanism (Folstad and Karter 1992; Zuk 1996; McKeen and Nunney 2005; Stoehr and Kokko 2006), emphasizing the importance of studying the impacts of these trade-offs in evolutionarily- and ecologically-relevant contexts.

One key immunological process that can be measured at both the local and systemic within-host scales is inflammation, broadly defined as the recruitment of immune cells and signaling molecules...
to the site of an infection (Sompayrac 2012). Inflammation occurs early in an infection, involves both the innate and adaptive arms of the immune system, initiates tissue repair or containment, and/or clearance of an infectious agent. Inflammation is maintained by positive feedbacks, which are then disrupted or deactivated by subsequent regulatory or anti-inflammatory signals upon clearance of infection (Sears et al. 2011). One key cytokine involved in controlling inflammation is tumor necrosis factor alpha (TNF-α), a pro-inflammatory cytokine which can activate both innate and adaptive immune responses (Vassalli 1992; Pasparakis et al. 1996; Graham et al. 2007; Bradley and Jackson 2008; Tumanov et al. 2010). In addition, it has been suggested that TNF-α levels indicate a host’s ability to respond to a broad range of parasites, such as worms, bacteria, viruses, or other microbial pathogens (Smith and Ovington 1996; Goff et al. 2002; Sjöwall et al. 2005; Zavala et al. 2018).

Here, we investigated the interactions between TNF-α as a marker of inflammation and parasite infection and coinfection in wild wood mice (Apodemus sylvaticus). Wood mice are commonly infected with a diverse community of parasites (Bown et al. 2006; Behnke et al. 2009; Knowles et al. 2013), including the gastrointestinal nematode Heligmosomoides polygyrus, which is closely related to H. bakeri, an established laboratory model for human nematode infections (Behnke et al. 2005; Maizels et al. 2004; Reynolds et al. 2012). Heligmosomoides bakeri is known to elicit strong immunomodulatory effects on the host’s immune system and skews the host response from a pro-inflammatory (Th1-type) immune profile to an anti-inflammatory (Th2/Treg-type) immune profile (Maizels et al. 2009; Maizels et al. 2012). Through the use of anthelmintic treatment, we have shown that in the wild, H. polygyrus negatively interacts with another common gastrointestinal parasite of wood mice, the coccidial Eimeria hungaryensis (Fenton et al. 2014; Clerc et al. 2019b), and that this strong negative effect of H. polygyrus on E. hungaryensis infection burden is localized to the small intestine (Knowles et al. 2013). However, we currently lack an understanding of how interactions between H. polygyrus and E. hungaryensis impact immune responses outside of the local site of infection. In addition, studies that experimentally investigate within-host variation in the immune response across different scales are rare, especially in wild populations. Researchers often generalize host immune reactivity from systemic measures, therefore there may be significant implications for understanding and predicting parasite community interactions at the host individual and population scales given the possible differences in immune activity between the local and systemic scales.

We used the production of the pro-inflammatory cytokine TNF-α in response to a panel of toll-like receptor (TLR) agonists in mouse mesenteric lymph node (MLN) cells as a proxy for the local inflammatory response surrounding the gastrointestinal tract, and in spleen cells as a proxy for the systemic inflammatory response, to elucidate how each immune response may mediate within-host interactions between H. polygyrus and E. hungaryensis. Furthermore, we used anthelmintic treatment to assess the effects of H. polygyrus removal/reduction on inflammation at the local versus systemic scales. We predicted that TNF-α production would differ between local and systemic sites due to different immune cell populations being involved and their proximity to the site of infection (Vassalli 1992). Because H. polygyrus modulates the host immune response toward an anti-inflammatory immune profile (Maizels et al. 2012; Wammes et al. 2016), we further hypothesized that anthelmintic-treated mice would have increased TNF-α production because of the reduced worm burden and subsequent reduced immunomodulatory effects. Due to the immunosuppressive effects of testosterone (Saino et al. 1995; Zuk 1996; Muehlenbein and Bribiescas 2005; Nunn et al. 2009), we also predicted that male mice would have lower TNF-α levels than females. Lastly, we expected older hosts to have higher concentrations of TNF-α, in line with the concept of immunosenescence (Cevenini et al. 2013). Even though TNF-α is not thought to be directly protective against H. polygyrus, it may be associated with the inflammatory immune response to E. hungaryensis infection (Smith et al. 1995; Pakandl et al. 2008; Maizels et al. 2009; Maizels et al. 2012).

**Methods**

**Field sampling**

We conducted two field experiments in wild wood mouse populations in Callendar Woods, Falkirk, Scotland (55.99°N, 3.77°W). We performed the experiment over two trapping sessions, each session lasting 8 weeks (Session 1: October to November 2014 and Session 2: June to July 2015). In 2014, the experiment consisted of two trapping grids in which two Sherman live traps (H. B. Sherman 2 × 2.5 × 6.5-inch folding trap, Tallahassee, FL, USA) were placed every 10 m in a 130 × 80 m grid (total 256 traps). In 2015, the large grid was split...
into two smaller grids that were separated by ~10 m, and an additional third grid was added ~50 m away from the other two grids. All three grids were 70 × 70 m in size and consisted of a total of 294 traps, leading to similar trapping effort in both years. Traps were baited with cotton bedding, mixed seeds, dried mealworms, and a piece of carrot in the late afternoon before each trapping night. The following morning, traps were checked for the presence of animals and all animals were processed and released at the site of capture. For both of the trapping sessions, we trapped three consecutive nights per week for a total of 24 trap-nights per session (Home Office Project License 60/4572).

At first capture, each wood mouse received a unique passive induced transponder tag injected subcutaneously in the scruff (AVID FriendChip), which allowed individual identification of mice at subsequent captures. Furthermore, all mice at first capture were randomly selected to receive either a single oral dose of a combination of anthelmintic drugs: ivermectin (which removes adult worms) at 100 mg/kg and pyrantril palmoate (which removes larval worms) at 9.4 mg/kg (Wahid et al. 1989; Wahid and Behnke 1993; Babayan et al. 2018a; Clerc et al. 2019a), or an equal volume of water as a control. For each mouse at each capture, we recorded the mouse’s trap location, sex, reproductive status (females: perforated vagina, pregnant, or lactating, males: non-reproductive, testes descended, or scrotal), body weight, and length. Fecal samples were collected from pre-sterilized traps each time an animal was captured, and then stored in 10% formalin at 4°C until analysis (Dryden et al. 2005). Heligmosomoides polygyrus and E. hungaryensis infection (infected/uninfected) and burdens (number of eggs/oocysts per 1 g of feces OPG/EPG) were determined by counting parasite transmission stages (eggs/oocysts) after salt floatation using a microscope at either 10× or 40× magnification (see Knowles et al. 2013).

Mice were sacrificed ~2 weeks after initial capture and treatment (anthelmintic or control), as previous work with wood mice has shown that anthelmintic drug treatments reduce H. polygyrus burdens for only about 10–16 days (Knowles et al. 2013; Clerc et al. 2019a). Spleen and MLNs were collected and stored in sterile RPMI media containing 1% Pen/Strep antibiotics for use in the TNF-α assay. We also collected the digestive tract of each mouse, separated into small intestine, cecum, and colon, to assess burdens of adult gastrointestinal parasites via dissection. The eyes of each mouse were also collected to use dried eye lens weight (milligrams) as a measurement of host age. This metric has been shown to be a reliable method for quantifying host age on a continuous scale, where older hosts have heavier eye lenses compared to younger hosts (Morris 1972; Hardy et al. 1983). Both eyes were stored in 10% formalin at 4°C for at least 4 weeks. After separating the eye lenses from the surrounding tissue, they were dried at 56°C overnight and weighed in pairs to the nearest milligram.

**Immunological methods**

To assess the ability of an individual mouse to respond to infection at both the systemic and local levels, we measured pro-inflammatory cytokine TNF-α production by spleen (systemic) and MLN (local) cells *in vitro*. Cells were stimulated with a panel of TLR agonists known to induce production of TNF-α (Akira et al. 2006) including in wild mice (Bradley and Jackson 2008). The agonists used were zymosan (cell surface fungal ligand, TLR2), endosomal oligonucleotide ODN2006 (ODN, TLR9), cell surface *Escherichia coli* K12 lipopolysaccharide (LPS, TLR4), and heat-killed *Listeria monocytogenes* (HKLM, TLR2; all acquired from InvivoGen, San Diego, CA, USA). TNF-α concentration was then measured by sandwich Enzyme-Linked Immunosorbent Assay (ELISA), a previously described, successful method in wild wood mice (Jackson et al. 2009).

Spleen and MLN samples were brought into the lab for cell culture on the same day as collection (tissues in media were kept on ice during transport until used in cell culture). Under sterile conditions, tissues were passed through 70 μm nylon mesh into RPMI media containing heat-inactivated fetal calf serum (10% final volume) and 1% Pen/Strep antibiotics. Cell suspensions were spun down at 1200 rpm, supernatant discarded, and resuspended in fresh media (5 mL for spleen cells, 1 mL for MLN cells). Before resuspension, spleen cells were incubated with red blood cell (RBC) lysis buffer for 3 min, inactivated with 10 mL of RPMI media. This suspension was spun down at 1200 rpm, supernatant discarded, and cells resuspended with 5 mL media. If all RBCs were not lysed, this previous step was repeated a second time. Cell concentrations for each sample were then counted (Cellometer Auto T4 Bright Field Cell Counter, Nexcelom Bioscience) and resuspended to 1 × 10^7 cells/mL for spleen cell suspensions, 5 × 10^6 cells/mL for MLN cell suspensions. Different final cell concentrations were used for these two tissue types due to fewer cells available from MLN than spleens; this difference was
accounted for in the statistical analyses by analyzing results from each tissue type separately (see below).

Cells from the spleen and MLN were then plated in 96-round bottom well plates for stimulation. Final volumes for samples from each individual mouse varied and low volume samples from individual mice did not receive all agonists. Controls and TLR agonists were run in triplicate, with three wells of cells receiving either an equal volume of media or agonist suspension (100 μL of cell culture, 100 μL of control or agonist added to each well). Concentrations of each agonist were as follows: Zymosan: 20 μg/mL, ODN: 10 μg/mL, LPS: 6 μg/mL, HKLM: 6.0e7 μg/mL (from Jackson et al. 2009). Cell cultures were incubated for 24 h at 37°C (5% CO2). Subsequently, plates were spun at 1200 rpm for 2 min and supernatant harvested for use in TNF-α ELISA.

Concentrations of TNF-α produced during culture were measured using a mouse sandwich ELISA according to the manufacturer’s instructions (DuoSet ELISA, Mouse TNF-α, R&D Systems). Pooled cell cultures from each mouse for each agonist were used in the ELISA to reduce variation among wells. Each individual mouse and agonist combination was run in triplicate. Plate absorbance was read at 450 nm, the peak wavelength for the plate’s indicator of cytokine concentration, and well absorbance converted to TNF-α concentration (pg/mL) based on the standard curve on each plate. As stated in Jackson et al. (2009), it is likely that affinity of A. sylvaticus to the TNF-α antibodies in these ELISA kits differs from Mus musculus, but the results should still accurately capture variation in this marker of inflammation among individuals of this species.

Statistical analysis

All analyses were performed in R (R Core Team 2018), version 3.5.1. Data on parasite infection and burden and in vitro inflammatory cytokine concentrations (pg/mL) were analyzed using general linear mixed models (GLMM) in the glmmADMB package. Initial data on parasite infection burden were determined measuring EPG/OPG of H. polygyrus or E. hungaryensis from samples collected at first capture, prior to the drug (or control) dosing (“Methods” section, see Clerc et al. 2018). Final measures of parasite infection were (1) final adult H. polygyrus burdens within the gut (number of adult worms) and (2) final E. hungaryensis infection (presence [1]/absence [0] of oocysts in feces). Analyses were performed using mean TNF-α concentrations for each individual and TLR agonist combination. Cytokine concentrations were log-transformed (log10 [1+TNF-α concentration]) to improve normality (Gaussian distribution in GLMM).

We measured the production of TNF-α from spleenocytes as an indicator of the systemic immune response and MLN cells as an indicator of the local immune response. There was variation in mean concentration of TNF-α produced in vitro in response to stimulation with different agonists, however, we analyzed TNF-α concentrations from all control and stimulated wells together, as sample size did not permit analyzing data from each agonist individually or inclusion of the agonist type as predictor variable in models. While this did not allow us to evaluate how a host may respond to a specific kind of infection (via the response to a single agonist), it did allow us to capture variation in TNF-α among individuals in response to anthelmintic treatment and simulate the complex innate immune stimuli that could be experienced by these mice (Jackson et al. 2009).

We included the following variables in each model as possible drivers of variation in the marker of inflammation (TNF-α production in response to the panel of agonists): anthelmintic treatment (treated/control), total H. polygyrus adult worm burden, E. hungaryensis infection (0/1), host sex (M/F), reproductive condition as determined by the presence of descended testes in males, and females that are visible pregnant or lactating (0/1), eye lens mass (proxy for age), and relevant biological interactions between treatment and other variables (Supplementary Table S1). We also separately analyzed initial H. polygyrus and E. hungaryensis infection burdens (EPG/OPG from first capture) as possible predictors of final TNF-α production (Supplementary Table S1). We performed model reduction of all full models using maximum likelihood ratios to compare the full model to reduced models using the “anova” function in the glmmADMB package. Interactions followed by main effects were not retained in the final model if the maximum likelihood ratio test had a P > 0.05, indicating that these terms did not significantly improve model fit. For TNF-α concentrations, random effects of mouse ID nested within ELISA plate, and TLR agonist were included in each model, to account for multiple measures for each individual mouse and variation in stimulation between TLR agonists. A summary of the final GLMMs for parasite burdens and TNF-α concentrations can be found in Table 1.
**Results**

**Impact of anthelmintic treatment and host characteristics on the gastrointestinal parasite community**

A total of 148 wood mice were captured during the two replicate field experiments (2014 and 2015). The overall recapture rate (percentage of hosts recaptured at least once) was over 90% for this population; 60% of control hosts, and 57% of drug-treated hosts were captured during the 12-18 day period post-treatment. TNF-α concentrations were measured from 19 mice in Fall 2014 and 30 mice in Summer 2015 (for a summary of demographics and sample size, see Supplementary Table S2). Wood mice were commonly infected with gastrointestinal parasites: 86% (42/49) with the nematode *H. polygyrus*, while 57% (29/49) were infected with the coccidian protozoan *E. hungaryensis*; 14% (7/49) hosts were coinfected when terminally sampled. There were no initial differences in *H. polygyrus* or *E. hungaryensis* infection intensity (EPG/OPG) between mice randomly allocated to anthelmintic treatment or control at first capture (first-capture parasite EPG/OPG: *H. polygyrus*: z = 0.47, P = 0.64; *E. hungaryensis*: z = −0.26, P = 0.796). Male mice, however, showed significantly higher initial *H. polygyrus* egg shedding than females at first capture (z = 3.17, P = 0.0015, Fig. 1A), but no such sex difference was found in initial *E. hungaryensis* oocyst shedding (Fig. 1B).

### Table 1 Summary of final GLMMs analyzing parasite burdens and TNF-α production

| Hypothesis                                                                 | Final model                                                                 | Variables                                      | z-value | P-value |
|----------------------------------------------------------------------------|----------------------------------------------------------------------------|-----------------------------------------------|---------|---------|
| Males will have higher initial *H. polygyrus* EPG than females.            | Initial *H. polygyrus* EPG ~ Trt + Sex + Trt × Sex                        | Treatment                                     | −0.17   | 0.8663  |
| Reproductively active hosts will have higher parasite burdens compared to non-reproductively active hosts. |                                                                              |                                               |         |         |
|drug treatment will lower *H. polygyrus* burden, increase *E. hungaryensis* oocyst shedding. | Final *H. polygyrus* burden ~ Trt + Reproductive Condition | Treatment                                      | −5.34   | <0.0001*** |
| Reproductively active hosts will have higher parasite burdens compared to non-reproductively active hosts. |                                                                              |                                               |         |         |
| Treated males will have higher TNF-α concentrations than control males.    | log10 (1 + TNF Concentration Spleen) ~ Trt + Sex + Final *H. polygyrus* burden + Trt × Sex | Final *H. polygyrus* burden                    | −1.59   | 0.111   |
| Old hosts will have higher TNF-α concentrations compared to young hosts.   | log10 (1 + TNF Concentration MLN) ~ Sex + Reproductive Condition + Age (eye lens mass) + Treatment + Final *H. polygyrus* burden + Trt × Age + Trt × Final *H. polygyrus* burden | Treatment                                      | 2.48    | 0.013*  |
| Reproductive condition will be lower in older hosts compared to young hosts. |                                                                              |                                               | 2.31    | 0.021*  |
| Age (eye lens mass) will be lower in older hosts compared to younger hosts. |                                                                              |                                               | −0.89   | 0.379   |
| Adult males will have higher TNF-α concentrations than adult females.      |                                                                              |                                               | −0.96   | 0.338   |
| Co-infected hosts will have higher TNF-α concentrations than single infections. |                                                                              |                                               | 1.46    | 0.144   |
| Adult males will have higher TNF-α concentrations than adult females.      |                                                                              |                                               | −2.72   | 0.006** |
| Co-infected hosts will have higher TNF-α concentrations than single infections. |                                                                              |                                               | 3.4     | 0.0006** |

Results are organized according to hypotheses presented and if the results support these hypotheses. Global (full) models, including main and random effects, are presented in Supplementary Table S1.
We found that anthelmintic treatment significantly reduced *H. polygyrus* adult worm counts when hosts were captured 2 weeks post-treatment ($z = -5.34, P < 0.0001$, Fig. 2A). While no mice were completely cleared of *H. polygyrus* worms, treated mice had a nearly 20-fold decrease in worm burdens compared to control mice. Anthelmintic treated mice also had significantly higher *E. hungaryensis* oocyst shedding (OPG, $z = 2.04, P = 0.042$, Fig. 2B), with OPGs being four-fold higher than control mice, which is consistent with our previous results (Knowles et al. 2013; Clerc et al. 2019a, 2019b). Even though we had limited statistical power to test for the effect of reproductive status and sex on parasite burdens following anthelmintic treatment (only one non-reproductive male in the dataset, see Supplementary Table S2), we found a trend toward a positive effect of reproductive activity on worm burdens. The *H. polygyrus* burdens of reproductively active females were on average double that of non-reproductive females, while reproductively active males had four times more adult *H. polygyrus* worms in their gut compared to non-reproductive mice ($z = 1.76, P = 0.078$, Fig. 3A), however sex as a main effect was not retained in the final model. There were no statistically significant differences in *E. hungaryensis* burden between reproductive and non-reproductive mice (Fig. 3B).

**TNF-α production**

Cell cultures stimulated with TLR agonists did produce more TNF-α than control cultures ($z = 10.78, P < 0.0001$; control: mean log10 [1 + TNF-α concentration] 2.05 ± 0.170, stimulated: mean log10 [1 + TNF-α concentration] 4.05 ± 0.099; Supplementary Fig. S1).

TNF-α production was significantly different between systemic (spleen) and the local (MLN) scales, with systemic TNF-α levels being consistently higher than those measured locally (Spleen: $4.2 ± 0.14, MLN: 3.1 ± 0.12, z = 5.63, P < 0.0001$; Fig. 4), likely due to the difference in TNF-α producing cell concentrations between tissues. While there was a correlation between mean TNF-α concentrations measured from spleen and MLN cells, only 7% of the variation in cytokine concentrations in one tissue was explained by concentrations in the other ($P = 0.068$, adjusted $R^2 = 0.072$; Supplementary Fig. S2). Therefore, we analyzed local and systemic TNF-α production separately in our subsequent analyses. In addition, we found no significant difference in TNF-α concentrations between field experimental replicates (spleen: $z = 0.74, P = 0.46$, MLN: $z = -0.01, P = 0.99$), so data were pooled from both years for analysis.

At the systemic level, we found a significant effect of anthelmintic treatment on TNF-α production ($z = -2.55, P = 0.011$) and an interaction between treatment and host sex (sex × treatment, $z = 1.82, P = 0.069$, Fig. 4B); systemic TNF-α in treated males was ~50% higher than those from control males. The opposite pattern appeared in females, where treated females showed ~50% lower systemic TNF-α concentrations than control females.

At the local scale, we also found a main effect of sex on local TNF-α production, with males producing ~25% more TNF-α than females ($z = 2.31, P = 0.021$; Fig. 4A). At the local scale, we found a significant interaction between anthelmintic treatment and host age (age × treatment, $z = -2.72, P = 0.006$, Fig. 5); lower production of local TNF-α was found in older, treated mice, while there was no effect of age on local TNF-α production in control mice. These interactions between host characteristics and anthelmintic treatment suggest that hosts are impacted differently by the reduction in *H. polygyrus* burden depending on their age or sex (see Clerc et al. 2019a). There was no relationship between pre-treatment *H. polygyrus* or *E. hungaryensis* burdens and post-treatment TNF-α production at either scale.

**Discussion**

In this study, we used targeted drug treatments to reduce *H. polygyrus* burdens in wild, demographically heterogeneous wood mice to determine how...
single and multiple infections with gastrointestinal parasites impacted the host immune response at two within-host scales. We found that anthelmintic treatment reduced burdens of the target parasite, *H. polygyrus*, by 20-fold 14 days after treatment (Fig. 2A). This reduction in *H. polygyrus* burden in turn led to a four-fold increase in oocyst shedding of the non-target coccidian parasite *E. hungaryensis* (Fig. 2B). Furthermore, using laboratory mouse reagents, we successfully measured levels of TNF-α production in spleen and MLN cell cultures, allowing us to use the levels of this important pro-inflammatory cytokine as a marker of inflammation and immune reactivity at different scales. We found that hosts responded differently to anthelmintic treatment depending on demographic factors (age and sex) and importantly, that the relationships between TNF-α production and infection status or demographics differed between the systemic (spleen) and local (MLN) scale. These results have implications for predicting the outcome of parasite coinfection, host health, and parasite transmission dynamics.

Perhaps more importantly, however, they increase our understanding of how host demography affects variation in immune markers, and the immune response more generally, depending on the scale at which they are measured.

Overall, we found that male mice had higher initial *H. polygyrus* infection burdens, as measured by egg shedding. Both local and systemic in vitro production of TNF-α also varied by host sex and anthelmintic treatment, where males had higher levels of this marker of inflammation than females. In addition, there was an interaction between host sex and drug treatment on systemic inflammation, where treated males showed higher systemic TNF-α production compared to control mice, while systemic TNF-α was lower in treated female mice. This was counter to our predictions, which were that males would demonstrate less immune activation than females, in line with the immunocompetence handicap hypothesis (Folstad and Karter 1992). However, a recent meta-analysis found that TNF-α was one of only a few immune markers significantly increased in
male mammals compared to females (Kelly et al. 2018). Our results suggest that males may maintain higher production of inflammatory cytokines following removal of adult *H. polygyrus* worms, which are known to stimulate an anti-inflammatory immune response, potentially leading to increased protection from subsequent infections (Lindstrom et al. 2004; Rynkiewicz et al. 2013; Parker et al. 2017). However, further research is needed to elucidate how this may impact costs of immunity, especially since we were unable to measure the immune reactivity before anthelmintic treatment, as the assay requires culling of animals.

We found that local TNF-α production varied with host age in response to anthelmintic treatment, with older, treated mice having lower TNF-α production in the MLN than younger treated mice, however, there was no effect of age on local TNF-α concentrations among untreated hosts (Fig. 5). In contrast, we previously found that *H. polygyrus*-specific IgG1 antibody titers increased with host age, possibly in response to repeated exposure throughout a host’s lifetime (Clerc et al. 2019a). We predicted that older hosts would have higher levels of pro-inflammatory cytokines following anthelmintic treatment, as older hosts often have higher parasite burdens and/or richness compared to younger hosts (Poulin 1996), as well as often experience chronic elevation of inflammatory markers due to immunosenescence (Martin et al. 2006; Cevenini et al. 2013). While the latter has been shown in wild populations of Soay sheep (Nussey et al. 2009), the processes related to ageing have not been extensively studied in wild mice, and we do not know at what age such effects would become detectable. Interestingly, we only saw variation in our marker of inflammation across different ages after anthelmintic treatment and subsequent *H. polygyrus* burden reduction. Chronic helminth infection may mask the effects of ageing on the wood mouse immune system, therefore making it difficult to see changes in immune strategy with age without removing this dominant gut parasite. The results presented here, in conjunction with those of Clerc et al. (2019a), show how measuring markers of host immunity in response to an experimental manipulation is a robust method for investigating the ecological and evolutionary implications of parasite infection.

The finite amount of resources available to an individual can lead to trade-offs between many important functions, such as growth, reproduction, and the immune response. Often, these trade-offs are context-dependent, based on season, food availability or nutritional content, reproductive state, age or parasite coinfection (Nelson 2004; French et al. 2007; Pedersen and Greives 2008). Several studies have shown that reproduction can redirect host resources away from the immune system in both males and females, via hormonal changes or seasonal changes in physiology (Nelson et al. 2002; Greenman et al. 2005). In our data, there was a slight trend toward reproductively active mice having higher final *H. polygyrus* burdens compared to non-reproductive
mice, which also depended on anthelmintic treatment status (Fig. 3). Males also showed increased levels of both local and systemic TNF-α production compared to females, which was again impacted by anthelmintic treatment (Fig. 4). This is consistent with what has been observed in captive zebra finches given an experimental immune challenge, where males always showed higher induced inflammation compared to females, even though both sexes had lower general inflammation when breeding compared to a non-breeding state (Love et al. 2008). Male wood mice had significantly higher initial H. polygyrus burdens compared to females, and also shed fewer E. hungaryensis oocysts. Recently it has also been found that age, H. polygyrus burden before anthelmintic treatment, and E. hungaryensis co-infection had significant effects on H. polygyrus specific antibody responses in this population of wild wood mice (Clerc et al. 2018, 2019a), so a combination of physiological- and parasite-driven challenges may be impacting the pro-inflammatory marker (TNF-α) measured here. Low sample sizes hindered our ability to analyze the effects of reproductive condition on local and systemic inflammation, but this is clearly a physiological process that can have significant impacts on allocation of within-host resources and immune response and is an avenue for future research.

By measuring the same marker of inflammation at two different within-host scales, we found that reducing H. polygyrus burden through anthelmintic treatment affected the host immune response differently at the local versus systemic scales. Many studies have found that gastrointestinal helminths can have significant effects on a host’s immune profile at the systemic scale (spleen/blood) by shifting the immune response toward an anti-inflammatory Th2/Treg type profile (McKay 2006; Maizels et al. 2009; Allen and Maizels 2011). However, how a host responds to the parasite at the site of infection, in our case the gastrointestinal tract, is also important for understanding the interaction between host and parasite and possible effects on locally-co-infecting parasites. Local inflammation has been measured by proxy using the MLNs surrounding the gut (Eberhardson et al. 2008), however measuring immune responsiveness in these tissues is difficult in rodents as it is a terminal procedure (Munoz et al. 2005). There was a correlation between the TNF-α concentrations produced from cells of each tissue within a host, but very little variation was explained ($R^2 = 0.07$), so it seems likely that the host characteristics we used in our analyses are more informative in explaining the local and systemic production of this pro-inflammatory cytokine.

TNF-α production by spleen and MLN cells varied significantly due to different aspects of host demography as well as anthelmintic treatment, as was seen by the impacts of age and treatment on local TNF-α concentrations but not systemic concentrations. These complementary, but distinct, measures of the pro-inflammatory response measured at a local and systemic scale allow us to better assess the wood mouse immune response to anthelmintic treatment, and perhaps apply this same approach to understand helminth infections and responses to parasite control measures in other systems. These results also emphasize the importance of rigorously studying the immune system in wild, demographically diverse hosts, to truly understand the ecological and evolutionary context of how hosts respond to immune challenges. In conclusion, considering a host’s immune system as uniform when studying within-host parasite interactions does not reflect the reality of individuals living in natural environments. Further research into utilizing the approach of measuring markers of immunity at multiple within-host scales will further aid in better understanding parasite-host interactions in the wild.

**Acknowledgments**

The authors would like to thank Godefroy Devevey, Aimeric Blaud, Romane Allanson, Paulina Cortes, Alex Garcia, Jillian Gordan, Ami Patrick, Laura Gordon, Nicola Dempster, and Sarah King for their assistance and support with sample collection and processing; Amy Buck and Divya Malik for guidance and support with immunological methods; and the anonymous reviewers for comments on previous versions of this manuscript.

**Funding**

The authors also thank the Division of Ecosystems and Disease Ecology, Division of Comparative Endocrinology, Division of Animal Behavior, and Division of Ecology and Evolution of the Society for Integrative and Comparative Biology as well as the Macrocology of Infectious Disease Research Coordination Network funded by the National Science Foundation (NSF DEB 1316223) for supporting the symposium “The Scale of Sickness: How Immune Variation Across Space and Species Affects Infectious Disease Dynamics” financially. This work was funded by the National Science Foundation (Postdoctoral Research Fellowship in Biology (DBI-1306608)) to E.C.R., a University of Edinburgh Torrance Bequest Scholarship to M.C., Welcome Trust ISSF grant [097821/Z/11/Z] to SB, a targeted...
Institute of Biodiversity, Animal Health & Comparative Medicine Research Fellowship to SB, a Wellcome Trust Strategic Grant (Centre for Immunity, Infection, and Evolution; 095831) to A.B.P., a NERC standard grant (NE/I026367/1) to A.B.P. and a University of Edinburgh Chancellors Fellowship to A.B.P.

Supplementary data

Supplementary data available at ICB online.

References

Abu-Raddad LJ, Patnaik P, Kublin JG. 2006. Dual infection with HIV and malaria fuels the spread of both diseases in sub-Saharan Africa. Science 314:1603–6.

Akira S, Uematsu S, Takeuchi O. 2006. Pathogen recognition and innate immunity. Cell 124:783–801.

Allen JE, Maizels RM. 2011. Diversity and dialogue in immune responses in wild wood mice (Peromyscus maniculatus). J Comp Physiol B 186:419–26.

Babayan SA, Liu W, Hamilton G, Kilbride E, Rynkiewicz EC, Allen JE, Maizels RM. 2011. Chronic adaptive responses to infection. Sci Rep 8(1):3802.

Beckman RJ, Bohnke M, Begon M, Bennett M, Birtles RJ, Burthe S, Lambin Behnke JM, Gilbert FS, Abu-Madi MA, Lewis JW. 2005. Do Behnke JM, Eira C, Rogan M, Gilbert FS, Torres J, Miquel J, Baucom RS, de Roode JC. 2011. Ecological immunology and Babayan SA, Sinclair A, Duprez JS, Selman C. 2018a. The immune and non-immune pathways that drive chronic gastrointestinal helminth burdens in the wild. Front Immunol 9(56):1–12.

Babayan SA, Sinclair A, Duprez JS, Selman C. 2018b. Chronic helminth infection burden differentially affects haematopoietic cell development while ageing selectively impairs adaptive responses to infection. Sci Rep 8(1):3802.

Baucom RS, de Roode JC. 2011. Ecological immunology and tolerance in plants and animals. Funct Ecol 25:18–28.

Behnke JM, Eira C, Rogan M, Gilbert FS, Torres J, Miquel J, Lewis JW. 2009. Helminth species richness in wild wood mice, Apodemus sylvaticus, is enhanced by the presence of the intestinal nematode Heligmosomoides polygyrus. Parasitology 136:793–804.

Behnke JM, Gilbert FS, Abu-Madi MA, Lewis JW. 2005. Do the helminth parasites of wood mouse interact? J Anim Ecol 74:982–993.

Bown KJ, Begon M, Bennett M, Birtles RJ, Burthe S, Lambin X, Telfer S, Woldehiwet Z, Ogden NH. 2006. Sympatric Ixodes trianguliceps and Ixodes ricinus ticks feeding on Calidris canutus, is enhanced by the presence of the intestinal nematode Heligmosomoides polygyrus. Parasitology 136:793–804.

Clerc M, Babayan SA, Fenton A, Pedersen AB. 2019a. Age affects antibody levels and anthelmintic treatment efficacy in a wild rodent. Int J Parasitol Parasites Wildl 8:240–7.

Clerc M, Deveye G, Fenton A, Pedersen AB. 2018. Antibodies and coinfection drive variation in nematode burdens in wild mice. Int J Parasitol 8:785.

Clerc M, Fenton A, Babayan SA, Pedersen AB. 2019b. Parasitic nematodes simultaneously suppress and benefit from coccidian coinfection in their natural mouse host. Parasitology 146(8):1096–1106.

Demas GE, Nelson RJ. 1998. Short-day enhancement of immune function is independent of steroid hormones in deer mice (Peromyscus maniculatus). J Comp Physiol B 168:985–8.

Dryden MW, Payne PA, Ridley R, Smith V. 2005. Comparison of common fecal flotation techniques for the recovery of parasite eggs and oocysts. Vet Ther 6:15–28.

Eberhardson M, Lindberg K, Karlsson M, Karlen P, Winquist O, Thorn M. 2008. The sentinel node technique is useful for studies of intestinal immunology in inflammatory bowel disease patients. Eur J Gastroenterol Hepatol 20:985–8.

Ezenwa VO, Jolles AE. 2015. Opposite effects of anthelmintic treatment on microbial infection at individual versus population scales. Science 347:175–7.

Fenton A, Knowles SCL, Petchey OL, Pedersen AB. 2014. The reliability of observational approaches for detecting interspecific parasite interactions: comparison with experimental results. Int J Parasitol 44:437–45.

Folstad I, Karter AJ. 1992. Parasites, bright males, and the immunocompetence handicap. Am Nat 139:603–22.

French SS, Johnston GH, Moore MC. 2007. Immune activity suppresses reproduction in food-limited female tree lizards Urosaurus ornatus. Funct Ecol 21:1115–22.

Goff WL, Johnson WC, Parish SM, Barrington GM, Elsasser TH, Davis WC, Valdez RA. 2002. IL-4 and IL-10 inhibition of IFN-gamma- and TNF-alpha-dependent nitric oxide production from bovine mononuclear phagocytes exposed to Babesia bovis merozoites. Vet Immunol Immunopathol 84:237–51.

Gorsich EE, Ezenwa VO, Jolles AE. 2014. Nematode–coccidia parasite co-infections in African buffalo: epidemiology and associations with host condition and pregnancy. Int J Parasitol Parasites Wildl 3:124–34.

Greenman CG, Martin LB, Hau M. 2008. The sentinel node technique is useful for studies of intestinal immunology in inflammatory bowel disease patients. Eur J Gastroenterol Hepatol 20:985–8.

Gorsich EE, Ezenwa VO, Jolles AE. 2014. Nematode–coccidia parasite co-infections in African buffalo: epidemiology and associations with host condition and pregnancy. Int J Parasitol Parasites Wildl 3:124–34.

Graham AL. 2008. Ecological rules governing helmint-microparasite coinfection. Proc Natl Acad Sci U S A 105:566–70.

Graham AL, Cattadori IM, Lloyd-Smith JO, Ferrari MJ, Bjornstad ON. 2007. Transmission consequences of coinfection: cytokines writ large? Trends Parasitol 23:284–91.

Greenman CG, Martin LB, Hau M. 2005. Reproductive state, but not testosterone, reduces immune function in male house sparrows (Passer domesticus). Physiol Biochem Zool 78:60–8.

Griffiths EC, Fairlie-Clarke K, Allen JE, Metcalf CEJ, Graham AL. 2015. Bottom-up regulation of malaria population dynamics in mice co-infected with lung-migratory nematodes. Ecol Lett 18:1387–96.

Griffiths EC, Pedersen AB, Fenton A, Petchey OL. 2014. Analysis of a summary network of co-infection in humans
reveals that parasites interact most via shared resources.

Proc Biol Sci 281(1782):1–9.

Hajnicky V, Vancova I, Kocakova P, Slovak M, Gasperik J, Slavikova M, Hails RS, Labuda M, Nuttall PA. 2005. Manipulation of host cytokine network by ticks: a potential gateway for pathogen transmission. Parasitology 130:333–42.

Hardy AR, Quy RJ, Huson LW. 1983. Estimation of age in the Norway rat (Rattus norvegicus berkenhout) from the weight of the eyelens. J Appl Ecol 20:97–102.

Jackson JA, Friberg JM, Bolch L, Lowe A, Ralli C, Harris PD, Behnke JM, Bradley JE. 2009. Immunomodulatory parasites and toll-like receptor-mediated tumour necrosis factor alpha responsiveness in wild mammals. BMC Biol 7:16.

Johnson PTJ, De Roode JC, Fenton A. 2015. Why infectious disease research needs community ecology. Science 349:1259504.

Kelly CD, Stoehr AM, Nunn C, Smyth KN, Prokop ZM. 2018. Sexual dimorphism in immunity across animals: a meta-analysis. Ecol Lett 21:1885–94.

Knowles SCL, Fenton A, Petchey OL, Jones TR, Barber R, Pedersen AB. 2013. Stability of within-host - parasite communities in a wild mammal system. Proc Biol Sci 280(1762):1–9.

Lafferty KD, Allesina S, Arim M, Briggs CJ, De Leo G, Dobson AP, Dunne JA, Johnson PTJ, Kuris AM, Margocliese DJ, et al. 2008. Parasites in food webs: the ultimate missing links. Ecol Lett 11:533–46.

Lehmer EM, Jones JD, Bego MG, Varner JM, Jeor SS, Clay CA, Dearing MD. 2010. Long-term patterns of immune investment by wild deer mice infected with Sin Nombre virus. Physiol Biochem Zool 83:847–57.

Lello J, Russell T. 2008. Functional group/guild modelling of inter-specific pathogen interactions: a potential tool for predicting the consequences of co-infection. Parasitology 135:825–39.

Lindstrom KM, Foufopoulos J, Parn H, Wikelski M. 2004. Immunological investments reflect parasite abundance in island populations of Darwin’s finches. Proc Biol Sci 271:1513–9.

Love OP, Salvante KG, Dale J, Williams TD. 2008. Specific variability in the immune system across life-history stages. Am Nat 172:E99–112.

Maizels RM, Balic A, Gomez-Escobar N, Nair M, Taylor MD, Allen JE. 2004. Helminth parasites - masters of regulation. Immunol Rev. 201:89–116.

Maizels RM, Hewitson JP, Murray J, Harcus YM, Dayer B, Filbey KJ, Grainger JR, McSorley HJ, Reynolds LA, Smith CA. 2012. Immune modulation and modulators in Heligmosomoides polygyrus infection. Exp Parasitol 132:76–89.

Maizels RM, Pearce EJ, Artis D, Yazdanabaksh M, Wynn TA. 2009. Regulation of pathogenesis and immunity in helminth infections. J Exp Med 206:2059–66.

Martin LB, Weil ZM, Nelson RJ. 2006. Refining approaches and diversifying directions in ecoinmunology. Integr Comp Biol 46:1030–9.

Martin LB, Weil ZM, Nelson RJ. 2008. Seasonal changes in vertebrate immune activity: mediation by physiological trade-offs. Philos Trans R Soc Lond B Biol Sci 363:321–39.

McKay DM. 2006. The beneficial helminth parasite? Parasitology 132:1–12.

McKeen KA, Nunney L. 2005. Bateman’s principle and immunity: phenotypically plastic reproductive strategies predict changes in immunological sex differences. Evolution 59:1510–7.

Morris P. 1972. A review of mammalian age determination methods. Mamm Rev 2:69–104.

Muelchenbein MP, Bibriescas RG. 2005. Testosterone-mediated immune functions and male life histories. Am J Hum Biol 17:527–58.

Munoz L, Albillos A, Nieto M, Reyes E, Lledo L, Monserrat J, Sanz E, de la Hera A, Alvarez-Mon M. 2005. Mesenteric Th1 polarization and monocYTE TNF-alpha production: first steps to systemic inflammation in rats with cirrhosis. Hepatology 42:411–9.

Mwatha JK, Kimani G, Kamau T, Mbugua GA, Ouma JH, Mumo J, Fulford AJC, Jones FM, Butterworth AE, Roberts MB, et al. 1998. High levels of TNF, soluble TNF receptors, soluble ICAM-1, and IFN-gamma, but low levels of IL-5, are associated with hepatosplenic disease in human schistosomiasis mansoni. J Immunol 160:1992–9.

Nelson RJ. 2004. Seasonal immune function and sickness responses. Trends Immunol 25:187–92.

Nelson RJ, Demas GE, Klein SL, Kreigsfeld LJ. 2002. Seasonal patterns of stress, immune function, and disease. New York (NY): Cambridge University Press.

Nunn CL, Lindenfors P, Pursall ER, Rolff J. 2009. On sexual dimorphism in immune function. Philos Trans R Soc Lond B Biol Sci 364:61–9.

Nussey DH, Kruuk LEB, Morris A, Clements MN, Pemberton JM, Clutton-Brock TH. 2009. Inter- and intrasexual variation in aging patterns across reproductive traits in a wild red deer population. Am Nat 174:342–57.

Nussey DH, Watt K, Pilkington JG, Zamonska R, McNeilly TN. 2012. Age-related variation in immunity in a wild mammal population. Aging Cell 11:178–80.

Pakandl M, Hlaskova L, Poplstein M, Neveceralova M, Vodicka T, Salat J, Mucksova J. 2008. Immune response to rabbit coccidiosis: a comparison between infections with Eimeria flavescens and E. intestinalis. Folia Parasitol (Praha) 55:1–6.

Palacios MG, Cunnick JE, Winkler DW, Vleck CM. 2007. Immunosenescence in some but not all immune components in a free-living vertebrate, the tree swallow. Proc Biol Sci 274:951–7.

Pap PL, Vagasi CI, Tokolyi J, Czirjak GA, Barta Z. 2010. Integration in haematological indices and immune function during the annual cycle in the Great Tit Parus major. Ardea 98:105–12.

Parker BJ, Barrieau SM, Laughton AM, Griffin LH, Gerardo NM. 2017. Life-history strategy determines constraints on immune function. J Anim Ecol 86:473–83.

Pasparakis M, Alexopoulou L, Episkopou V, Kollias G. 1996. Immune and inflammatory responses in TNF alpha-deficient mice: a critical requirement for TNF alpha in the formation of primary B cell follicles, follicular dendritic cell networks and germinal centers, and in the maturation of the humoral immune response. J Exp Med 184:1397–411.
Pedersen AB, Fenton A. 2007. Emphasizing the ecology in parasite community ecology. Trends Ecol Evol 22:133–9.

Pedersen AB, Greives TJ. 2008. The interaction of parasites and resources cause crashes in a wild mouse population. J Anim Ecol 77:370–7.

Poulin R. 1996. Richness, nestedness, and randomness in parasite infracommunity structure. Oecologia 105:545–51.

Poulin R. 2001. Interactions between species and the structure of helminth communities. Parasitology 122:S3–11.

Restif O, Graham AL. 2015. Within-host dynamics of infection: from ecological insights to evolutionary predictions. Philos Trans R Soc Lond B Biol Sci 370:20140304.

Reynolds LA, Filbey KJ, Maizels RM. 2012. Immunity to the model intestinal helminth parasite Heligmosomoides polygyrus. Semin Immunopathol. 34:829–846.

R Core Team (2018). R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. URL. https://www.R-project.org/.

Rynkiewicz E, Hawlena H, Durden L, Hastriter M, Demas G, Clay K. 2013. Associations between innate immune function and ectoparasites in wild rodent hosts. Parasitol Res 112:1763–70.

Rynkiewicz EC, Pedersen AB, Fenton A. 2015. An ecosystem approach to understanding and managing within-host parasite community dynamics. Trends Parasitol 31:212–21.

Saino N, Moller AP, Bolzern AM. 1995. Testosterone effects on the immune system and parasite infestations in the barn swallow (Hirundo rustica): an experimental test of the immunocompetence hypothesis. Behav Ecol 6:397–404.

Seabloom EW, Borer ET, Gross K, Kendig AE, Lacroix C, Mitchell CE, Mordecai EA, Power AG. 2015. The community ecology of pathogens: coinfection, coexistence and community composition. Ecol Lett 18:401–15.

Sears BF, Rohr JR, Allen JE, Martin LB. 2011. The economy of inflammation: when is less more? Trends Parasitol 27:382–7.

Semenya AA, Sullivan JS, Barnwell JW, Secor WE. 2012. Schistosoma mansoni infection impairs anti-malaria treatment and immune responses in mosquito-borne Plasmodium coatneyi infected rhesus macaques. Infect Immun 80:3821.

Sjöwall J, Carlsson A, Vaarala O, Bergström O, Ernerudh J, Forsberg P, Ekerfelt C. 2005. Innate immune responses in Lyme borreliosis: enhanced tumour necrosis factor-α and interleukin-12 in asymptomatic individuals in response to live spirochetes. Clin Exp Immunol 141:89–98.

Smith NC, Ovington KS. 1996. The effect of BCG, zymosan and Coxella burnetii extract on Eimeria infections. Immunol Cell Biol 74:346–8.

Smith NC, Ovington KS, Deplazes P, Eckert J. 1995. Cytokine and immunoglobulin subclass responses of rats to infection with Eimeria nieschulzi. Parasitology 111:51–7.

Sompayrac LM. 2012. How the immune system works. 4th ed. New York, NY: Wiley-Blackwell.

Stoehr AM, Kokko H. 2006. Sexual dimorphism in immunocompetence: what does life-history theory predict? Behav Ecol 17:751–6.

Telfer S, Birtles R, Bennett M, Lambin X, Paterson S, Begon M. 2008. Parasite interactions in natural populations: insights from longitudinal data. Parasitology 135:767–81.

Vandegrift KJ, Raffel TR, Hudson PJ. 2008. Parasites prevent summer breeding in white-footed mice, Peromyscus leucopus. Ecology 89:2251–8.

Vassalli P. 1992. The pathophysiology of tumor necrosis factors. Annu Rev Immunol 10:411–52.

Wahid FN, Behnke JM. 1993. Immunological relationships during primary infection with Heligmosomoides polygyrus (Nematospiroides dubius) - parasite-specific IgG1 antibody-responses and primary response phenotype. Parasite Immunol 15:401–13.

Wali FN, Behnke JM, Conway DJ. 1989. Factors affecting the efficacy of ivermectin against Heligmosomoides polygyrus (Nematospiroides dubius) in mice. Vet Parasitol 32:325–40.

Wammes LJ, Hamid F, Wiria AE, May L, Kaisar MMM, Prasetyani-Gieseler MA, Djuardi Y, Wibowo H, Kruize YCM, Verweij JJ, et al. 2016. Community deworming alleviates geohelminth-induced immune hyporesponsiveness. Proc Natl Acad Sci U S A 113:12526–31.

Zavala GA, García OP, Camacho M, Ronquillo D, Campos-Ponce M, Doak C, Polman K, Rosado JL. 2018. Intestinal parasites: associations with intestinal and systemic inflammation. Parasite Immunol 40:e12518.

Zuk M. 1996. Disease, endocrine-immune interactions, and sexual selection. Ecology 77:1037–42.