What drives population-level effects of parasites? Meta-analysis meets life-history

Maggie J. Watson *

School of Animal & Veterinary Sciences, Charles Sturt University, Boorooma Street, Wagga Wagga, NSW 2642, Australia

Abstract

Parasites are considered drivers of population regulation in some species; unfortunately the research leading to this hypothesis has all been conducted on managed populations. Still unclear is whether parasites have population-level effects in truly wild populations and what life-history traits drive observed virulence. A meta-analysis of 38 data sets where parasite loads were altered on non-domesticated, free-ranging wild vertebrate hosts (31 birds, 6 mammals, 1 fish) was conducted and found a strong negative effect of parasites at the population-level (g = 0.49). Among different categories of response variables measured, parasites significantly affected clutch size, hatching success, young produced, and survival, but not overall breeding success. A meta-regression of effect sizes and life-history traits thought to determine parasite virulence indicate that average host life span may be the single most important driver for understanding the effects of parasites. Further studies, especially of long-lived hosts, are necessary to prove this hypothesis.

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1. Introduction

A central goal of population ecology is to identify factors controlling population dynamics. In wild populations, predation and competition are well studied, with some theoretical and empirical investigations focusing on the effects of parasites. Population regulation by parasites has been identified in Red Grouse Lagopus lagopus scoticus (Hudson et al., 1998), Svalbard Reindeer Rangifer tarandus platyrhinchus (Albon et al., 2002) and Soay Sheep Ovis aries (Gulland, 1992); unfortunately, these examples represent managed populations, and therefore may not reflect true effects of parasites on wild populations. Thus, the question remains—are parasites significant drivers of population-level effects and what host life-history traits drive observed virulence (sensu lato Casadevall and Pirofski, 1999—the capacity of a parasite to cause damage to a host)?

The modern view of parasitism is predicated on the assumption that ‘every parasitic organism... imposes a cost on its host’ because resources, however slight, are being diverted from host to parasite (Combes, 2005). These costs can be couched in two evolutionary trajectories: (1) the ‘mutual aggression model, (Holmes, 1983) which suggests that parasites evolve to be as virulent as possible, and thus are a primary regulatory force; and (2) the ‘prudent parasite model’ (Holmes, 1983; Renaud and de Meeus, 1991) which suggests that parasites evolve towards a balance between short- and long-term needs conferring a range of benefits to the infected host that may or may not offset the costs (Michalakis et al., 1992; Schmidt-Hempel, 2003).

Several researchers have argued that the only way to assess the true effects of parasites is by altering the parasite population of the host in situ (Møller, 2005). Alterations of parasite loads are easy to effect in domestic and laboratory animals, and even wild animals in the laboratory (Diamond, 1983; McCallum, 1995). However, relatively little parasite work on wild, free-ranging hosts incorporates this technique due to logistical difficulties surrounding field work and obtaining sufficient sample sizes to detect differences between infected and non-infected hosts. Therefore, much ecological work on the effects of parasites ends up being correlative (Poulin, 2007a). It is unclear if the differences detected between parasitized and non-parasitized hosts are due to indirect effects or pre-existing differences (i.e., prior to infection; Bize et al., 2008 or host-quality; Lailvaux and Kasumovic, 2011). Field experimentation is necessary to quantify actual costs of parasites on hosts due to the many problems associated with extrapolating laboratory results on individuals or populations to real effects in the field (Seitz and Ratte, 1991).

In order to understand if parasites are truly a driver of host populations, reviews of the effects of parasites to wild hosts need to be conducted. Reviews to date of both observational and
experimental work on the cost of parasites to wild hosts (birds, mammals, fish and insects: Lehmann, 1993; birds: Möller, 1997; birds and mammals: Tompkins and Begon, 1999; mammals: Irvine, 2006) have implied that parasites are costly, but the implications of that cost are unreliable, due to the methods used to synthesize results (Stewart, 2010). A recent meta-analytical synthesis of parasite induced mortality of nestlings showed an overall small effect (12% mean parasite-induced mortality, range 0–89%, n = 117), with parasite-induced mortality determined by latitude, nesting site, probability of host survival and parasite prevalence (Møller et al., 2009). However, this meta-analysis only considered studies of nesting birds and may be fundamentally flawed because it includes observational data as well as experimental data (Borenstein et al., 2009).

The objectives of the present analyses were to review quantitatively experimental studies of wild, free-ranging hosts that measure parasite-induced changes in population-level traits (i.e. measures of fecundity and mortality); then, to evaluate this effect of parasites using life-history traits. Based on those host life-history traits that Möller et al. (2009) found to be significant, the following predictions are made: (1) cavity-nesting species (includes burrowing mammals as well as hollow nesting birds) will experience increased parasite density and intensity and thus more virulent effects than ground or open nesting species (Ewald, 1983); (2) colonial species will experience increased parasite density and intensity and thus more virulent effects than less gregarious species (Ewald, 1983); (3) tropical species will encounter more virulent parasites than temperate species because the absence of seasonality maintains higher parasite abundance (Møller, 1998); and (4) higher virulence will evolve in hosts with shorter life-spans because of the fewer opportunities there are for dispersal to a new host in search of a mate, and as a consequence, the parasites become more virulent (Lehmann, 1993; Nidelet et al., 2009).

2. Materials and methods

2.1. Data collection and inclusion criteria

The studies considered for use in the meta-analysis were obtained from a survey of the primary literature. The initial search was directed using reviews by Möller (1997), Newton (1998), Tompkins and Begon (1999) and Irvine (2006) followed by a comprehensive search of ISI Web of Science and Google Scholar up to and including January 2012. The following search terms and their combinations were used: “parasites”, “experiment”, “manipulation”, “cost”, “effects”, “mortality”, “survival”, “fitness”, “hosts”, “life-history”. Older literature (pre-1985) was identified through Literature Cited sections of recent papers and unpublished theses (the same search terms were used in ProQuest Dissertations and Theses, Theses Canada and Trove). Only papers written in English were included. When reference to unpublished work was encountered, attempts were made to solicit raw data from the author(s). A large number of studies were screened using abstracts only (<2000); 89 full-text articles were assessed for eligibility. Of these, 51 were excluded due to a lack of numerical data, lack of sample size and/or variance, untranslatable test statistics, duplication of dataset from a previous paper or reported results not relevant to the selection criteria (e.g. behavioural or physiological/individual responses). Studies were selected if (a) host species were wild (not domesticated), free-ranging (not held in captivity) and the study was conducted under field-conditions (not laboratory conditions); (b) parasite species were experimentally manipulated (increased or decreased); and (c) the parasite was naturally occurring and not introduced, thus avoiding the ‘suicide king’ issue of parasites infecting hosts outside their normal range and becoming more virulent in the process (Dybå and Storfer, 2003). Of these, a study was included in the final meta-analyses if it provided (a) the means and standard errors or standard deviations (or any other statistic whereby means and standard errors could be derived) of at least one population-level parameter measuring the cost of parasitism for experimental and control groups, and (b) the sample sizes associated with the means.

2.2. Response variables and calculation of effect sizes

Response variable and effect size data were extracted from the text and tables for all studies except Cheney and Coté (2003), Fitze et al. (2004), Pap et al. (2005), Slomczynski et al. (2006) (additional information requested and received from the authors); and Bize et al. (2004) and Hillegass et al. (2010) (data extracted from graphs using DataThief; Tummers, 2006). Statistics were converted to effect sizes in the form of Hedges’ g (Hedges and Olkin, 1985) in the program Comprehensive Meta-analysis (CMA; Borenstein, 2006). Hedges’ g was chosen as the effect size over the more commonly used Cohen’s d because Hedges’ g pools variance using n – 1 instead of n and thus provides a better estimate for smaller sample sizes (Grisson and Kim, 2005). Studies that reported only F statistics (Møller, 2002; Vandegrift et al., 2008) were not converted to effect sizes due to issues surrounding the overestimation of effect sizes identified by Hultell and Levine (2003) and lack of accurate sample size data in the respective articles. The type of response variable was coded into the data set to enable subgroup analyses. The response variables used were: clutch size (number of eggs in the clutch), percent hatching success (percentage of eggs that hatched from a single clutch), number of young produced (total brood size), percent breeding success (percentage of young produced, fledged or survived during the study period), and survival rate (survival during the study period or between one breeding season and the next).

2.3. Meta-analytic procedures

All meta-analyses were performed in CMA (Borenstein, 2006). A random-effects model was used for all tests because variability was expected in the effects being measured across different species and hosts. Many articles included multiple effect sizes from different measures of the effects of parasitism, so rather than combining all the effect sizes within a study (which may have obfuscated the true effect), in the overall meta-analysis one effect size was chosen at random from each of the forty-three studies (Gurevitch and Hedges, 1999). Separate random-effects meta-analyses were conducted grouped by effect being studied—so any given study might have data in several meta-analyses (sub-analyses) thus maintaining the independence of the data (Gurevitch and Hedges, 1999). One study (Roby et al., 1992) considered the responses of two host species to the same anti-parasitic treatment, so the two hosts were considered as independent studies.

2.4. Heterogeneity and publication bias

Publication bias, or the ‘file drawer problem’, where non-significant results are relegated to the file drawer rather than to the published literature (Rosenberg, 2005), is an ongoing issue affecting meta-analyses, leading to bias via the selective publication of statistically significant results (Hedges and Olkin, 1985). To guard against this issue, publication bias was assessed using three methods: funnel plot (plot of effect size and precision to search for asymmetry), Q-rank correlation (a test for publication bias; Beggs and Mazumdar, 1994), and trim-and-fill (Duval and Tweedie, 2000). Heterogeneity indicates the presence of effect-modifiers, and the Q-test for heterogeneity was calculated for the overall
meta-analysis. However, because the meta-analysis spans many classes of organism, both as hosts and as parasites, heterogeneity is expected to be high.

2.5. Meta-regression

Meta-regression analyses, a tool used to examine the impact of moderator variable on effect sizes, were performed to assess the degree to which certain life-history traits influence the virulent effects of parasites. These life-history traits (living in cavities/hollows, coloniality, latitude, and host life-span) have been identified as possible predictors to parasite virulence (Møller et al., 2009). Cavity living was coded as 0 for species that lived in the open and 1 for species that lived or bred in tree hollows or underground. Coloniality was coded by maximum recorded colony size: 0 for solitary, 1 for 2–10 pairs, 2 for 11–100 pairs, 3 for 101–1000 pairs, 4 for 1001+ pairs, and 5 for unknown. Latitude was recorded as the maximum latitude where the species was found. Host life-span was recorded as the maximum recorded life-span of the host species. These life-history traits were included in the meta-regression as potential predictors of parasite virulence.

Table 1

Studies used in the meta-analysis investigating using parasite load manipulations the effect of parasites on their wild, free-ranging hosts. Hosts are listed in taxonomic order using common name and parasites are listed by common name. Data for the meta-regression are coded in the following order: living in cavities/hollows (0 = open; 1 = cavity/hollow), coloniality (0 = solitary; 1 = 2–10 pairs; 2 = 11–100 pairs; 3 = 101–1000 pairs; 4 = 1001+ pairs), latitude (in cases where the same species is used from different locations, just the latitude is recorded), and host life-span (maximum recorded; average).

| Host | Parasite | Response variable | Source paper | Meta-regression |
|------|----------|-------------------|--------------|-----------------|
| Piscine Perciformes | Isopods | Clutch size | Cheney and Côté (2003) | 0, 0,13.10, 6, 6 |
| Longfin Damselfish | Nematode | Clutch size, hatching success, # young produced | Hudson (1986) | 0, 0, 53.95, 8, 3 |
| Avian Galliformes | Nematodes | # Young produced | Redpath et al. (2006) | 0, 4, 27.55, 17, 8 |
| Red Grouse | Lice | Hatching success | Clayton et al. (1999) | 0, 3, 41.15, 36, 6 |
| Columbiaforms | Ticks | # Young produced | McKilligan (1996) | 0, 4, 27.55, 17, 8 |
| Rock Dove | Ticks | Hatching success | Norcross and Bolen (2002) | 0, 3, 33.56, 27, 25 |
| Columbiformes | Nematodes, trematodes, cestodes | Clutch size, hatching success, # young prod., % succ. | Van Oers et al. (2002) | 0, 0, 53.29, 43, 12 |
| Crested Tern | Lice, ticks | # Young produced | Watson unpub. Data | 0, 4, 38.31, 32, 15 |
| Alpino Swift | Louse-flies | # Young produced | Bize et al. (2004) | 1, 3, 47.12, 26, 6 |
| Barn Swallow | Mites | Clutch size, hatching success, # young produced | Møller (1990) | 1, 2, 57.12, 10, 6 |
| Cliff Swallow | Bugs | # Young Produced | Brown and Brown (1986) | 1, 4, 41.13, 11, 5 |
| Tree Swallow | Blow fly | % Successful | Roby et al. (1992) | 1, 2, 42.59, 12, 3 |
| House Martin | Bugs | Hatching success, % successful, # young produced | de Lope and Møller (1993) | 1, 3, 38.50, 15, 2 |
| Purple Martin | Mites | Hatching success, % successful | Moss et al. (1966) | 1, 2, 38.58, 13, 8 |
| Sand Martin | Ticks | # Young produced | Szép and Møller (1999) | 1, 3, 48.08, 10, 2 |
| European Starling | Mites | Hatching success, % successful | Roby et al. (1992) | 1, 0, 42.59, 10, 6 |
| Blue Tit | Flies | Clutch size, # young produced | Richner and Tripet (1999) | 1, 0, 46.31, 15, 3 |
| Haematazoa | Flies, blow fly | % Successful | Merino et al. (2000) | 40.48 |
| Haematazoa | Flies, blow flies, mites | % Successful | Boušlama et al. (2002) | 36.42 |
| Great Tit | Flies | # Young produced | Richner et al. (1993) | 1, 0, 46.31, 15, 3 |
| Fleas | Clutch size, # young produced | Opplinger et al. (1994) | 46.31 |
| Fleas | Clutch size | Fitze et al. (2004) | 46.54 |
| Mammalian Lagomorpha | Nematodes | # Young produced, survival rate | Bloomer et al. (1995) | 0, 0, 44.18, 5, 1 |
| Mountain Hare | Nematodes | Survival rate | Newey and Thirgood (2004) | 0, 2, 57.00, 9, 4 |
| Townsend's Vole | Bot fly, mites, nemaotes | Survival rate | Steen et al. (2002) | 1, 0, 49.04, 1, 1 |
| Rodentia (Sciuridae) | Ticks, lice, fleas, worms | # Young produced | Hillegass et al. (2010) | 1, 2, 27.35, 6, 4 |
| Cape Ground Squirrel | Fleas | # Young produced | Neuhaus (2003) | 1, 2, 50.00, 11, 3 |
| Cetartiodactyla | Nematodes | Survival rate | Schmidt et al. (1979) | 0, 2, 37.34, 24, 6 |

M.J. Watson / International Journal for Parasitology: Parasites and Wildlife 2 (2013) 190–196
101–1000 pairs and 4 for 1001+ pairs. Latitude was entered using information provided in the source article. Host life-span was coded as maximum recorded (from a wild individual where possible; Carey and Judge, 2002) and average (age an individual which reaches breeding age can be expected to live; Robinson, 2005); any gaps in these two databases were filled using the Encyclopedia of Life.

3. Results

The comprehensive literature search yielded 38 studies that reported the required statistics (or the required statistic was supplied by the author) and so were used in the meta-analysis (Table 1). These studies comprised 31 on avian hosts, 6 on mammalian hosts and 1 on a fish host. There were no studies of reptiles or amphibians that fitted the parameters of this meta-analysis.

3.1. Overall effect and publication bias

The data set were first analysed as a whole (termed combined in Table 1). In order to avoid replication for this combined analysis, one sample was randomly chosen from each study. A forest plot of the effect sizes for all studies showed that twenty-seven reported negative effects of parasites, and eleven showed positive or nil effects (Fig. 1). The mean effects size across all studies was 0.489 (95% CI, 0.220–0.759, n = 38) and was statistically significant from a null effect size (Z = 3.56, p = 0.0004). Additionally, to determine what impact the random sampling might have had on the power of the combined analysis, a meta-analysis was performed on the entire data set resulting in a mean effect size of 0.47 (95% CI, 0.282–0.658, n = 60).

A funnel plot suggested some publication bias (Fig. 2), however, a rank correlation test between precision and effect size (Begg and Mazumdar, 1994) was not statistically significant (Kendall’s τ = 0.105, P = 0.176; one-tailed, with continuity correction). Additionally, the trim-and-fill analysis of the random effects model imputed no missing studies. Despite some evidence to suggest that there was a publication bias, there is much support for the negative effects of parasites at the population level.

3.2. Subgroup analyses

As expected, there was significant heterogeneity in effect sizes across studies (Q = 749.54, df = 37, p < 0.0001), thus, subgroup analyses on response variables were analysed, both to examine possible variables influencing the results and to examine the data from a biologically relevant point of view. The complete data set (not the randomly selected subset) were separated into the response variable measured—clutch size (n = 12), percent hatching success (n = 11), number of young produced (n = 20), percent breeding success (n = 12), and survival rate (n = 7). A meta-analysis was performed on each group (Table 2) and revealed significant negative effects of parasites on clutch size, hatching success, and young produced, but not overall breeding success or survival rate.

3.3. Meta-regression of life history traits

The effect sizes from the complete data set (n = 60; not the randomly sampled subset used in the overall meta-analysis) were regressed against five moderator variables—cavities/hollow living, coloniality, latitude (chosen from Möller et al. (2009) paper on variables that significantly impact chick survival), maximum host life-span, and average host life-span (chosen as a more precise measure of intrinsic host mortality than survival rate used by Möller et al., 2009). The proportion of variance explained by all...
These moderator variables was nonsignificant except for average age (significant at p = 0.05) (Table 3; Fig. 3).

4. Discussion

A meta-analysis of 38 experimental studies of the costs of parasites to population-level measures of natural, free-ranging hosts revealed an overall effect size, $g$, of 0.49. These parasites, therefore, produced a moderate (as defined by Cohen, 1988) negative effect. This effect size can be interpreted as an average difference of 0.49 standard deviations between parasitised and non-parasitised individuals. This effect is as strong as the reported effect sizes of predators on populations. Côté and Sutherland (1997) quantified the combined effects of predation and competition on population sizes of anurans in field experiments. In this case, the summary effect size is obfuscated by interactions between predation and competition, but the authors state that “the average over-all effects of predator exclusion were very large” and the average effect of competitor removal in the presence of predators on survival was $-0.35$ (negatively effecting survival). Given the summary effect size reported in this study, the indication is that parasites may be at least as important as predation (or predation combined with competition) in their effects on populations. It is important to include parasites in future research that records demographic information. By incorporating parasite removal experiments into predation- or competition-based experiments, the interactions between these demographic forces will yield a broader understanding of host–parasite interactions (e.g. Holt and Roy, 2007).

Meta-analyses of the response variables indicated that clutch size, hatching success, and number of young produced were all significantly reduced in parasitised individuals. The only response variables that was not significantly affected by parasites was breeding success (defined variously as percentage of young produced, fledged or survived during the study period) and survival (although sample size was very low for survival rate, $p = 0.053$, thus more studies may indicate that survival is indeed significantly reduced by parasites). Further analyses using meta-regression to determine the key life-history traits that may be driving the observed population-level effects of parasites revealed that the most likely key driver of effects of a parasite on its host was host lifespan. Other variables previously suggested (Moller et al., 2009) as determinants of virulence—cavity/hollow living, coloniality or latitude were not significant.

There are strongly suggestive mathematical models (Anderson, 1978; Anderson and May, 1978; May and Anderson, 1978) and some empirical evidence (Gregory, 1991; Hudson et al., 1998) that parasites can regulate host populations and cycles. The large effect size revealed in this meta-analysis supports the idea that parasites are a major force in population regulation in wild, free-ranging vertebrate host populations. Support for the idea of the importance of parasites has far-reaching implications in both ecological and conservation-based science. If parasites are such strong regulators, why are some species seemingly untouched by the effects of parasites despite large parasite loads? Are parasites simply another form of predation? Should parasites be removed in populations needing conservation regulation, or, conversely, added to invasive populations? Unfortunately, before drawing a longbow based on these strong suggestions, it is necessary to understand the full extent of the effects of parasites on natural populations.
around this meta-analysis, it is important to examine the taxonomy of the host species that went into the model. While this meta-analysis does include studies from twenty-four different species, they are mainly clustered within one order of one class of host study organism. The taxonomic bias of host species studied must temper any conclusions—these results may only be applicable to the hosts used in the meta-analysis. It may be that the effects of parasites as measured here are inflated compared to other sorts of hosts and the actual overall effects of natural parasites are much less.

Given these caveats, the information provided from this meta-analysis still lends itself to hypothesis generation and testing around the life-history traits that drive the evolution of a parasite to cause damage to its host. The results of the meta-regression, which refuted all previous predictions regarding the determinants of parasite virulence, suggest that intrinsic host mortality (life-span) is the key to understanding why and how parasites evolve to either harm or be benign to their host. Although the significance of this result is not high, it may suggest why observed parasite virulence varies so widely between host species. The evolutionary idea that background host mortality may explain an increase in virulence has been demonstrated in the laboratory (Nidelet et al., 2009). Additionally, longer host lifespan have been shown to evolve under high condition dependent morality (Williams and Troy, 2003; Dowling, 2012). Combining these two ideas with the results of this meta-analysis and meta-regression suggests that we should expect long-lived hosts to have non-virulent parasites except when the host’s condition is poor. Moreover, short-lived hosts should show increased responses to parasites regardless of their own condition. Therefore, in order to increase our understanding of the effects of parasites on wild populations, focused studies on long-lived hosts are vital. Especially, where long-lived and short-lived examples can be found within the same taxonomic group.

The surprising lack of support for the hypotheses that parasites should be more virulent with cavity/hollow living, higher numbers of individuals living or breeding together, and in tropical latitudes is illustrative of the conundrum of mixing the results of observational studies with experimental ones. Intuitively, one expects that species living or breeding in hollows, underground or in a colony would experience greater abundance and prevalence of parasites due to repeated use of the living area. However, the number of parasites on an individual host does not necessarily indicate an increased effect of that parasite on the host. Additionally, when one considers observational data, the researcher is asking if parasitised individuals are more affected by parasites than those individuals who are naturally uninfected, while an experimental study seeks to understand if parasites have more effects regardless of the chance or reason for having become infected in the first place.

In conclusion, this study provides a quantitative test of the effects of parasites to their wild, free-ranging vertebrate hosts. The results suggest that parasites are indeed drivers of population-level life-history traits, as has been advocated for over thirty years. However, the results are heavily taxonomically biased, and therefore, may only be applicable to a limited number of species and scenarios. Interrogation of the data using meta-regression revealed that the one life-history trait that appears to drive the capacity of a parasite to negatively impact its host is intrinsic host lifespan. However, it is difficult to separate the relative importance of these life-history traits in shaping parasite virulence, and more data are needed to understand these phenomena. An examination of the effects of parasites on long-lived hosts is warranted to understand fully the magnitude and extent of the effects of parasites in natural situations.

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