Host Patch Traits Have Scale-Dependent Effects On Diversity In A Stickleback Parasite Metacommunity

Daniel I. Bolnick  
*University of Texas at Austin*

Emlyn J. Resetarits  
*University of Texas at Austin*

Kimberly Ballare  
*University of Texas at Austin*

Yoel E. Stuart  
*Loyola University Chicago, ystuart@luc.edu*

William E. Stutz  
*University of Texas at Austin*

Follow this and additional works at: [https://ecommons.luc.edu/biology_facpubs](https://ecommons.luc.edu/biology_facpubs)

Part of the Biology Commons

**Recommended Citation**

Bolnick, Daniel I.; Resetarits, Emlyn J.; Ballare, Kimberly; Stuart, Yoel E.; and Stutz, William E.. Host Patch Traits Have Scale-Dependent Effects On Diversity In A Stickleback Parasite Metacommunity. *Ecography, 43, 7: 990-1002, 2020.* Retrieved from Loyola eCommons, Biology: Faculty Publications and Other Works, [http://dx.doi.org/10.1111/ecog.04994](http://dx.doi.org/10.1111/ecog.04994)

This Article is brought to you for free and open access by the Faculty Publications and Other Works by Department at Loyola eCommons. It has been accepted for inclusion in Biology: Faculty Publications and Other Works by an authorized administrator of Loyola eCommons. For more information, please contact ecommons@luc.edu.

This work is licensed under a [Creative Commons Attribution 3.0 License](http://creativecommons.org/licenses/by/3.0/). © Bolnick et al., 2020.
Many metacommunities are distributed across habitat patches that are themselves aggregated into groups. Perhaps the clearest example of this nested metacommunity structure comes from multi-species parasite assemblages, which occupy individual hosts that are aggregated into host populations. At both spatial scales, we expect parasite community diversity in a given patch (either individual host or population) to depend on patch characteristics that affect colonization rates and species sorting. But, are these patch effects consistent across spatial scales? Or, do different processes govern the distribution of parasite community diversity among individual hosts, versus among host patches? To answer these questions, we document the distribution of parasite richness among host individuals and among populations in a metapopulation of threespine stickleback *Gasterosteus aculeatus*. We find some host traits (host size, gape width) are associated with increased parasite richness at both spatial scales. Other patch characteristics affect parasite richness only among individuals (sex), or among populations (lake size, lake area, elevation and population mean heterozygosity). These results demonstrate that some rules governing parasite richness in this metacommunity are shared across scales, while others are scale-specific.

Keywords: diet, helminth, infection, macroparasite, metacommunity, threespine stickleback

**Introduction**

A long-standing question in ecology is, ‘why are some communities more diverse than others?’ To address this question, biologists have sought to identify and quantify the biotic and abiotic factors that affect community diversity, and which differ across a landscape. A clear lesson from such work is that the salient factors depend on the spatial scale being considered (Cotterie 2005, Chave 2013, Meynard et al. 2013, Leibold and Chase 2017). For instance, ecosystem productivity is often negatively associated with community diversity at small spatial scales, but the trend can be reversed at large scales (Chase and Leibold 2002, McBride et al. 2014). Similarly, here we show that factors affecting parasite
in many animal species, co-occurring individuals actively pre-

Prediction 1. Host diet has a scale-independent effect on parasite richness

In many animal species, co-occurring individuals actively prefer different prey, resources or microhabitats (Bolnick et al. 2003). Host populations also diverge in diet and resource use, often along the same ecological axes as the individual-level diet variation (Schluter and McPhail 1992, Matthews et al. 2010a). Because many parasites are trophically transmitted (Wilson et al. 1996, Johnson et al. 2009, Cirtwill et al. 2016), host diet should correlate with parasite richness. This correlation should be consistent across spatial scales, assuming each parasite’s complex life cycle is relatively invariant across the metacommunity.

Prediction 2. Host ecomorphology has a scale-independent effect on parasite richness

Diet variation often reflects variation in trophic morphology that affects prey capture and processing (Wainwright and Richard 1995). A corollary of Prediction 1 is that trophic morphology should be correlated with parasite richness. Indeed, morphology might be a more reliable covariate of parasite richness because diet measurement is noisy and often restricted to short time scales. If the diet–morphology relationship is consistent across host populations, morphology effects might therefore also act similarly across spatial scales.

Prediction 3. Host heterozygosity has a scale-independent, negative effect on parasite richness

Parasite diversity should be inversely related to the host’s ability to recognize and eliminate various parasites (given equal exposure risk). Such host immune competence can be related to genetic diversity at particular loci (e.g. MHC; Wegner et al. 2003, Kalbe et al. 2009, Oliver et al. 2009), but immunity will often depend on the collective action of many loci. Thus, another measure of immunogenetic diversity might be genome-wide genetic heterozygosity (Coltman et al. 1999, Arkush et al. 2002, Šimková et al. 2008, Whitehorn et al. 2011). We predict that genome-wide heterozygosity is negatively related to parasite richness among both scales. Alternatively, parasite richness might be controlled by host immune genotypes at specific loci that affect resistance to many taxa at once. If such loci evolved in parallel across replicate populations, we should find richness associated with certain SNPs’ genotypes, at both spatial scales.

Prediction 4. Host sexual dimorphism contributes only to within-population variation in parasite richness

In many species, sexes differ in feeding ecology (Shine 1991) and immunity (Zuk 1996, Nunn et al. 2009), which should influence individual-level parasite richness (Morand and Bordes 2015). Because host populations have a roughly equal sex ratio, we do not predict that dimorphism will influence among-population variation in parasite richness.

Prediction 5. Habitat contributes to among-population differences in parasite richness

Elevation, lake size and distance from the ocean are geographic features that should be experienced by all individuals living in...
Methods

Collection

In June 2009 we collected threespine stickleback from 46 sites on Vancouver Island, British Columbia, Canada, in the historical lands of the Kwakwaka’wakw First Nations. Our sample sites included five estuaries with anadromous fish and 33 lakes and eight streams from nine watersheds (Supplementary material Appendix 1 Table A1, Fig. A1). At each site, we placed unbaited 0.5-cm gauge wire minnow traps along ~200 m of shoreline in 0.5–3 m deep water until we took 60–100 fish per site (Supplementary material Appendix 1 Table A1). Fish were euthanized in MS-222 and preserved in 10% buffered formalin after saving a fin clip in ethanol for DNA. Specimens were later rinsed and stored in 70% isopropyl alcohol after staining with Alizarin Red.

Parasite diversity

We scanned each fish under a dissection stereomicroscope to count and identify macroparasites (helminths, crustaceans, mollusks and microsporidia) to the lowest feasible taxonomic unit, following Stutz and Bolnick (2017). We focus on two parasite diversity statistics (Supplementary material Appendix 1 Fig. A2): 1) the number of uniquely identifiable parasite taxa for each fish, \( \alpha_i \), i.e. parasite richness in individual host \( i \) from host population \( j \); 2) the average per-individual parasite richness, \( \bar{\alpha}_j \), for all fish in population \( j \).

Ecomorphology

Before dissection, we weighed all fish within 0.01 g and used digital calipers to measure standard length, body depth and body width at the pectoral fins (mm). For a random subset of 30 individuals per population, we measured three trophic traits – gape width, gill raker number and longest gill raker length – and averaged left- and right-side armor plate number. We sexed all individuals by visually inspecting gonads. Linear measurements were log transformed and size-standardized by regression on log standard length. We calculated trait means per population for population-scale analyses. Note that none of the populations exhibited a multimodal size or weight distributions that would imply a multi-year age structure, consistent with previous work on stickleback concluding that the vast majority die at the end of their first breeding season (Woottton 1984).

Diet

For a random subset of 28 populations, we analyzed fish stomach contents in the same 30 fish used for ecomorphology, recording presence of prey to the lowest feasible taxonomic level to calculate the number of observed prey taxa per fish (prey richness). Stickleback individuals and populations vary in their relative consumption of benthic invertebrates versus pelagic zooplankton (Lavin and McPhail 1986, Ingram et al. 2011), so we use the proportion of benthic prey taxa as another diet metric. This measure was highly correlated with the first axis of a non-metric multidimensional scaling (NMDS) analysis using fish stomach contents (per the Jaccard index in the R package vegan; Supplementary material Appendix 1 Table A2). Several studies have confirmed that stomach contents are correlated with direct observations of feeding behavior, with long-term measures of diet using stable isotopes, and with trophic morphology (Matthews et al. 2010b, Snowberg et al. 2015).

Genetic diversity

With DNA from a subsample of 12 fish per population for four marine sites, 31 lakes and six streams, we used ddRAD-seq (Peterson et al. 2012) to obtain genotypes for 175 350 SNPs in 336 fish (mean 107 698 SNPs/individual). The protocol and bioinformatic pipeline are described in Stuart et al. (2017). We calculated genome-wide mean heterozygosity (\( H_i \)) for each fish \( i \) in population \( j \), and mean heterozygosity for each population (\( \bar{H}_j \)).

Statistical analyses

All statistical models were implemented in R ver. 3.5.2 (Table 1). Our primary goal in this study is to understand why host individuals and host populations differ in parasite diversity. We fit a Poisson general linear model (GLM) evaluating how per-fish parasite richness varies as a function of host population nested within watershed (both random effects, see model M1 in Table 1). Within the majority of lakes, and in the dataset overall, a Poisson distribution was an effective description of among-individual variation in parasite richness. For each source of variation, we calculated its effect size using \( \eta^2 \). Note that a companion paper (Bolnick et al. 2019) found little effect of geographic proximity (e.g. spatial autocorrelation) on the parasite community composition. So, we do not here consider spatial autocorrelation except to note that between-lake difference in mean parasite richness is uncorrelated with geographic distance as-the-crow-flies (Mantel \( r = 0.05, p = 0.20 \)).

What host traits affect parasite richness variation among individual hosts?

Within each host population, we tested for correlations between individuals’ phenotypic traits and their parasite richness. Due to a shared environment and ancestry, any infection differences among fish within a population should stem from individual hosts’ phenotypic characteristics or genotype (i.e. Predictions 1–4). We focused exclusively on lake fish to exclude habitat differences among populations. We omitted watershed, which had
Table 1. Models tested in this study. A summary of the statistical results of each model is provided in Supplementary material Appendix 1.

<table>
<thead>
<tr>
<th>Name</th>
<th>Model:</th>
<th>Does parasite richness depend on:</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1:</td>
<td>(\logit(\alpha_i) \sim \lambda_j) \text{watershed}</td>
<td>host population, nested within watershed</td>
</tr>
<tr>
<td>M2:</td>
<td>(\logit(\alpha_i) \sim \lambda_j + \text{sex}_i) \text{length}_j</td>
<td>individual host sex and size</td>
</tr>
<tr>
<td>M3:</td>
<td>(\logit(\alpha_i) \sim \lambda_j + \text{sex}_i + \text{length}_j + \text{GRN}_i + \text{GRL}_i + \text{PN}_i + \text{GW}_i)</td>
<td>individual host ecomorphology</td>
</tr>
<tr>
<td>M4:</td>
<td>(\logit(\alpha_i) \sim \lambda_j + \text{sex}_i + \text{length}_j + \text{NMDS}_1 + \text{NMDS}_2 + \text{DB}_i)</td>
<td>individual host diet</td>
</tr>
<tr>
<td>M5:</td>
<td>(\logit(\alpha_i) \sim \lambda_j + \text{set}_i + \text{length}_j + \text{SNP}_i)</td>
<td>individual host genotype at SNP (k) (GWAS)</td>
</tr>
<tr>
<td>M6:</td>
<td>(\logit(\alpha_i) \sim \lambda_j + \text{sex}_i + \text{length}_j + \text{Het})</td>
<td>individual host mean genome-wide heterozygosity</td>
</tr>
<tr>
<td>M7:</td>
<td>(t[\alpha_i] = t[\text{length}] + t[\text{GRN}] + t[\text{GW}] + t[\text{NMDS}_1] + t[\text{NMDS}_2])</td>
<td>sexual dimorphism in ecomorphology and diet</td>
</tr>
<tr>
<td>M8:</td>
<td>(\alpha_{ij} \sim \text{habitat} + \text{watershed})</td>
<td>habitat</td>
</tr>
<tr>
<td>M9:</td>
<td>(\alpha_{ij} \sim \log \text{lake size} + \text{elevation} + \text{ocean distance})</td>
<td>lake geography</td>
</tr>
<tr>
<td>M10:</td>
<td>(\alpha_{ij} \sim \text{mean length} + \text{mean GW} + \text{mean GRN} + \text{mean PN} + \text{mean GRL})</td>
<td>host population ecomorphology</td>
</tr>
<tr>
<td>M11:</td>
<td>(\alpha_{ij} \sim \text{mean NMDS}_1 + \text{mean NMDS}_2 + \text{mean DB})</td>
<td>host population diet</td>
</tr>
<tr>
<td>M12:</td>
<td>(\alpha_{ij} \sim \text{freq(SNP)}_i + \text{watershed})</td>
<td>host population allele frequency at SNP (k) (GWAS)</td>
</tr>
<tr>
<td>M13:</td>
<td>(\alpha_{ij} \sim \text{mean Het})</td>
<td>host population mean heterozygosity</td>
</tr>
<tr>
<td>M14:</td>
<td>(\text{mean Het} \sim \log \text{lake size} + \text{elevation} + \text{ocean distance})</td>
<td></td>
</tr>
</tbody>
</table>

**KEY:** \(\lambda\); population-specific Poisson rate (random effect intercept); Subscript \(j\) on any of the following traits indicates we fit a main effect and population random effect of a given trait (e.g. population by trait interactions); length: log transformed standard length; GRN: gill raker number; GRL: size-corrected gill raker length; PN: armor plate number; GW: size-corrected gape width; NMDS: diet ordination axes 1 and 2; DB: diet breadth; SNP: genotype at a particular SNP \(k\); \(t[\cdot]\): \(t\)-statistic measure of sexual dimorphism; Het: genome-wide mean heterozygosity. Population trait means or allele frequencies are indicated as such.

We carried out two analyses of the effects of host individuals’ genotype on individual parasite richness. First, we used genome-wide association (GWAS) to test each SNP’s effect on parasite richness (M5). Specifically, we used Poisson GLMMs to evaluate how individual parasite richness depends on population, sex, length and focal SNP genotype. We retained a population-specific random effect of sex, but not length, because only the former was supported in models M2–4. We applied M5 separately for each of 39,039 SNPs (restricting our attention to SNPs scored in at least 50 fish, with minor allele frequency exceeding 10%). We applied Holm corrections to the \(p\)-values.

Second, to test effects of whole-genome genetic diversity on parasite richness, we used Poisson GLMM (M6) to relate parasite diversity to individual hosts’ genome-wide mean heterozygosity, with population as a random slope and intercept.

We next tested for covariates of among-population variation in mean per-fish parasite richness \(\alpha_{ij}\). Although population constitutes the level of replication in this analysis, model M7 seeks to explain the magnitude of between-population variation (e.g. a source of within-population variation).
We focus on mean per-fish richness within populations, rather than aggregate parasite richness of a whole sample, because they are tightly correlated \((r = 0.58, p < 0.0001;\) Supplementary material Appendix 1 Fig. A3), and the former is not biased by sample size and so does not require rarefaction. First, we tested for variation among habitats (lake, stream, estuary) while including a watershed random effect \((M8).\) Subsequently, we focus on lakes as the level of replication (omitting stream and marine sites). We tested the effect of geographic traits on \(\alpha_{ij}\) \((Prediction 5),\) using lake elevation, ocean distance and log lake size (surface area) as fixed effects \((M9).\) Ocean distance is as-the-fish-swims distance along the river basin between the focal lake and the ocean. For the subset of lakes with ecomorphology data, we used an additional regression to test whether \(\alpha_{ij}\) depends on population mean length, mean gape width, mean gill raker number, mean gill raker length and mean armor plating \((M10).\) For the subset of lakes that also have diet data, model \(M11\) compared \(\alpha_{ij}\) to mean diet \((\%\) benthic prey, prey richness and NMDS1 and 2). To test for genetic correlates of parasite richness \((Prediction 3),\) we started with a population-scale GWAS analysis, using regression to relate \(\alpha_{ij}\) to the population allele frequency of a focal SNP \((\text{with watershed as a random effect, } M12).\) Again, we used Holm corrections for multiple comparisons. We regressed \(\alpha_{ij}\) on population mean heterozygosity, \(H_j\) \((M13).\)

Previous analyses of microsatellites suggest that heterozygosity in these lakes depends on lake characteristics \((\text{Caldera and Bolnick 2008}),\) so we tested a final linear model relating mean heterozygosity to lake elevation, log area and ocean distance \((M14).\) Based on the results of models \(M9–M14,\) we switched to a path analysis to account for causal relationships between predictor variables. To build the path analysis model \(\text{(shown in Fig. 3),}\) we retained significant predictor variables from the linear models, keeping at least one variable per model \(M9–M14.\)

Results

Among-host and among-population variation in parasite richness

Per-fish parasite richness \((\alpha_j)\) varied from 0 to 10 species, with a mean of 2.28 species per host \((SD = 1.69, n = 4375\) fish; Fig. 1). Host population explained slightly less than half the variation \((\eta^2 = 0.426; M1: \text{host population effect, } p < 0.0001;\) Supplementary material Appendix 1 Table A3), with no significant watershed effect. The remaining variation arose from among-individual differences in richness. The prevalence and intensities of each parasite species, and variation in parasite community composition, are described in a separate paper \((\text{Bolnick et al. 2019}).\)

Parasite richness covaries with individual ecomorphology

Individual hosts’ parasite richness was correlated with individual traits. AIC for \(M2\) favored retaining fish length \((\text{standard normal; } Z = 11.02, p < 0.0001),\) sex \((Z = -1.22, p = 0.22),\) a random effect of population, and a sex×population random slope. Larger fish tended to have higher parasite richness. Sex was not a significant main effect in \(M2,\) but there was a strong and significant sex×population interaction \((p = 0.0011),\) i.e. the degree and direction of sexually dimorphic parasite richness varies among populations. Models with a length×population interaction were not favored by AIC model selection \((\text{Supplementary material Appendix 1 Fig. A4}).\)

At the individual host level, we found little evidence for ecomorphology or diet effects on parasite richness. Neither gill raker length, gill raker number, nor armor plate number correlated significantly with among-individual variation in parasite richness \((M3;\) Supplementary material Appendix 1 Table A3). Nor were there interactions between population and ecomorphology \((\text{LRT tests } p > 0.05).\) No diet trait affected host-level parasite richness \((M4;\) Supplementary material Appendix 1 Table A3). There were no diet trait×population interactions. Only host size had significant effects.

Genomic analyses of individual parasite richness yielded no significant SNP-richness associations. Of 39 039 SNPs tested with model \(M5, 1265\) were significant at \(p < 0.05\) and 21 at \(p < 0.001,\) but no more than false discovery rate expectations. We found no effect of individual genome-wide heterozygosity on parasite richness \((M6, \text{Poisson GLMM; } Z = -1.36, p = 0.174).\)

Sexual dimorphism in parasite richness

Parasite richness was significantly dimorphic in 17 of the 42 populations \((\text{Fig. 2A}).\) Of these 17 dimorphic populations, females had higher richness than males in 13 sites \((M2, \text{sex×site interaction, } \chi^2 = 21.68, \text{df} = 2, p < 0.0001)\). The magnitude and direction of sex dimorphism in parasite infection was predictable using ecologically relevant traits \((\text{Fig. 2B–C; } M7).\) Parasite richness was higher in whichever sex had larger body size \((t = 3.759, p = 0.0009)\) and whichever sex had lower diet NDMS2 scores \((t = -2.48, p = 0.021).\) These lower NDMS2 scores indicate greater chironomid larval consumption but fewer large prey like Gammarus, ceratopogonid larvae and stickleback eggs \((\text{Supplementary material Appendix 1 Table A3}).\) Diet NDMS1, which negatively covaries with the proportion of benthic prey \((p = -0.51, t = -16.8, p < 0.0001)\), was significantly dimorphic but did not predict dimorphism in parasite richness \((t = -1.2, p = 0.25).\)

Among-population differences in parasite richness

Mean per-fish parasite richness \((\alpha_j)\) spanned an order of magnitude across fish populations \((\text{Fig. 1C; } M1 p < 0.001)\). – 0.44 taxa per fish captured at Campbell River Point, to
4.72 in Gray Lake. Richness varied by habitat (M8, habitat effect $\chi^2 = 19.0$, df = 2, $p < 0.0001$), being on average 2.8-fold higher in lake stickleback than in anadromous fish ($\chi^2 = 14.29$, df = 1, $p = 0.0002$) and 1.8-fold higher than stream fish ($\chi^2 = 7.79$, df = 1, $p = 0.0052$). Stream stickleback had, on average, 1.5 times as many parasite taxa as marine fish ($\chi^2 = 3.63$, df = 1, $p = 0.056$). Watershed had no detectable effect, so we do not consider it further.

Lake geography partly influenced among-lake variation in mean parasite richness (M9). We found no support for effects of elevation ($t = -0.58$, $p = 0.56$) or lake depth ($t = 0.6$, $p = 0.55$), a weak support for an effect of log surface area ($t = -1.84$, $p = 0.076$), and a significant positive effect of ocean distance ($t = 2.56$, $p = 0.016$; Supplementary material Appendix 1 Fig. A5). A few post hoc observations are worth noting. First, although lake depth had no linear effect...
on parasite richness, visual inspection of the data suggested a quadratic relationship (depth $t = 4.18$, $p = 0.0001$, depth$^2$ $t = -3.7$, $p = 0.0006$, Supplementary material Appendix 1 Fig. A6). Log lake surface area likewise had a quadratic effect on mean richness (area $t = 2.11$, $p = 0.043$; area$^2$ $t = -2.64$, $p = 0.013$, Supplementary material Appendix 1 Fig. A7). Last, although elevation had no main effect in M9, inspection of the data suggested an interaction with lake area, which was confirmed with a post hoc linear model (elevation $t = 3.82$, $p = 0.0006$, log surface area $t = 3.00$, $p = 0.0055$; elevation $\times$ area $t = -3.46$, $p = 0.0016$, Supplementary material Appendix 1 Fig. A8).

We found support for some host-population trait means explaining among-lake differences in parasite richness (M10, fish length $t = 2.8$, $p = 0.008$; size-adjusted gape width $t = 2.06$, $p = 0.048$; gill raker number $t = 1.87$, $p = 0.072$; size-adjusted gill raker length $t = -0.026$, $p = 0.979$). AIC supported a simpler model in which mean parasite richness increased with
mean fish length and gape width, and decreased with mean gill raker number (all typical benthic ecomorph traits). More benthic-feeding populations (higher NMDS1) tended to have more parasites per fish but this was marginally significant ($p = 0.07$, M11). NDMS2 and mean prey richness had no effect ($p = 0.42$ and 0.41 respectively).

Genome-wide association mapping (M12) revealed no significant (after Holm correction) SNP effects on host-population parasite richness. Mean genome-wide heterozygosity had no detectable association with population mean parasite richness (M13, $r = 0.925$, $p = 0.364$).

The variables considered above are likely to be inter-related. For example, heterozygosity increases with lake area (Caldera and Bolnick 2008) but decreases with distance from the ocean and elevation (M14; $p = 0.029$, 0.012, <0.001 respectively).

To account for this covariance among predictors, we used path analysis to partition direct and indirect effects (Fig. 3, 4, Supplementary material Appendix 1 Table A4), taking the significant effects from the $\mathbf{R}_{ij}$-based linear regressions above. The path analysis explained 45.6% of the variation in mean per-fish parasite richness among lakes. Lake area had no significant direct effect on parasite richness ($r = 0.053$, $p = 0.822$), but had indirect effects via fish heterozygosity and diet. Specifically, larger lakes have more genetically diverse fish ($r = 0.366$, $p = 0.011$), and heterozygosity has a positive effect on parasite richness ($r = 0.360$, $p = 0.038$). Stickleback in larger lakes also consumed a smaller proportion of benthic prey ($r = -0.777$, $p < 0.001$), which conferred higher parasite richness ($r = 0.570$, $p = 0.013$). The indirect negative effect of lake size via fish diet exceeded its positive effect via fish heterozygosity ($r = -0.311$ versus 0.130). In simple bivariate correlation tests, lake distance from ocean had a positive effect on parasite richness ($r = 0.310$, $p = 0.017$). However, this positive correlation is mediated via an indirect positive effect of ocean distance on heterozygosity (Supplementary material Appendix 1 Table A4; $r = 0.553$, $p = 0.002$), which has a positive partial correlation with parasite richness. Lakes farther from the ocean also had larger fish ($r = 0.52$, $p = 0.001$), but in this analysis, mean per-lake fish size had no further relevance to fish diet ($r = 0.16$, $p = 0.111$) or parasite richness ($r = 0.075$, $p = 0.635$). Last, higher elevation lakes had less heterozygous fish ($r = -0.74$, $p < 0.001$), indirectly reducing richness.

**Discussion**

Parasites form diverse multi-species assemblages in nested metacommunities across individual hosts, and among host populations (Seabloom et al. 2015). Key questions in parasite ecology, and ecology more broadly, are 1) what processes dictate this metacommunity diversity, and 2) are these processes scale-dependent? Here, we present a case study using macroparasites from a stickleback metapopulation to show that parasite richness varies widely among host individuals and across host populations. In both cases, the variation in richness spans an order of magnitude. We have identified ecological factors contributing to this variation in parasite diversity and have shown that mostly different factors act at small and large spatial scales (Fig. 5).

**Individual scale: why does parasite richness differ among host individuals?**

Individual stickleback hosts within a population experience approximately similar abiotic conditions and biotic prey communities. Thus, variation in parasite richness among host individuals is either stochastic or due to differences in individual traits. We confirmed that stickleback individual size, gape width, diet and sex were correlated with individual parasite richness, in sometimes subtle ways, but did not find any effect of genotype or heterozygosity.

Larger individual stickleback carried more parasite taxa, as in other fish species (Calhoun et al. 2018). Larger fish may be older, having more time to accrue infections or having senescing immune systems (Zelmer 2014). Or, larger stickleback might be capable of eating more prey due to larger stomachs, increasing overall intake rates of parasites. A third possibility is that larger fish eat particular kinds of prey that promote parasite diversity. For instance, in stickleback and many other fish, larger individuals are more benthivorous. Although we do not have an a priori expectation that benthic-feeding fish would have greater parasite richness, we do also find that fish with larger gaps (a typical benthic trait) also have higher parasite diversity. Moreover, our sexual dimorphism analysis suggests that parasite richness is higher in whichever sex has the more benthic diet and is larger. Thus, we conclude that individual size and trophic ecology affect individual hosts’ parasite richness. This confirms that habitat patch characteristics (i.e. individual host traits) modify community assembly, likely by modulating the rate of colonization.

Diet effects are consistent with previous studies in other species that found that individual diet influences...
In stickleback, we found that sex-biased parasite richness covaries with sexual dimorphism in ecomorphology traits, implying a role for encounter filters. Across all lakes, females tended to be larger than males, consume more large prey, and have higher parasite richness. In lakes with larger dimorphism in body size or diet, parasite richness was also more dimorphic. The exceptions support our conclusions: in a few lakes where diet or size dimorphism is reversed, parasite richness dimorphism is also reversed. Sexual dimorphism in immune function (Zuk 1996, Nunn et al. 2009), which we did not measure, could contribute to the residual variation in sex dimorphism of parasite richness.

We found no single-locus or genome-wide genetic control of per-fish parasite richness ($\alpha_i$). This is perhaps because parasite richness is an aggregate measure arising from many parasite species’ interactions with the host, which might be controlled by separate gene(s). In the companion study (Bolnick et al. 2019), we report many genetic associations with infection by each common parasite. But, because these associations tend not to involve the same genes, there is no overall genetic effect on parasite richness.

**Population scale: why does parasite richness differ among host populations?**

Parasite richness varied substantially among host stickleback populations with habitat, geography, ecomorphology and genetic diversity. The strongest effect was habitat: parasite richness was nearly three times higher for lake stickleback...
than their anadromous relatives; stream fish were intermediate, consistent with previous studies (MacColl and Chapman 2010, Eizaguirre et al. 2011, 2012). Given that stickleback are relatively recent (post-glacial) colonists of freshwater, this higher infection rate indicates that the colonization process increased infection risks (Weber et al. 2017), rather than providing a means of escape from historical enemies (Grunberg et al. 2019). It remains unclear to what extent these habitat differences are a function of abiotic conditions, availability of intermediate or terminal hosts, or habitat differences in dilution effects (e.g. higher non-host fish species richness in the ocean; Becker et al. 2014).

For lake populations, lake geography affected parasite richness, which was higher in mid-sized lakes (with intermediate depths), farther from the ocean, and at higher elevation. Some geographic effects acted indirectly via host population traits. For example, stickleback in larger lakes have more limnetic diets on average (Lavin and McPhail 1986), and this diet shift is associated with reduced parasite richness; i.e. lake size had a net negative indirect effect on parasite richness via diet. At the time we sampled parasites, average water temperature in the top 10 m of water was not correlated with lake size, depth and elevation, but we cannot rule out that more generally water temperature might be a causal link between lake geography and parasite community diversity.

This negative effect was lessened somewhat by a positive but weaker indirect effect of lake size through host heterozygosity. Larger lakes support more genetically diverse stickleback populations (Caldera and Bolnick 2008). This increase in mean heterozygosity was associated with higher mean per-fish parasite richness, inconsistent with the oft-cited immunological benefit of genetic diversity (Poulin et al. 2000, Joly et al. 2008, Kaunisto et al. 2013). This surprising effect might be explained if we consider a reversed cause-effect relationship: richer parasite communities might select for greater genetic diversity. But, such selection is unlikely to affect genome-wide heterozygosity, because 1) most SNPs should be effectively neutral, and 2) heterozygosity is mostly a function of lake geography.

Heterozygosity was also influenced by lake elevation and distance to ocean. Higher-elevation lakes tend to have lower heterozygosity, consistent with expectations of a stronger bottleneck during colonization. Path analysis suggests that elevation reduces parasite richness indirectly via reduced heterozygosity, but there was no direct elevation-richness
correlation. Likewise, lake distance from the ocean only had an indirect effect on richness via heterozygosity.

Our results do not conform to general predictions from island biogeography and basic metacommunity theory. All else equal, communities should be more diverse on habitat patches that are larger, or closer to other such patches (MacArthur and Wilson 1963), as demonstrated for human parasites on islands in Jean et al. (2016). Larger lakes are akin to larger islands, but in our study, lake size had no net relationship to parasite richness due to conflicting effects via host diet and heterozygosity. Furthermore, we find that more isolated stickleback populations with longer distances to the ocean have richer, rather than poorer, parasite communities. We hypothesize that these unexpected effects could be driven by host evolutionary genetics.

Scale-dependent and -independent factors affecting parasite richness

To what extent do the variables structuring parasite diversity generalize across scales (Fig. 5)? Only two traits showed similar effects across scales. Larger individual fish had richer parasite infections, and populations with larger average fish had higher mean parasite richness. We also detected a marginally significant positive effect of gape width on parasite richness at both scales, which suggests the trend at each scale is real rather than type I error. These effects of size and gape confirm Prediction 1.

Direct measures of diet had inconsistent effects across spatial scales. At the larger spatial scale, more benthivorous (lower NDMS1) stickleback populations had richer average per-fish parasite communities. No such trend was found at the individual scale. Diet sexual dimorphism (acting within populations) did correlate with parasite richness dimorphism, but this involved NDMS2. Thus, diet does influence parasite richness, but does so inconsistently across scales. This does not support Prediction 2.

A scale-specific relationship was also seen between genomic heterozygosity and parasite richness. Some studies suggest that heterozygosity might increase a host’s repertoire of immunological tools, thereby lowering richness (Coltman et al. 1999, Arkush et al. 2002, Whitehorn et al. 2011). Indeed, fish species with higher heterozygosity had fewer parasite species (Poulin et al. 2000, Joly et al. 2008, Kaunisto et al. 2013). Our data did not support such a trend at the individual-level, despite high statistical power, refuting Prediction 3. The lack of an individual-level effect undermines the notion that heterozygosity acts through individual immunogenetic diversity. We did, however, find a population-level effect of heterozygosity, though in an unexpected direction (parasite richness increased with host population mean heterozygosity).

So, why does heterozygosity matter at the population scale? One possibility is that larger host populations can both support more diverse parasite communities and maintain higher heterozygosity. Parasite diversity can be related to host population size, following a species–area relationship (Bagge et al. 2004, Zelmer 2014). However, our path analysis found no support for a direct effect of lake size on parasite richness. An alternative explanation is that fish species exposed to more diverse parasites might be subject to stronger balancing selection and evolve higher heterozygosity (Hamilton 1982, Poulin et al. 2000, Berenos et al. 2010). But, this adaptationist interpretation should not impact genome-wide heterozygosity, only those loci involved in immune defense and sites linked to them.

Conclusions

Because parasites are such a large component of biological diversity and have large effects on the communities in which they are embedded (Lafferty et al. 2008), the field of ecology needs to understand the biological processes regulating the distribution, abundance and diversity of parasites. Here, we present a case study illustrating the highly multivariate and scale-dependent nature of these processes. A companion paper shows similarly scale-dependent and multivariate effects on the prevalence, and covariance, between particular parasite species that constitute the communities examined here (Bolnick et al. 2019). The macroparasite community of threespine stickleback is structured by among-individual variation in host sex, size, morphology and diet, and among-population differences in host size, morphology, diet, genetic diversity and habitat. Some of these variables (size, gape width) act consistently across individual- and population-scales. In general, it seems that more benthivorous stickleback have richer parasite communities, a trend observed between sexes, and among populations. Other variables are scale-dependent and contribute to parasite differences only among individual hosts (sex) or only among host populations (heterozygosity, habitat). Such scale-dependent effects are expected in metacommunities of all kinds (Leibold et al. 2004). This scale-dependence may help explain inconsistent results among studies conducted at disparate scales. The implication of these findings is that studies of species distributions and abundances need to draw scale-specific inferences.

Data availability statement


Acknowledgements – We thank C. Harrison, T. Rodbumrung and T. Ingram, for assistance with field work. J. Day assisted with parasite data collection. R. Grunberg commented on the manuscript. Funding – The research was supported by the Howard Hughes Medical Institute through an Early Career Scientist position to DIB, NSF (DEB-1144773) to DIB, and a NSF Graduate Research Fellowship to EJHR. Conflict of interest – No conflicts of interest. Author contributions – DIB conceived of and designed the study, analyzed data and initially drafted manuscript. EJR contributed to writing and figure design. KB, YES and WES contributed to field sampling, data collection, and writing.
References

Bagge, A. M. et al. 2004. Fish population size, and not density, as the determining factor of parasite infection: a case study. – Parasitology 128: 305–313.
Appendix 1.

04994 at Supplementary material (available online as Appendix ecog-04994 at <www.ecography.org/appendix/ecog-04994>). Appendix 1.