Title: What controls the range of hosts a fish parasite infects?

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1 Abstract

² Keywords

- 3 FishPEST, species distribution model, boosted regression tree,
- 4 Take-home messages
- 1. It is possible to predict parasite niche breadth using either parasite community
- similarity. This means that freshwater fish parasites are not random assemblages,
- but that parasites with similar niches infect similar host species. It is possible
- that parasite community information condenses host trait variation, (perhaps)
- evolutionary history, and geographic location (to some extent).
- 2. Predictive accuracy does not vary as a function of host specificity (though I only
- consider parasites with 20 or more occurrence records, but these could all be on
- the same host species).
- 3. It may be possible to predict parasite spillover from invasive hosts to native host
- communities, or to predict biotic resistance of a community to invasion.

Introduction

- 16 Host-parasite relationships are complex, intimate (non-neutral) interactions
- with lots of impacts

- Most of parasite population and community ecology is about looking for
 patterns, and lots of folks don't find them, and have called host-parasite
 interactions neutral, or random Parasite communities are sometimes conserved
 across host species, such that the presence of one parasite may increase the likelihood
 of finding another parasite species.
- Being able to predict parasite occurrence is pretty important, for a number of reasons (species invasions/biotic resistance, spillover to human hosts, etc.
- **Previous work and knowledge gap** One of the largest factors holding predictive models of parasite distributions among potential hosts is the relative paucity of data (but see). However, this barrier is being overcome both by scientists Strona and Laf-27 ferty (2012); Nunn and Altizer (2005) and museums (Gibson et al. (2005)). Studies uti-28 lizing these large datasets have largely asked questions about parasite co-occurrence 29 patterns Strona et al. (2013), or the factors influencing parasite sharing Braga et al. (2014); Dallas and Presley (2014). These studies largely examine parasite community 31 composition, and determine the influence of host traits or phylogenetic relationships on 32 parasite community composition. However, almost no studies have attempted to predict what host species a parasite will infect. Previously, Strona et al. (2013) developed a framework to predict parasite co-occurrence likelihood given host community, and habitat variables. This approach is predicated on the idea that information on hosts and geography are what determine the likelihood of a parasite infecting a given host.
- Thesis (what I did, what I found) Here, we test this idea by quantifying the predictive capability of models trained on host traits, geographic variables, or parasite community variables. To do this, we examined a large dataset on interactions between freshwater fish and their parasite communities Strona and Lafferty (2012). The number and identity of host species that a given parasite could infect may be constrained by geographic location, host trait variables (i.e. patch quality), or the existing parasite community of the given host species (i.e. parasite community structure). We trained

boosted regression models on each of these three variable types, predicting the potential distribution of 238 parasite species. Parasite species distributions were most constrained by the existing parasite community, as this model allowed for highly accurate
prediction of parasite occurrence on a set of hosts. This suggests that either parasite
community composition contains information about host susceptibility to infection,
and potentially information on the relevant ranges of host and parasite, serving as a
measure of geographic location. Taken together, this suggests that parasite community
composition can determine parasite occurrence in a host community. This has important applications to the study of invasive species, as it may be possible to predict what
parasites will spillover to other hosts in the case of a non-native host introduction, or
the biotic resistance of the native host community, as the native parasite community
may be capable of infecting the non-native host.

57 Methods

Data and processing We use an existing global database of fish-parasite associations (hereafter referred to as FishPest; citepstrona2013) consisting of over 38000 helminth parasite records spanning a large diversity of parasites (Acanthocephala, Cestoda, Monogenea, Nematoda, Trematoda). Many of these entries represent isolated occurrences of parasites on hosts. In order to allow for cross-validation and accurate prediction, we constrained our analyses to parasites with a minimum of 20 host records. In other words, we only examined parasites that had been recorded more than 20 times, but these occurrences could be on fewer than 20 host species. The inclusion of duplicate occurrences was only permitted if the parasite was recorded on a host in a different geographic location, based on latitude and longitude values. This resulted in a total of 238 parasite species. Our response variable was parasite occurrence (binary), and was predicted using three classes of variables, representing host life history traits, geographic location, and parasite community similarity (Table 1).

Values of predictor variables were obtained largely through the FishPest database Strona

and Lafferty (2012); Strona et al. (2013), supplemented by variables obtained from FishBase Froese and Pauly (2010). However, there were still some missing predictor variable values. Missing predictor variable values were imputed using the imputation 74 procedure in the randomForest R package (Liaw and Wiener (2002)). Details of host 75 trait and geographic variable determination are provided in Strona et al. (2013) and Table 1. Parasite community similarity was considered as a the first five principal components from a principal components analysis on the host-parasite matrix. This matrix 78 contains information on all parasites infecting all host species, except with the parasite species of interest removed. Thus, the principal components represent a measure 80 of parasite community similarity among host species without any information about host range of the parasite species under consideration. In addition, we included para-82 site species richness of a host as a predictor variable.

Predictive model formulation Here, we used boosted regression trees to predict parasite occurrence among potential host species for each of our 238 parasite species. Regression tree analysis is an extremely powerful tool for prediction and feature selec-86 tion, bypassing many of the issues of simple regression models (e.g. multicollinearity, nonlinear relationships) Elith et al. (2008); Dallas and Drake (2014). Boosting refers to 88 the process of creating a large number of regression trees, and weighting them by their predictive power to extract general weak rules, which are then combined to enhance 90 predictive ability. The optimal number of trees was determined using the out-of-bag (00B) estimation procedure, with the upper limit set to 50000. Other pertinent param-92 eters include the learning rate (l = 0.001), which controls the degree each new tree contributes to the overall model, and interaction depth (id = 4), which allows for up to 94 four-way interactions among predictor variables. The absence of a recorded interaction between host and parasite does not mean that the parasite does not infect that host. Borrowing from the idea behind Maximum Entropy modeling, we sampled the data to obtain background interactions, which we define here as a set of possible interactions between host and parasites. This background set was not composed of the entire dataset, but rather a sample of five times the num-100

a training set, 70% of the data used to train the boosted regression tree model, and 102 a test set, the remaining 30% of the data used to test the predictive accuracy of the 103 trained model. 104 From the final boosted regression tree models, we are able to extract variable relative 105 contribution (RC) measures, which provide information about the importance of each 106 variable to the final model predictions. Relative contribution values for each predic-107 tor variable was determined by permuting each predictor variable and quantifying the 108 reduction in model performance, a method that is free of classical assumptions about 109 normality and equal variance (Anderson 2001). Relative contribution estimates were then based on the number of times a given predictor variable was selected for splitting, 111 weighted by the degree the split improves model performance, and scaled between 0 (no contribution) to 100 (maximum contribution). 113 All models were compared to a random null model, which randomized occurrence values in the test dataset, but kept them constrained to the total number of occurrences. 115 Model performance was assessed using receiver operating characteristic curves, which 116 relate true positive and false positive (type I error) rates graphically. The area between 117 the curve generated by true and false positives and the 1:1 line from the origin gives a measure of predictive accuracy. 119

ber of positive occurrence records for a given parasite. These data were subset into

Results

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Importance of host traits, geographic variables, and parasite community similarity. All models performed better than our null predictions (null model \overline{AUC} = 0.50). With varying degrees of accuracy, models were able to predict parasite occurrence for the 238 parasite species examined using host traits (\overline{AUC} = 0.66), geographic variables (\overline{AUC} = 0.79), and parasite community similarity (\overline{AUC} = 0.88). The full model containing all variables was able to successfully predict parasite occurrences in the hold-out test dataset with high accuracy (\overline{AUC} = 0.89), only marginally more ac-

curate than the model trained with only parasite community similarity variables (Figure 1). The relative contribution (RC) values for each separate model, and the model 129 trained on all available data are provided in Figure 2. 130 In the full model, variables of different classes were allowed to have different relative 131 contributions, which allows for the determination of variables driving the predictive ac-132 curacy of the full model. For instance, relative contribution values were largest for the 133 parasite community similarity values obtained from the principal components analysis 134 on the host-parasite network with the parasite species of interest removed (Figure 2), 135 with the five PCA vectors comprising around 52% of the relative contribution values, 136 and four of the top 5 predictive variables. On the other side of the predictive spectrum, 137 host trait variables, specifically host age at maturity, lifespan, and growth rate, con-138 tributed very little to model performance.

Was predictive ability influence by parasite ecology? The relative importance 140 of variable classes, or the general predictive power of the trained model, may differ as a function of parasite taxonomic group or host specificity. We tested for variation in pre-142 dictive power among parasite taxonomic group (Acanthocephalans, Cestodes, Monogeneans, Nematodes, and Trematodes) and as a function of the number of host specificity. 144 Here, we defined parasite host specificity as the number of hosts a parasite infects. We failed to detect evidence that parasite taxonomic group (Figure S1) influenced predic-146 tive power in any of our trained models. We did, however, observe an effect of host specificity (Figure S2), as predictive accuracy became more variable as host specificity 148 increased (i.e. the number of hosts a given parasite infected became smaller). Despite this variability, the the mean predictive accuracy over a range of host specificity values 150 remained constant (Figure S2). 151

Discussion

53 State most important findings

- what does this mean? Tie into concepts of invasion and biotic resistance. Also emphasize that this suggests parasites are likely not random assemblages, but can be
- predicted with limited accuracy based on host traits, and with much greater accuracy
- using only data on parasite community composition.
- PSR was lowest predictor of that class of variable (parasite community vars). If PSR
- was driving patterns, then neutrality could still be on the table, since new parasites
- would be predicted to occur on hosts with a large number of parasites. This preferen-
- tial attachment could be based on random network (small world) formation, and psr is
- 162 likely linked to host abundance.

163 even more general implications

164 Acknowledgements

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189 Tables

Table 1: Description and units of variables used to predict parasite occurrences.

Variable	Units	Description	Range
Max length	cm	Maximum fish species length	1 - 2000
Trophic level	_	1 + mean trophic level of food	2-5
Age at maturity	years	Age at sexual maturity	0.1 - 34
Life span	years	Estimated maximum age	0 - 145
Growth rate	$years^{-1}$	Rate to approach asymptotic length	0.02 - 9.87
Geographic region	_	Biogegraphic region	_
Area of occupancy	No. $1x1 \circ cells$	Global host distribution	1 - 1610
Latitude	max - min degrees	Latitudinal distribution	1 - 148
Longitude	max - min degrees	Longitudinal distribution	1 - 359
Principal components	_	PCA axes of host-parasite network	-11.7 - 9.8
Parasite species richness	#	No. parasite species of host species	0 - 89

Figures 190

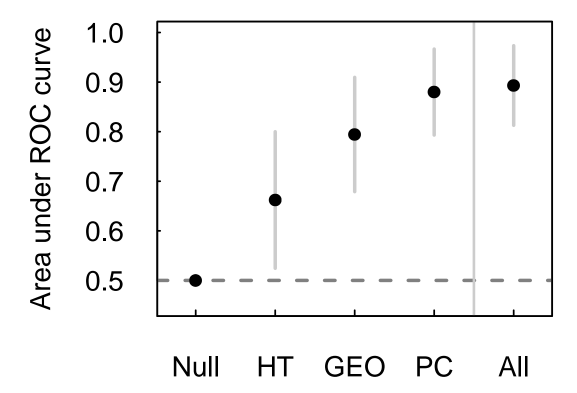


Figure 1: Accuracy (Area under Receiver operating characteristic (ROC) curves) for our predictive models incorporating host traits ('HT'), our null model ('Null') that maintained interaction number (number of occurrence records), but assigned occurrences equiprobably among potential interactors.

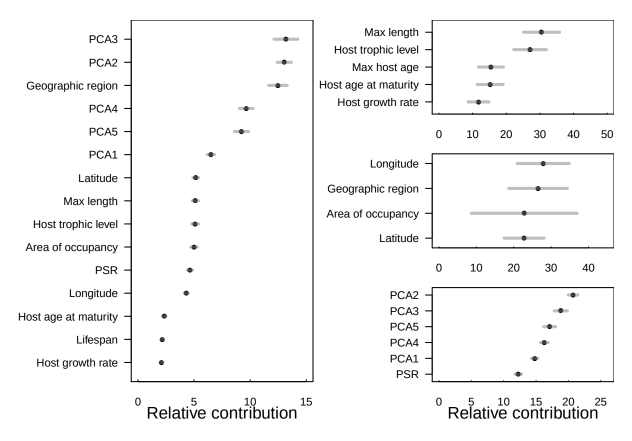


Figure 2: The average relative contribution values from the boosted regression tree models trained on all available data (left), host trait data (top right), geographic variables (middle right), and parasite community similarity (bottom right). Variables named "PCA" are principal components axes, and "PSR" refers to parasite species richness. Other variable definitions and units are available in Table 1.

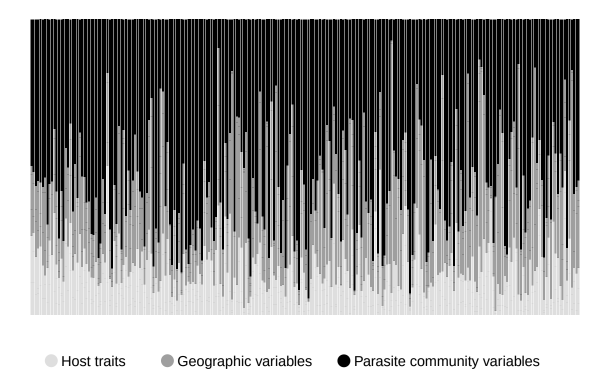


Figure 3: Relative contribution values for variables of one of three classes; host traits, geographic variables, or parasite community variables. Each column represents a model trained on occurrence data for a single parasite species.