The influence of variability in species trait data on community-level ecological prediction and inference

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Abstract
Species trait data have been used to predict and infer ecological processes and the responses of biological communities to environmental changes. It has also been suggested that, in lieu of trait, data niche differences can be inferred from phylogenetic distance. It remains unclear how variation in trait data may influence the strength and character of ecological inference. Using species-level trait data in community ecology assumes intraspecific variation is small in comparison with interspecific variation. Intraspecific variation across species ranges or within populations may lead to variability in trait data derived from different scales (i.e., local or regional) and methods (i.e., mean or maximum values). Variation in trait data across species can affect community-level relationships. I examined variability in body size, a key trait often measured across taxa. I collected 12 metrics of fish species length (including common and maximum values) for 40 species from literature, online databases, museum collections, and field data. I then tested whether different metrics of fish length could consistently predict observed species range boundary shifts and the impacts of an introduced predator on inland lake fish communities across Ontario, Canada. I also investigated whether phylogenetic signal, an indicator of niche-conservatism, changed among measures. I found strong correlations between length metrics and limited variation across metrics. Accordingly, length was a consistently significant predictor of the response of fish communities to environmental change. Additionally, I found significant evidence of phylogenetic signal in fish length across metrics. Limited variation in length across metrics (within species), in comparison with variation within metrics (across species), made fish species length a reliable predictor at a community-level. When considering species-level trait data from different sources, researchers should examine the potential influence of intraspecific trait variation on data derived by different metrics and at different scales.

Introduction
Ten years ago, McGill et al. (2006) suggested “rebuilding” community ecology by examining how species traits control the relationship between a species’ niche and its environment. Since then, species traits have been used to build mechanistic inferences on topics ranging from community assembly (HilleRisLambers et al. 2012), to biodiversity patterns (Meynard et al. 2011; Swenson et al. 2011), to ecosystem functioning (de Bello et al. 2010; Flynn et al. 2011), to the impacts of environmental change (Angert et al. 2011). Further, despite criticism, phylogenetic distance has been used as a proxy for differences in species traits or niche differences (HilleRisLambers et al. 2012; Gerhold et al. 2015). This use assumes phylogenetic signal or phylogenetic niche-conservatism (although meta-analyses have suggested phylogenetic signal is not as common as previously believed; Kelly et al. 2014).

Intraspecific trait variation is known to have significant ecological and evolutionary effects (Bolnick et al. 2011) and not incorporating such variation into trait-based studies can influence the accuracy of predictions in community ecology (Albert et al. 2012; Violle et al. 2012). However, data quantifying community-wide intraspecific trait variation are often not available or difficult to attain,
particularly for studies at regional scales. Therefore, species-level trait values are often substituted. The use of species-level trait data for community-level inference relies upon the assumption that intraspecific variation is low, particularly in comparison to interspecific variation (McGill et al. 2006). We know, however, this is not always the case, particularly for species distributed across environmental gradients (Albert et al. 2010). In part due to intraspecific trait variation, species trait data may vary significantly when gathered from field observations, experiments, literature (e.g., field guides or atlases) or from taxonomically or regionally distinct online databases. Few studies have examined the consistency of species trait data across sources (Fitzsimmons 2013; Kazakou et al. 2014); although some work has been done on sensitivity to missing data and variable sampling effort (Pake- man 2014; Sandel et al. 2015). Trait values are often presented as “mean,” “common,” or “maximum” values and sensitivity to type of metric for single traits remains unclear. Further, individual sources offer little detail on variation in species traits through space or time, although the importance of this variation has been recognized (Fitzsimmons 2013; Kazakou et al. 2014).

I use species data for a single trait, body size, collected from six sources (online, print, collection and field based) to examine how variability in species-level trait data influences trait-based predictions in community ecology. I also test whether variability in trait data influences our ability to detect phylogenetic signal and infer the usefulness of phylogenetic relatedness as a proxy for ecological relatedness. Size (e.g., length, volume or mass) is perhaps the most commonly measured individual trait across taxa. It influences many aspects of ecology and evolution and is correlated with other species traits including those which determine metabolism, reproductive rate, dispersal, and trophic interactions (LaBarbera 1989; Brown et al. 2004; Woodward et al. 2005). Further, body size has been used to infer the direct and indirect impacts of human activities including hunting and fishing (Jennings and Blanchard 2004; Fenberg and Roy 2008), land-use changes (Mulder and Elser 2009), introduced species (Ness et al. 2004; Alofs and Jackson 2015), and climate warming (Gardner et al. 2011; Alofs et al. 2014).

Here, I focus on how variation in freshwater fish body size, particularly species length metrics, may influence community-level inference. Many freshwater fishes demonstrate significant intraspecific trait variation within populations, across species ranges and along environmental gradients (e.g., Einum and Fleming 2002; Heins et al. 2004; Blanck and Lamouroux 2006; Gutowsky and Fox 2012). Across species, length is related to a range of life-history traits including fecundity and longevity (Wine-miller and Rose 1992; Alofs et al. 2014). Alofs et al. (2014) demonstrated there was a significant relationship between average body length in Ontario, Canada (as reported by Holm et al. 2009), and the magnitude of northern range boundary shifts by warm- and coolwater-adapted fishes in that province over about 30 years. Specifically, larger predatory fishes demonstrated greater poleward shifts than smaller prey species (including both warm- and coolwater-adapted fishes). In Ontario, resident populations of small prey fishes also appear to be more vulnerable to the introductions of these large range-expanding predators as they establish in northern lakes (Alofs and Jackson 2015). Particularly, average Ontario length is significantly related to the relative risk imposed by introductions of Amboplites rupestris (Rock Bass), a gape-limited predator.

Intraspecific variation across species ranges and within populations can produce variability in trait data measured at different scales or by different methods. The relative amount of intraspecific variation, in contrast to interspecific variation, may control the usefulness and reliability of trait data. I used data on inland lake fish communities across Ontario (from Alofs et al. 2014 and Alofs and Jackson 2015) and a phylogeny of Ontario fish communities (Doyle 2013) to test the influence of variability in species length metrics on community-level inference. I hypothesized that, given variability along length metrics, the strength of ecological prediction and inference would vary by the method and the scale of data collection. Specifically, prediction and inference would be strengthened by using (1) species averages or common estimates of traits rather than species maximums or records, which may represent outliers, and (2) using species data derived in Ontario rather than at larger scales. I test this idea using 12 metrics of fish length (collected from a variety of sources) to examine (1) the relationship between size and the extent of northern range boundary shifts by fishes, (2) the relationship between size and the vulnerability of common species to the introduction of a predatory fish, and (3) evidence of phylogenetic signal in size.

Materials and Methods

Data

I collected 12 different metrics of fish length for 40 species (Table 1, Supporting Information, Appendix S1; species were selected based on community-level analysis described below) from two online sources, two print sources, the Royal Ontario Museum collection (Fig. 1) and field data collected by the Ontario Ministry of Natural Resources and Forestry during their Broad-scale Monitoring (BsM) program (Sandstrom et al. 2010). The BsM field data include length data for more than 250 thousand
individual fishes from standardized sampling of 745 lakes across Ontario. The 12 length metrics included five “average,” “common,” or “median” and seven “maximum” or “record” metrics. Metrics referred to as “average” or “common” in print and online databases were not clearly described in sources and are likely to be based on expert opinion rather than calculated from sampled data (E. Holm, Royal Ontario Museum, pers. comm.). Most metrics were based on measures of total length (from tip of snout to tip of closed caudal fin); however, during BsM, fork length (from tip of snout to center of open caudal fin) was measured. Fork length is correlated with total length (Carlander and Smith 1945) although shorter than total length particularly for species with lobed caudal fins, and at most equal to total length.

Five missing values were substituted from alternative sources to complete the data set: world record length for Coregonus artedi was not reported in the Freshwater Fishes of Ontario, and thus, the Ontario record, maximum in the data set, was used. The common length of Osmerus mordax was not reported in Fishbase and the average common length from the Freshwater Fishes of Canada was substituted. The Freshwater Fishes of Canada did not report a maximum length for Chrosomus neogaeus, Notropis heterodon and N. volucellus and the world record reported in the Freshwater Fishes of Ontario was substituted.

I collected the relative northern range boundary shift of 13 species calculated as the change in mean latitude of most northern 20% of occurrences in a data set of 1527 lakes across Ontario that were sampled during a historical and contemporary period ~30 years apart (from Alofs et al. 2014; Table 1, Supporting Information, Appendix S1). I also collected data on the vulnerability of 29 species to Rock Bass introductions; the relative risk ratio associated with these predator introductions was calculated from two-by-two contingency tables created from a data set of 1551 Ontario lakes also sampled in two time periods (from Alofs and Jackson 2015; Table 1, Supporting Information, Appendix S1). The relative risk ratio measures the impact of introduction as the probability of each resident species “loss” (presence during a historical survey and absence during a contemporary survey) given introduction and establishment of Amboplites rupestris (Rock Bass) over the probability of loss given no establishment, that is, background variation. Finally, I used the phylogenetic tree of 26 species occurring in lakes on Manitoulin Island, Ontario, published by Doyle (2013);

Table 1. Description of 12 measures of fish length collected from various sources.

| Code     | Source                  | Database type | Measure type | Length measure |
|----------|-------------------------|---------------|--------------|----------------|
| FB.MAX   | FishBase                | Online        | Maximum      | Total length   |
| FB.COM   | FishBase                | Online        | Common       | Total length   |
| FT.MAX   | FishTraits Database     | Online        | Maximum      | Total length   |
| FC.MAX   | Freshwater Fishes of    | Online        | Maximum      | Total length   |
| FC.COM   | Canada (Scott and Crossman 1998) | Print         | Common (or maximum of common range) | Total length |
| FO.WORLD.MAX | Freshwater Fishes of    | Print         | World record | Total length   |
| FO.ON.MAX | Ontario (Holm)         | Print         | Ontario record | Total length   |
| FO.ON.AVG | et al. 2009            | Print         | Ontario average | Total length   |
| ROM.MAX | Royal Ontario Museum    | Collection    | Maximum in all catalogued lots | Total length |
| BSM.MAX | OMNRF Broad-scale Monitoring | Field   | Absolute maximum of all measured fish | Fork length |
| BSM.AVG  | Field       | Fork Length   | Maximum per lake averaged across lakes | Fork Length |
| BSM.MED  | Field       | Fork Length   | Median of all measured fish | Fork Length |

1Fork Length was reported for Walleye.
species indicated in Table I, Supporting Information, Appendix S1).

**Analysis**

I calculated correlations between each of the 12 metrics of fish length. To compare variation among metrics of length to interspecific variation in length, I calculated the coefficient of variation (CV) for each species (across metrics) and each metric (across species). For 13 species, I fitted separate linear regression models of relative northern range boundary shifts against each of the 12 length metrics. For 29 species, I fitted similar linear regression models of the relative risk associated with predator introductions against each of the length metrics. For all of these models, I examined standardized (z-score) effect sizes, $R^2$, and $P$-values. All size metrics and the relative risk ratios were natural-log-transformed for analysis. All analyses were performed with R 3.1.2 (R Development Core Team 2014).

To test for evidence of phylogenetic signal across 26 species, I calculated four commonly used indices (described and compared in Münkemüller et al. 2012): Moran’s $I$, 

![Figure 2. Correlation matrix for 12 metrics of fish species length for 40 species.](image-url)
Abouheif's $C_{\text{mean}}$ Pagel's $\lambda$ and Blomberg's $K$. Moran's $I$ and Abouheif's $C_{\text{mean}}$ are autocorrelation indices and not based on an evolutionary model. Stronger phylogenetic signal is indicated by greater deviation from zero by these estimates. Pagel’s $\lambda$ and Blomberg’s $K$ are based on a Brownian motion model of trait evolution. Similar to $I$ and $C_{\text{mean}}$, $\lambda$ and $K$ approach zero with phylogenetic independence. The upper limit of $\lambda$ is near one, while $K$ may exceed one indicating greater phylogenetic signal than expected with Brownian motion. Moran’s $I$ and Abouheif’s $C_{\text{mean}}$ were estimated and their significance in comparison with random trait variation tested with Monte Carlo simulations (999 randomizations) using the abouheif.moran function in the adephylo package. Pagel’s $\lambda$ and Blomberg’s $K$ were estimated and their significance was tested (999 randomizations) using the phylodig function in the phytools package. The significance of $\lambda$ was tested using a randomization test, while that of $\lambda$ was tested by a likelihood ratio test.

**Results**

Length data were highly correlated between the 12 metrics and across the 40 fish species I considered (Fig. 2). Correlations ranged from $r = 0.9089$ between the common length reported on FishBase and median length recorded in BsM field data to $r = 0.9998$ between the maximum length reported on FishBase and the world record reported in the *Freshwater Fishes of Ontario*. Variation among metrics was greater for larger species (higher CV across metrics; Supporting Information Appendix S1). The mean CV of length among metrics within species (range 0.15–0.52, mean = 0.33, SD = 0.09) was less than half the variation in length among species within metrics (range 0.80–0.93, mean = 0.86, SD = 0.05).

All metrics but the common length reported on FishBase were significant predictors of both range shifts and the impact of predator introduction (at the $\alpha = 0.05$ level; Table 2, Fig. 3). Length was positively related to range shifts with standardized effect sizes ($\beta$s) ranging from 0.55 to 0.98. Length had the strongest effect on predicted range shifts when calculated with field BsM data. Length was negatively related to the impact of predators; standardized effect sizes varied between −0.22 and −0.27, with the exception of common length reported on FishBase ($\beta = -0.15$, but nonsignificant). Aside from this metric, there was little change in the amount of variation in either response explained by length metrics ($R^2$ values in Table 1). Average Ontario length reported in the *Freshwater Fishes of Canada* was the strongest predictor of (explained the most variation in) northern range boundary shifts. Average maximum length reported in the *Freshwater Fishes of Ontario* was the strongest predictor of the impact of *A. rupestris* introductions.

### Table 2. The $R^2$ and P-values from regressions of 12 measures of fish species length as predictors of range boundary shifts and the impact of predator introductions.

| Range boundary shift | Impact of predator introduction |
|----------------------|---------------------------------|
| (df = 11)            | (df = 27)                       |
| $\beta$ | $R^2$ | $P$ | $\beta$ | $R^2$ | $P$ |
|----------------------|----------------------|----------------------|----------------------|----------------------|
| FO.ON.AVG            | 0.76                | 0.62                | 0.001               | 0.25                | 0.26                | 0.005               |
| BSM.AVG              | 0.78                | 0.56                | 0.003               | 0.23                | 0.22                | 0.010               |
| BSM.MED              | 0.86                | 0.59                | 0.002               | 0.23                | 0.21                | 0.012               |
| FC.COM               | 0.67                | 0.59                | 0.002               | 0.27                | 0.30                | 0.002               |
| FB.COM               | 0.55                | 0.30                | 0.051               | 0.15                | 0.09                | 0.115               |
| FO.ON.MAX            | 0.76                | 0.56                | 0.003               | 0.22                | 0.20                | 0.015               |
| FO.WORLD.MAX         | 0.69                | 0.55                | 0.004               | 0.24                | 0.23                | 0.008               |
| FC.MAX               | 0.64                | 0.54                | 0.004               | 0.24                | 0.22                | 0.010               |
| FB.MAX               | 0.67                | 0.52                | 0.005               | 0.24                | 0.23                | 0.008               |
| FT.MAX               | 0.66                | 0.54                | 0.004               | 0.26                | 0.25                | 0.006               |
| BSM.MAX              | 0.98                | 0.61                | 0.002               | 0.25                | 0.22                | 0.010               |
| ROM.MAX              | 0.79                | 0.56                | 0.003               | 0.23                | 0.20                | 0.016               |

Maximum species length recorded during field BsM indicated the weakest phylogenetic signal by all four indices (Table 3). This was the only metric not significantly different from random expectations by all indices. Average maximum length reported in the *Freshwater Fishes of Canada* and the average Ontario length reported in the *Freshwater Fishes of Ontario* were consistently among the metrics with the strongest phylogenetic signal across these indices. All metrics of length, other than the maximum length in BsM, indicated significant phylogenetic signal by all four indices.

**Discussion**

Variability between sources in species trait data may influence the strength of traits as predictors and the ecological inferences we make using trait data. Intraspecific variation in trait values, both across a species range and within populations, could lead to variability in data collected at different scales and by different methods. Here, fish length varied with scale and type of metrics (e.g., average or maximum). Variation among metrics, however, was small in comparison with variation between species. Consistent with this finding, Blanck and Lamouroux (2006) reported smaller intra- than interspecific variation in length for European freshwater fishes. I also found consistent relative lengths across species reflected in high correlations between metrics (Fig. 2). Accordingly, changes in the type of metrics or the scale of the data source did not change the significance of length as an ecological predictor (with the exception of one metric, further discussed below). Average length in Ontario, however, explained the most variation in range boundary shifts across the province. Common length in Canada...
Figure 3. The strongest (left) and weakest (right) relationships between fish species length and (A) the magnitude of northern range boundary shifts or (B) the relative risk on resident populations imposed by predatory Rock Bass introductions. Followed by (C) traitgrams for the measures with relatively strong (left) and weak (right) phylogenetic signal. Traitgrams plot species by their relative size along the y-axis and connect species with an underlying phylogenetic tree. More crossed lines indicate a trait is more randomly distributed indicating less phylogenetic signal.
explained the most variation in the impacts of predator introductions to Ontario lakes. In my analysis, fish length is a reliable predictor of community-level responses to environmental change. But, this may not be universal; trait reliability should be tested for other taxa or with additional species traits (Blanck and Lamouroux 2006; Kazakou et al. 2014).

I found relatively consistent phylogenetic signal in size data; however, this must be interpreted with care. First, while phylogenetic signal is significant, phylogenetic distance is clearly an imperfect predictor of fish size (Fig. 3C). Second, the species pool in the analyzed phylogeny, that of Manitoulin Island, is not a complete sample of the Great Lakes Basin or Ontario regional pool. Finally, while closely related species may share similar functional traits at a regional scale, indicated by phylogenetic signal, this may not hold true in a local community (Gerhold et al. 2015). Thus, the assumption of niche-conservatism, which underlies the use of phylogenetic relatedness as a proxy for trait dispersion in community ecology, may best be tested using trait data sampled from local communities.

Several factors may influence the variability in species trait data between sources. Cordlandwehr et al. (2013) suggest the accuracy of species-level traits retrieved from trait databases depends on three factors: the level of aggregation (scale), the plasticity of chosen traits, and habitat (environment). I add sampling bias as a fourth possible factor influencing the accuracy of trait data. Sampling bias may be location based or influenced by abundance, catchability or historical, cultural or economic interest in the species. For example, Sandel et al. (2015) found frequently measured plant species had higher trait values than rarely measured species. I found the maximum size of small prey fishes with intensive field sampling was often larger than the maximum recorded by other sources. In contrast, the maximum size of larger sportfishes in field sampling was often smaller than the maximum recorded by other sources. I suspect this is the result of historically poor sampling of small fishes with low catchability, the comparatively large samples of sportfish data by anglers and resource management agencies, and the reporting of large sportfish by anglers often during angling competitions. Ultimately, the variability in species trait data through all of these mechanisms is driven by intraspecific variation in traits.

Researchers should carefully evaluate species trait data for potential sampling or measurement biases and errors. Familiarity with the species included in community-level analyses facilitates evaluating trait data. Additionally, outliers in the relationship between trait values taken from two different sources can indicate sampling bias or errors in transcription. For example, the median length of Micropterus salmoides (Largemouth Bass) in BsM field sampling (140 mm) is an outlier when compared to other metrics (Fig. 2, Supporting Information Appendix S1). In this case, the smaller than expected value for M. salmoides is due to a large number of juveniles sampled for this species. Average or median size metrics can be strongly influenced by juvenile life stages, whereas maximum observed metrics can reflect rarely observed sizes, those individuals on the tail end of the size distribution. In my analysis, the only metric by which length was not significantly related to range shifts or the impact of introduced predators was the common length reported in FishBase. The common length of M. dolomieu (Smallmouth Bass) reported by this database is 80 mm, which is clearly erroneous when compared with the values from other source (Fig. 2, Supporting Information Appendix S1). Tomelleri and Eberle (1990) are cited as the source of this value; however, they report a common length of 20 inches, suggesting a transcription error in the FishBase value.

|                  | FO.ON.AVG | BSM.AVG | BSM.MED | FC.COM | FB.COM | FO.ON.MAX | FO.WORLD.MAX | FC.MAX | FB.MAX | FT.MAX | BSM.MAX | ROM.MAX |
|------------------|-----------|---------|---------|--------|--------|-----------|--------------|--------|--------|--------|---------|---------|
| Moran’s I        | 0.28      | 0.35    | 0.32    | 0.32   | 0.20   | 0.25      | 0.28         | 0.25   | 0.29   | 0.30   | 0.13    | 0.27    |
|                  | 0.012     | 0.011   | 0.015   | 0.010  | 0.042  | 0.023     | 0.013        | 0.024  | 0.011  | 0.010  | 0.117   | 0.014   |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
| Abouheif’s Ikm   |           |         |         |        |        |           |              |        |        |        |         |         |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
| Pagel’s λ       |           |         |         |        |        |           |              |        |        |        |         |         |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
| Blomberg’s K     |           |         |         |        |        |           |              |        |        |        |         |         |

|                  | FO.ON.MAX | BSM.AVG | BSM.MED | FC.COM | FB.COM | FO.ON.MAX | FO.WORLD.MAX | FC.MAX | FB.MAX | FT.MAX | BSM.MAX | ROM.MAX |
|------------------|-----------|---------|---------|--------|--------|-----------|--------------|--------|--------|--------|---------|---------|
|                  | 0.25      | 0.27    | 0.32    | 0.32   | 0.20   | 0.25      | 0.28         | 0.25   | 0.29   | 0.30   | 0.13    | 0.27    |
|                  | 0.014     | 0.010   | 0.015   | 0.010  | 0.042  | 0.023     | 0.013        | 0.024  | 0.011  | 0.010  | 0.117   | 0.014   |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
|                  |           |         |         |        |        |           |              |        |        |        |         |         |
Replacing this value with 510 mm (~20 inches) would make the relationship between common length from FishBase and both range shifts and the impacts of introductions significant ($R^2 = 0.42$, $P = 0.016$ and $R^2 = 0.25$, $P = 0.006$, respectively).

As ecologists continue to incorporate species traits into analyses, it is important they consider potential sources of variability in data. Plant databases have begun to include quantitative environmental data for the location of sampled populations as well as population trait mean and variance (Violle et al. 2007; Kattge et al. 2011). This effort should be expanded across taxa and within aquatic ecosystems. Recently developed data-collection tools and citizen-science programs facilitate collecting trait data and location information with common standards (Duputié et al. 2014). In a period of rapid environmental change, species traits will continue to be important for predicting future changes in ecological communities. Accumulating measures of species traits from populations across environmental gradients will be needed to evaluate the relative importance of intraspecific variation and to understand how individual species will respond to environmental changes.

**Data Accessibility**

Data are included in Supporting Information (See Below).

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**Conflict of Interest**

None declared.

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## Supporting Information

Additional Supporting Information may be found online in the supporting information tab for this article:

**Appendix S1.** Excel spreadsheet of twelve metrics of fish length collected from various sources for forty species (organized by length), range shift (degrees latitude) and predator risk (risk ratio) data, which species where included in the analyzed phylogeny and coefficient of variation (CV) across species and measures; see Table 1 for details of metrics and abbreviations.