

# Latitudinal gradients in biotic niche breadth vary across ecosystem types.

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## Abstract

Several properties of food webs—the networks of feeding links between species—are known to vary systematically with the species richness of the underlying community. Under the “latitude-niche breadth hypothesis”, which predicts that species in the tropics will tend to evolve narrower niches, one might expect that these scaling relationships could also be affected by latitude. To test this hypothesis, we analysed the scaling relationships between species richness and average generality, vulnerability, and links per species across a set of 196 empirical food webs. In estuarine, marine, and terrestrial food webs there was no effect of latitude on any scaling relationship, suggesting constant niche breadth in these habitats. In freshwater communities, on the other hand, there were strong effects of latitude on scaling relationships, supporting the latitude-niche breadth hypothesis. These contrasting findings indicate that it may be more important to account for habitat than latitude when exploring gradients in food-web structure.

# 1 Introduction

2 Food webs –networks of feeding links between species– have been used for several decades  
3 to summarise the structure of ecological communities [1–3] and to understand how that  
4 structure relates to environmental variables such as habitat type [4, 5], primary produc-  
5 tivity [6–8], and climate [9, 10]. The latter variables in turn have strong gradients over  
6 latitude, with productivity and temperature both being higher in the tropics while climate  
7 is more variable at high latitudes [11]. These variables affect both the resources available  
8 and species’ metabolisms [12–15], and have been proposed as determinants of the strength  
9 of interspecific interactions [16–18]. By modulating interactions between species, latitudi-  
10 nal gradients may also shape food-web structure. Indeed, these latitudinal environmental  
11 gradients have been put forward as potential drivers for the latitudinal gradient in species  
12 richness, one of the most general and robust patterns in ecology [16, 19, 20].

13 One proposed link between species richness and latitude is the “latitude-niche breadth  
14 hypothesis” [21]. This hypothesis predicts that decreased seasonality in the tropics should  
15 lead to more stable populations, which in turn should evolve smaller niches [21]. These  
16 narrow niches should therefore allow more species to coexist in the tropics than at higher  
17 latitudes. Alternatively, the higher productivity of the tropics [22] may result in a broader  
18 niche space [23] which could also sustain greater biodiversity even if niche sizes are globally  
19 similar. Although the assumptions of the latitude-niche breadth hypothesis are only  
20 equivocally supported [21], it remains a compelling potential mechanism for the latitudinal  
21 gradient in species richness [24–26].

22 If the latitude-niche breadth hypothesis is correct, there should also be direct relation-  
23 ships between latitude and the degree of specialisation (i.e., the breadth of the Eltonian  
24 niche; [27, 28]) of species within food webs. Specifically, narrower niches in the tropics

would equate to greater specialisation (narrower niches) while constant niche sizes but greater productivity would translate to constant specialisation and niche width across latitude (Fig. 1). Attempts to unravel these effects, however, are complicated by known relationships between species richness and many other network properties [29]. For example, narrower niches imply fewer links per species (i.e., greater specialisation) in the tropics ([30, 31]; but see [18]). However, average numbers of links per species tend to increase in larger food webs [29, 32]. This means that a latitudinal effect on specialisation may be obscured by a latitudinal gradient in species richness.

If this is the case, it may still be possible to uncover effects of latitude on specialisation by examining the shape of the scaling relationship between specialisation and species richness over changing latitude. By testing whether latitude affects the scaling of each property with species richness, we test for the effects of latitude on specialisation predicted by the “latitude-niche breadth hypothesis” (Fig. 1). If the scaling of specialisation with species richness is weaker in the tropics (i.e., if species gain fewer links, predators, or prey as the size of the network increases), this will indicate narrower niches at the tropics. If, however, the scaling of specialisation with species richness does not vary over latitude, this will indicate that niches are similarly-sized worldwide but that there is a broader niche space in the tropics. Additionally, as food webs describing different ecosystem types may differ in their topology [5, 33], we also explored the differences in scaling relationships across ecosystem types. Here, we use three measures of specialisation; mean links per species, mean generality (number of prey), and mean vulnerability (number of predators).

## Methods

### Data Set

We compiled a list of 196 empirical food webs from multiple sources (see *Appendix S1* for web origins and selection criteria). We recorded study site latitude from the original source where possible or, where study sites were described but exact coordinates were not given, obtained estimated coordinates using Google Earth [34]. If a range of latitudes (e.g.  $42 - 49^{\circ}N$ ) was provided, we used the midpoint of this range. We grouped food webs by ecosystem type (stream,  $N=71$ ; lake,  $N=47$ ; marine,  $N=28$ ; estuarine,  $N=18$ ; and terrestrial,  $N=31$ ) according to their designation in previous aggregations of food webs (i.e., [35–37]).

As the food webs in this dataset are derived from a variety of sources and were compiled over many decades, it is likely that they vary in their resolution and in the amount of sampling effort invested in their assembly. Many analyses of food-web structure attempt to reduce this variation by using food webs comprised of “trophic species” –aggregations of species with identical sets of predators and prey– rather than species *per se* [8, 33, 37, 38]. As our study is concerned directly with the number of species at a particular latitude, however, we did not wish to ignore species with identical sets of interactions. We therefore analysed both original (i.e., without aggregating any species) and trophic-species (i.e., after aggregating species with identical predators and prey) versions of the dataset; in each case using the number of species and feeding links in each web to calculate the mean link density (number of links per species), mean generality (number of prey per species), and mean vulnerability (number of predators per species) of the web. The version of the dataset used did not qualitatively change the results, suggesting that the scaling relationships between species richness, other food-web properties, and latitude are very

similar whether or not species with identical sets of predators and prey are included. For simplicity, here we present only the results for the original (un-aggregated, original species) webs.

## Gradients over Latitude

To put our dataset in the context of other research on latitudinal gradients in species richness, we first examined simple linear relationships between latitude and each of species richness, links per species, generality, vulnerability, and proportions of basal resources, intermediate consumers, and top predators. We fit models of the form

$$S_i = \alpha_0 + \alpha_1 L_i + \alpha_2 E_i + \alpha_3 L_i E_i + \epsilon_i, \quad (1)$$

where  $S_i$  is the species richness of web  $i$ ,  $L_i$  its absolute latitude (degrees north or south regardless of direction),  $E_i$  is a categorical variable indicating the ecosystem type of network  $i$  (comprising terms for stream, marine, lake, and terrestrial networks with estuarine networks corresponding to  $E_i = 0$ ) and  $\epsilon_i$  is a residual error term. We next calculated the AIC of the maximal model as well as the AIC's of a suite of candidate simplified models identified using the R [39] function dredge from package MuMIn [40]. Simplified models were obtained by systematically removing all possible combinations of terms from the full model. The best-fitting model was then determined to be the model with the fewest terms where  $\Delta\text{AIC} < 2$ , as this model is the least likely to suffer from over-fitting.

## Scaling Relationships with Species Richness

The scaling relationship between link density ( $Z$ ) and species richness ( $S$ ) has been shown to be a power law [29] of the form

$$Z_i \sim \alpha S_i^\beta, \quad (2)$$

90 which is often re-expressed in logarithmic form

$$\log Z_i \sim \log \alpha + \beta \log S_i. \quad (3)$$

91 As the two forms imply a statistical fit of the data to different error distributions, neither  
 92 of which has strong *a priori* justification in our dataset, we followed the recommendations  
 93 in [41] to compare the two model formulations explicitly (see *Appendix S2* for details). The  
 94 logarithmic form (equation 3) provided the better fit to the data, as did the logarithmic  
 95 forms of similar models for the scaling of generality and vulnerability. We therefore used  
 96 and present logarithmic models throughout the rest of the analyses.

## 97 **Effect of Latitude on Scaling**

98 After determining the appropriate form of the scaling relationship, we then assessed the  
 99 impact of latitude on the scaling relationships between species richness and link density,  
 100 generality, vulnerability. In the context of the scaling relationships above, note that this  
 101 implies that we wished to determine the effect of latitude on the scaling exponent  $\beta$ . We  
 102 included a categorical variable for ecosystem type (stream, lake, terrestrial, marine, or  
 103 estuary), as well as interactions between food web type and latitude.

104 We therefore began by considering models of the form

$$Z_i = \alpha S_i^{\beta_0 + \beta_1 L_i + \beta_2 E_i + \beta_3 L E_i} + \epsilon_i, \quad (4)$$

105 where  $S_i$  is the species richness of web  $i$ ,  $L_i$  its absolute latitude (degrees north or south  
 106 regardless of direction),  $E_i$  is a categorical variable indicating the ecosystem type of net-  
 107 work  $i$  (comprising terms for stream, marine, lake, and terrestrial networks with estuarine

networks corresponding to  $E_i = 0$ ) and  $\epsilon_i$  is a residual error term. The logarithmic formulation of this model is

$$\log Z_i = \log \alpha + \beta_0 \log S_i + \beta_1 L \log S_i + \beta_2 E \log S_i + \beta_3 LE \log S_i + \epsilon_i. \quad (5)$$

We then simplified versions of model 5 for link density, generality, and vulnerability following the procedure described above. As a supplemental check to ensure that variation in sampling effort across food webs was not responsible for the trends we observed, we then repeated our analyses using jackknifed data sets in which we 1) sequentially removed each food web in the dataset and 2) sequentially removed sets of food webs that shared a common author. The first jackknife essentially controls for the influence of any single outlier, while the second controls for the influence of particular research groups, some of which contributed large numbers of food webs (up to 27) to the dataset. Parameter estimates for the simplified models varied very little across either jackknife test (see *Appendix S3* for details), indicating that the trends we observed were not due to either strong outliers or to substantial differences in sampling effort across research groups.

## Results

Link density (mean number of feeding links per species), generality (mean number of prey per species), and vulnerability (mean number of predators per species) were strongly and positively correlated ( $R^2=0.891$  for link density and generality,  $R^2>0.999$  for link density and vulnerability, and  $R^2=0.890$  for generality and vulnerability). Contrary to the expected latitudinal gradient, the best-fit version of equation (1) did not include a significant effect of latitude for any ecosystem type. Further, there were no significant relationships between link density, generality, or vulnerability with latitude for any ecosystem type.

Each measure of specialisation increased with increasing species richness ( $\beta_0=0.637$ ,  $p<0.001$ ;  $\beta_0=0.553$ ,  $p<0.001$ ; and  $\beta_0=0.637$ ,  $p<0.001$ , respectively; Fig. 2). For estuarine, marine, and terrestrial food webs the strength of this scaling did not vary with latitude ( $\beta_{Latitude}=-0.001$ ,  $p=0.365$  for link density;  $\beta_{Latitude}=-0.001$ ,  $p=0.535$  for generality; and  $\beta_{Latitude}=-0.001$ ,  $p=0.366$  for vulnerability; Fig. 3). In lake food webs, however, the scaling of each property was stronger towards the poles ( $\beta_{Latitude:Lake}=0.004$ ,  $p=0.019$ ;  $\beta_{Latitude:Lake}=0.005$ ,  $p=0.004$ ; and  $\beta_{Latitude:Lake}=0.004$ ,  $p=0.018$ , respectively). In stream food webs, generality increased more rapidly towards the poles ( $\beta_{Latitude:Stream}=0.007$ ,  $p=0.001$ ) while link density and vulnerability did not vary with latitude (i.e., the interaction term  $\beta_{Latitude:Stream}$  was not retained in the best-fit models).

## Discussion

The tendency of food-web structure to exhibit scaling relationships with species richness has been well-established [29, 33]. As species richness in particular is also known to vary systematically over latitude [16, 19, 20, 42], intuitively one might suspect that any relationship between food-web properties such as generality might be due to the latitudinal gradient in species richness. In this dataset, however, we found no evidence to support latitudinal gradients in species richness, links per species, generality, or vulnerability.

The lack of a latitudinal gradient in species richness in this dataset contrasts strongly with other studies [16, 19, 20, 42]. As numbers of species and links included in a food web vary strongly with sampling effort as well as with the underlying diversity of the study area, it is possible that the lack of latitudinal trends here is a result of researchers tending to expend similar amounts of sampling effort across studies. This could result in food webs describing species-rich tropical communities omitting more species and links than studies of species-poor arctic communities if research groups spend similar person-hours



assembling webs and can observe similar numbers of species and links per person-hour. In addition, it is worth noting that gradients in species richness are generally measured for a single taxonomic group at a time [16, 19, 20, 42]. It is possible that these taxa are not well-represented in our food webs and that the dominant taxa that are represented do not have an underlying latitudinal gradient in richness. In either case, the lack of a strong association between species richness and latitude in any ecosystem type means that any effect of latitude on other scaling relationships is not being driven by an underlying latitudinal gradient in species richness. This is fortunate since the lack of confounding effects of latitude allows us to more clearly assess effects of latitude on scaling with species richness.

Scaling of links per species, generality, and vulnerability with species richness varied strongly across ecosystem types. In estuarine, marine, and terrestrial food webs scaling of each property varied little with latitude. This is consistent with the idea that species' niche breadths do not vary systematically with temperature and productivity but that the niche space might be larger in species-rich communities [23]. Rather than niche space depending on temperature and productivity, it may be that species diversity itself affects the biotic niche space available to species (although climate may affect speciation rates and therefore the diversity in a region [43]). For example, as the plant diversity of a community increases both the variety of food available to herbivores and the structural variety of the habitat will also increase.

Unlike other ecosystem types, the scaling of generality in lake and stream food webs was stronger (i.e., generality increased more steeply with increasing species richness) in higher-latitude food webs. In lake food webs, this trend was echoed in the scaling relationships between species richness and vulnerability and links per species. This means that species in tropical freshwater communities gain fewer additional feeding links per

178 additional species in the web and that species in tropical lakes also gain fewer preda-  
179 tors, and fewer links in general, per additional species than species in high-latitude lakes.  
180 These trends are consistent with the hypothesis that greater stability in the tropics leads  
181 to narrower niches [21] and a higher proportion of specialists.

182 That freshwater food webs supported the hypothesis of narrower niches in the tropics  
183 –while other ecosystem types did not– is noteworthy given that these ecosystems (es-  
184 pecially streams) are known for being highly variable and that seasonal variability is  
185 one of the proposed drivers of the latitude-niche breadth hypothesis [21]. Both streams  
186 and lakes can experience severe changes in water temperature and volume (e.g., floods,  
187 drying, freezing) that remove food or other resources (notably oxygen during freezing  
188 events) [44, 45]. These events are often linked to seasonal events such as snowmelts or  
189 summer drought [44]). Further, both temperate streams and lakes tend to experience sea-  
190 sonal strong pulses of allochthonous inputs (e.g., fallen leaves, terrestrial invertebrates [46–  
191 48]. These trends combined mean that, relative to estuarine and marine communities,  
192 freshwater food webs may experience high turnover in both community composition and  
193 productivity [49–51]. Notable exceptions from the above trends are New Zealand stream  
194 communities (representing 31 of the 71 stream food webs in our dataset), which experi-  
195 ence unpredictable flooding and drying throughout the year and do not receive seasonally  
196 pulsed subsidies [44, 52]. However, as this subset of webs is very tightly grouped in lati-  
197 tude ( $44.64 - 46.41^{\circ}\text{S}$ , within an overall range of  $23 - 69.02^{\circ}$  for stream communities), it is  
198 unlikely that they have greatly influenced our results (see also *Appendix S3*). Moreover,  
199 just as in highly-variable communities where said variation is more seasonal, New Zealand  
200 communities are dominated by ecological generalists [44, 52] implying that they appear  
201 to fit the general pattern of streams worldwide.

202 Importantly, while terrestrial communities are also strongly seasonal at high latitudes

and can receive significant allochthonous inputs [46], terrestrial consumers tend to be morphologically specialised for feeding on particular prey [53]. This means that primarily gape-limited aquatic consumers tend to be more generalist across all types of aquatic habitats than terrestrial consumers [5, 53]. The key to this explanation of the differences between freshwater and marine and estuarine ecosystems is whether the former experience more severe seasonal variation. Although we are not aware of any study explicitly comparing seasonal variation in freshwater and saltwater or brackish food webs in a similar location, we believe that freshwater ecosystems are indeed likely to experience more severe changes because of their small size. While oceans and estuaries certainly vary in terms of water temperature and nutrients over the course of a year [51], these changes are likely to be slower and milder than in freshwaters because marine and estuarine communities are buffered by being open to the ocean rather than isolated in the midst of a terrestrial matrix. Net primary productivity in particular is much more variable over the course of a year in non-marine communities [11], suggesting that niche breadths may also be more variable over the course of the year.

## Conclusion

Overall, our results were inconsistent with the latitude-niche breadth hypothesis in estuarine, marine, and terrestrial communities but consistent with the hypothesis of greater specialisation in the tropics in stream and lake food webs. This suggests that different mechanisms may structure food webs in different habitat types and that freshwater food webs in particular may be strongly affected by seasonal variation. In addition, different relationships between latitude and niche breadth in different habitat types goes some way towards explaining the equivocal support for the opposing hypotheses of narrower niches in the tropics [21] and broader niche space in the tropics [23]. Our study indicates that

both have merit but would appear to apply to different systems.

## Data Accessibility

Food webs used in this study were retrieved from the University of Canberra’s GlobalWeb database ([35]; [www.globalwebdb.com](http://www.globalwebdb.com)) and from two papers ([36]; <http://dx.doi.org/10.1111/j.1461-0248.2010.01568.x> and [37]; <http://dx.doi.org/10.1371/journal.pbio.1001579>). Original sources for the food webs are given in *Appendix S1*.

## Competing interests

We have no competing interests.

## Authors’ contributions

ARC, DBS, and TNR designed the study, ARC collected published data, performed the analyses, and wrote the first draft. DBS and TNR substantially revised the article; all authors approved the final version.

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## Figure Captions

**Figure 1:** In A) we show the known scaling relationship between link density (links per species) and species richness. This scaling relationship is a power law and therefore linear in a log-log plot. In B) we show two versions of the latitudinal-niche breadth hypothesis that have been proposed to explain this gradient. Hypothesis 1 posits that greater environmental stability in the tropics will allow species to evolve narrower niches (indicated by parabolas) than those at the poles. Hypothesis 2 suggests that species will have constant niche sizes over latitude but that greater primary productivity in the tropics creates a larger niche space such that each species still occupies a smaller proportion of the total niche space. These two hypotheses have different implications for the scaling of food-web properties such as the number of feeding links per species with species richness. C) If hypothesis 1 is true, then the exponent of the scaling relationship between link density and species richness should be larger towards the poles, where each additional species in the food-web will have a larger niche (i.e., more feeding links). If hypothesis 2 is true, then the exponent of this distribution should not vary significantly over latitude.

**Figure 2:** Scaling relationships for re-scaled link density, generality, and vulnerability relative to the species richness of a food web. Link density, generality, and vulnerability were each re-scaled to remove the effects of latitude and ecosystem type. As these relationships take the form of power laws, we did this by dividing the food-web property (e.g. link density) by species richness raised to an exponent including the effects of latitude and, where applicable, ecosystem type and the interaction between ecosystem type and latitude. Note that in all cases estuarine food webs were treated as the baseline ecosystem type, but that at most two ecosystem types had interactions between ecosystem type and latitude retained in the best-fit model (see *Results* for specifics). For each relationship, we show the re-scaled values (white circles) as well as the overall scaling relationship using estuarine ecosystems as a baseline (black line, N=196 food webs). For a figure with the uncorrected values, see Fig. *S7*, *Appendix S4*.

**Figure 3:** Changes to the scaling of link density, generality, and vulnerability with species richness across ecosystem types and over latitude. We show the estimated scaling exponent for species richness (black line) with its 95% confidence interval (in grey), based on N=196 empirical food webs. Latitude is given in degrees from the equator regardless of direction.