


LETTER

Globally consistent reef size spectra integrating fishes and invertebrates

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Abstract

The frequency distribution of individual body sizes in animal communities (i.e. the size spectrum) provides powerful insights for understanding the energy flux through food webs. However, studies of size spectra in rocky and coral reef communities typically focus only on fishes or invertebrates due to taxonomic and data constraints, and consequently ignore energy pathways involving the full range of macroscopic consumer taxa. We analyse size spectra with co-located fish and mobile macroinvertebrate data from 3369 reef sites worldwide, specifically focusing on how the addition of invertebrate data alters patterns. The inclusion of invertebrates steepens the size spectrum, more so in temperate regions, resulting in a consistent size spectrum slope across latitudes, and bringing slopes closer to theoretical expectations based on energy flow through the system. These results highlight the importance of understanding contributions of both invertebrates and fishes to reef food webs worldwide.

Keywords

Benthic, biomass equivalence rule, coastal, community, fish, macroecology, size distribution, size spectrum, trophic pathways.

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INTRODUCTION

Body size is arguably the most important single factor determining an individual's vital rates and how it interacts with its environment (Brown *et al.*, 2004). Body size distributions therefore provide rich insights into size-dependent relationships between animals and underlying energy flow of communities. One such distribution links individual body size and abundance in a community (the community size spectrum). This relationship has been extensively studied in both marine and terrestrial realms (e.g. Reuman *et al.*, 2008), following early conjectures of a 'biomass equivalence rule': that biomass is approximately equal across logarithmic size bins spanning sizes of the smallest to the largest creatures (Ghilarov, 1944; Sheldon *et al.*, 1972). This results in a negative power-law relationship between abundance concentration (N) and body size (M) (Andersen and Beyer, 2006), $N \propto M^\lambda$, where $\lambda \approx -2$. Because of the important information concerning system-wide energy movements (Brown and Gillooly, 2003; Trebilco *et al.*, 2013), methods used to estimate the power law exponent have been extensively evaluated in the literature (White *et al.*, 2008; Edwards *et al.*, 2017).

Although remarkable consistencies in empirical size spectra have been observed (Sprules *et al.*, 2016), substantial deviations can also occur. These deviations provide important information about ecosystem structure and perturbations. For example the selective removal of larger individuals through fishing has been shown to steepen the negative slope of the size spectrum in both pelagic (Pope and Knights, 1982; Daan *et al.*, 2005; Blanchard *et al.*, 2005) and reef ecosystems (Dulvy *et al.*, 2004; Graham *et al.*, 2005; Wilson *et al.*, 2010; Robinson *et al.*, 2017). By contrast, seasonal competition for resources (Edgar, 1994) and energy subsidies from outside the reef ecosystem (Trebilco *et al.*, 2013, 2016; Morais and Bellwood, 2019) can potentially result in shallower size spectra, while habitat

complexity can cause deviations of the size spectra from the expected power law (Rogers *et al.*, 2014). For a community of individuals feeding on a common resource, that is at a single trophic level, such as herbivorous fishes (Robinson *et al.*, 2016), abundance may also scale less steeply with body size, following the allometric scaling of body size with metabolic rate and energetic equivalence (Kleiber, 1932; Damuth, 1981; Nee *et al.*, 1991). However, most aquatic communities are comprised of a trophic chain or web, whereby individuals feed upon one another as well as the basal resource. Consequently, due to inefficiencies in the transfer of energy between trophic levels (Lindeman, 1942), fewer individuals can be sustained when feeding at higher trophic levels. Given the strong relationship between an individual's size and its trophic position (Jennings *et al.*, 2001), this is consistent with fewer large-bodied individuals in a community arising from individuals feeding in a size-based way (i.e. a food chain or web) (Brown and Gillooly, 2003; Jennings and Mackinson, 2003; Trebilco *et al.*, 2013; Andersen, 2019). Although the general pattern of declining abundance with body size holds in many places, particularly at very large spatial scales, there has been no global test of the 'biomass equivalence rule' at the community scale for reefs or any other large system (Polishchuk and Blanchard, 2019).

Global data sets available to test the 'biomass equivalence rule' for marine systems have been previously lacking. The Reef Life Survey (RLS) program has quantified the abundance and size distribution of all conspicuous species on reef habitats globally (Edgar and Stuart-Smith, 2014) and provides the best available means for exploring biomass equivalence at this scale. It is the largest single database, terrestrial or marine, in terms of its taxonomic, spatial and temporal coverage with a basis of standardised quantitative methods. The high resolution yet global coverage of the data enables us to investigate size spectra at varying spatial scales.

Another challenge relates to the major missing component of reef community size spectra: benthic invertebrates. Whilst most previous empirical work on reef size spectra has focused solely on fish communities, large mobile benthic invertebrates can play fundamental roles in reef ecosystems, even to the point of dominating the animal biomass present. For example in some temperate reefs, we observed communities in which over 90% of individuals >1cm body size, were invertebrates (see also Edgar *et al.*, 2017). Furthermore, considerable overlap exists in resource use between fishes and invertebrates, with overlap in the diets of many fishes and invertebrates, and many fish predators relying heavily on invertebrate prey (i.e. fishes and invertebrates do not necessarily occupy separate energy pathways) (Barneche *et al.*, 2014). As such, to better understand the size structure of whole reef communities and food webs that are not artificially constrained by taxonomic group, data on both fishes and invertebrates are needed. Several previous studies have recognised the potential importance of invertebrates in reef size spectra (e.g. Donovan *et al.*, 2018), but body size data were lacking. Here, we use invertebrate body size data to test the ‘biomass equivalence rule’ for size spectra of reef communities, comparing fish-only data and fish and invertebrate data for the same sites globally.

We hypothesise that: (1) The inclusion of invertebrates will change the slope (i.e. exponent) of the community size spectrum (Figure 1). If invertebrates are relatively smaller bodied than their fish counterparts in a community (e.g. Figure 1a), we would expect their inclusion in the size spectrum to have a steepening effect (Figure 1b). Likewise, if invertebrates are relatively larger bodied than the fishes in the community (e.g. Figure 1c), we would expect a shallowing effect when they are included (Figure 1d). This also might correspond to a situation where herbivorous or detritivorous invertebrates occupy a single trophic level, which would result in shallower slopes (Dinmore and Jennings, 2004; Maxwell and Jennings, 2006). We further hypothesise that: (2) This invertebrate inclusion effect will be greater in temperate communities compared to tropical communities due to a relatively greater proportion of invertebrates in temperate reefs (Edgar *et al.*, 2017). (3) The broad geographic span and fine transect-level grain allow us to consider multiple spatial scales, and thereby test our third hypothesis; spatial scale of sampling contributes to variation around slope estimates. A λ of -2 is expected in the absence of human impacts, such as fishing. Because few reefs worldwide are beyond the reach of fishers, we expect to find a steeper (more negative) slope overall. This study provides improved understanding on the variability of reef size spectrum slopes globally, which is crucial for the development of size spectra as indicators for reef ecosystem health (e.g. Nash and Graham, 2016; Trebilco *et al.*, 2016; Zgliczynski and Sandin, 2017; Morais *et al.*, 2020a).

Survey data

Applying the RLS protocol (available at <https://www.reeflife-survey.com/>), trained divers swim along a 50 m transect and identify to species level the fishes and invertebrates they encounter (Edgar and Stuart-Smith, 2014). A single survey ($n = 11936$ surveys) consists of two separate methods undertaken on the same transect line. Method 1 involves recording any fish species ($n = 2608$ species) within 5m wide blocks either

side of the line, whereas method 2 involves searching along the bottom, underneath kelp and in cracks in 1m wide blocks either side of the line, recording invertebrates ($n = 1184$ species) and cryptic fishes ($n = 951$ species). The abundance of each species within the defined block area is counted directly or estimated when necessary for highly abundant species. Size is estimated for all fishes, and by experienced biologists for invertebrates at some sites. Animals are estimated to belong to one of 13 size categories: 2.5, 5, 7.5, 10, 12.5, 15, 20, 25, 30, 35, 40, 50, and 62.5cm. Lengths greater than 62.5 cm are estimated to the nearest 12.5cm. For a full description of the survey methods, see RLS (2020). The abundance from method 2 records was standardised to the equivalent area covered by method 1 by multiplying abundance by five, standardising all records as densities per 500 m². A site ($n = 3369$ sites) usually contained multiple surveys undertaken along at least two depths on the same day. Sites are nested in ‘locations’, which are nested within ecoregions ($n = 91$ ecoregions), as defined by the Marine Ecoregions of the World (Spalding *et al.*, 2007).

Estimation of invertebrate body length distributions

All invertebrates encountered on surveys were identified to species level (or the highest taxonomic resolution possible) and counted within 1m wide blocks on either side of each 50 m transect line surveyed for fishes. At a small subset of surveys, body length of the invertebrates was estimated or measured. Species body length distributions with sufficient observations ($n > 10$ per species, spanning a sufficient range of body length bins for distribution fitting) were therefore available for only 167 invertebrate species ($\approx 14\%$ of total invertebrate species in the data) from seven taxonomic classes. For these species, individual body lengths were best described by a lognormal distribution, consistent with the body length distributions of the fish species and previous body length distribution literature (e.g. Blackburn and Gaston, 1994). For each species, we fitted a lognormal distribution to the body lengths using the ‘fitdistrplus’ package (Delignette-Muller and Dutang, 2015) in R (R Core Team, 2020). We then fitted two linear regression models estimating the two parameters of the lognormal distribution (mean and variance) using the asymptotic length of the species and its taxonomic class as predictor variables (Equations S1.2, S1.3). For the remaining species with only asymptotic length available, we were then able to reconstruct the lognormal body length distribution by estimating the two lognormal distribution parameters using these two regression models. Asymptotic sizes for all invertebrate species were obtained from SealifeBase (Palomares and Pauly, 2019).

From body length to body mass

Conversion to individual body mass distributions was achieved using published length-weight allometric relationships derived from SealifeBase (Palomares and Pauly, 2019) and FishBase (Froese and Pauly, 2010) and observed (where available) or estimated individual body length. For each species, we calculated the asymptotic mass (M_∞) given asymptotic body length (L_∞) and the species’ length-weight relationship. Where species-specific individual length-weight

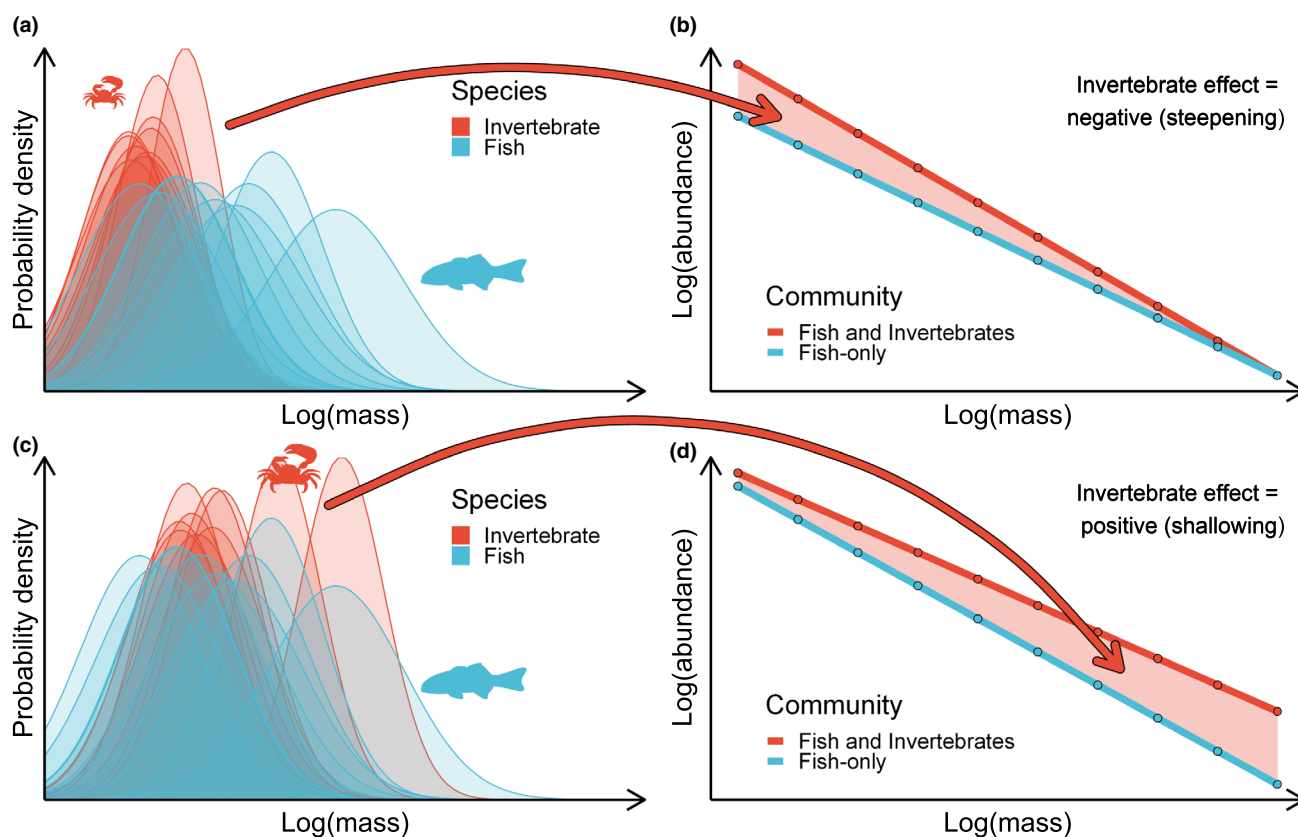


Figure 1 Hypothesised effect of including invertebrates in the size spectrum: (1) A steepening effect (a and b), and (2) a shallowing effect (c and d). The steepness of the size spectrum arises from the relative abundances of larger and smaller-bodied individuals. If invertebrates have a steeper size spectrum slope (i.e. relatively fewer large-bodied individuals) compared to their co-located fish (a), we would expect the slope of the size spectrum of the combined community (fish and invertebrates) to be steeper than the slope of the fish only (b). A shallowing effect (d) would be expected if invertebrates have a relatively greater number of large-bodied individuals compared to the fish-only community (c).

information was unavailable, body mass was estimated from one of two linear regression models: a class-level and an overall length–weight regression model (Supplementary material S2).

To assess the effect of including invertebrates into the size spectrum on the estimation of the slope, all further analyses were carried out firstly with only fish species included, and secondly with invertebrates also included. Differences in the size spectrum slopes between these two analyses are referred to as the ‘invertebrate inclusion effect’ ($\Delta\lambda$).

Fitting the normalised abundance size spectrum

Relationships between N and M are generally estimated from a linear regression of binned size data on a log–log scale (Newman 2005). Size spectrum analyses often ‘normalise’ the y-axis by dividing the abundance within each mass bin by the actual width of the x-axis bin to account for varying bin widths. This normalisation procedure has the effect of reducing the size spectrum slope by 1 and results in the slope being comparable with the power law exponent λ . Here we use the slope of the normalised abundance size spectrum to estimate the exponent λ . We chose a linear regression method over a maximum likelihood estimation of the exponent (see Edwards

et al., 2017), due to the simplicity of incorporating the spatially hierarchical nature of the data (sites nested within ecoregions).

For each survey, individuals were binned into \log_2 mass bins, and the abundance within each bin is calculated as the number of individuals in each bin. Ackerman and Bellwood (2000) found that the abundances of 75% of fish smaller than 5cm were underestimated in reef visual census data. To avoid biases associated with under-sampling of small individuals, we applied a lower bound cut-off of 32g body mass, which represented the modal \log_2 mass bin (Supplementary material S3, see also Ackerman *et al.*, 2004). Abundances were divided by 500 to obtain abundance per m^2 .

We normalised the abundance by dividing by the width of the logarithmic mass bin (Supplementary material S4). We then fitted linear mixed-effects models of \log_2 abundance (N) as a function of the \log_2 mass bin mid (M) and with ecoregion (e) and site (s) as random effects, both having a random slope and intercept, and with site nested within ecoregion (Equation 1).

$$\log_2(N) = \beta_0 + u_{0,e} + u_{0,s|e} + (\beta_1 + u_{1,e} + u_{1,s|e}) \cdot \log_2(M) + \varepsilon \quad (1)$$

where, $u_{0,e}$, $u_{0,s|e}$, $u_{1,e}$ and $u_{1,s|e}$ are normally distributed random effects, and where β_1 represents the overall (global-level)

slope, $u_{1,e}$ is the ecoregion-level variation and $u_{1,s|e}$ the site level variation (given the ecoregion variation) in the slope estimates of the model (Supplementary material S4). Linear mixed models were fitted using the lme4 package (Bates *et al.*, 2015) in R (R Core Team, 2020). Confidence intervals around the overall slope estimate were estimated using the Wald method in the ‘confint’ function of the lme4 package (Bates *et al.*, 2015).

RESULTS

For fish-only communities, we estimated the overall mean site-level slope of the normalised abundance size spectrum (λ) as $-1.88 (\pm 0.06, 95\% \text{ CI})$. The inclusion of invertebrates steepened (i.e. decreased) λ from -1.88 to $-2.04 (\pm 0.06, 95\% \text{ CI})$ (Figure 2, One sample *t*-test: mean $\Delta\lambda = -0.07$, d.f. = 3377, $P < 0.001$).

Absolute latitude explained 13% of the variation in the invertebrate inclusion effect ($\Delta\lambda$), with a greater steepening at higher latitudes (linear regression model: $\Delta\lambda \sim \text{abs}(\text{latitude})$; $R^2 = 13\%$, $P < 0.001$) (Figure 3b and c). Slopes for fish-only communities were shallower at high latitudes, whereas slopes for the combined fish and invertebrate data were remarkably consistent across latitudes (Figure 3a) (see also S5). This greater steepening by invertebrate inclusion in higher latitude regions, was also observed in sites with the greatest protection from fishing pressure (see Supplementary material S6).

Variation in the slope estimates was explained at both the ecoregion and site (given the ecoregion) scales (Figure 4). More of the variation in the slope was evident across ecoregions (Combined community: $\sigma_e = 0.2$, 14% total variation), than among sites within ecoregions (Combined community: $\sigma_{s|e} = 0.17$, 9% of total variation). The total variation explained, across all sites and ecoregions, is the sum of these two variation components, and hence shows that variation declines with increasing spatial scale overall.

DISCUSSION

This study provides the first global test of the generality of the ‘biomass equivalence rule’ for reef communities, analysing size spectra of 3369 reef communities worldwide. Our analyses resulted in three key findings: (1) The inclusion of invertebrates, as opposed to a purely fish-centric approach generally used previously, brought the global estimate of size spectrum slopes closer to the theoretical exponent of -2 , the value expected under the biomass equivalence rule; (2) The effect of including invertebrates was most marked for temperate reefs, where invertebrates contribute a substantial fraction of reef animal biomass; and (3) The contributions to variance in slope estimates were comparable at both the ecoregion (14%) and site scales (9%). Many studies of size spectra aggregate observations to larger spatial scales, whereas our work shows that accounting for hierarchical sampling at the local community scale is important for informing the overall processes driving estimates of size spectra as well as testing the generality of theoretical expectations.

Size spectrum theory, that encompasses detailed mechanistic models describing size-based feeding and physiological constraints (Blanchard *et al.*, 2017; Andersen, 2019) to simple scaling theory that summarises these processes via transfer efficiency and predator–prey mass ratios (Brown and Gillooly, 2003; Jennings and Mackinson, 2003) both predict normalised abundance size spectrum slopes of approximately -2 . However, many processes can affect both of these assumptions and could contribute to the variation around this theoretical value, even in the absence of fishing (Trebilco *et al.*, 2016; Eddy *et al.*, 2020). The empirical consistency of the size spectrum slope across many different aquatic ecosystems (Sprules *et al.*, 2016), and sensitivity to the effects of impacts such as fishing (Shin *et al.*, 2005; Petchey and Belgrano, 2010), has led to its proposed use as an ecological indicator of ecosystem health for reefs (Nash and Graham, 2016). However, its uptake for

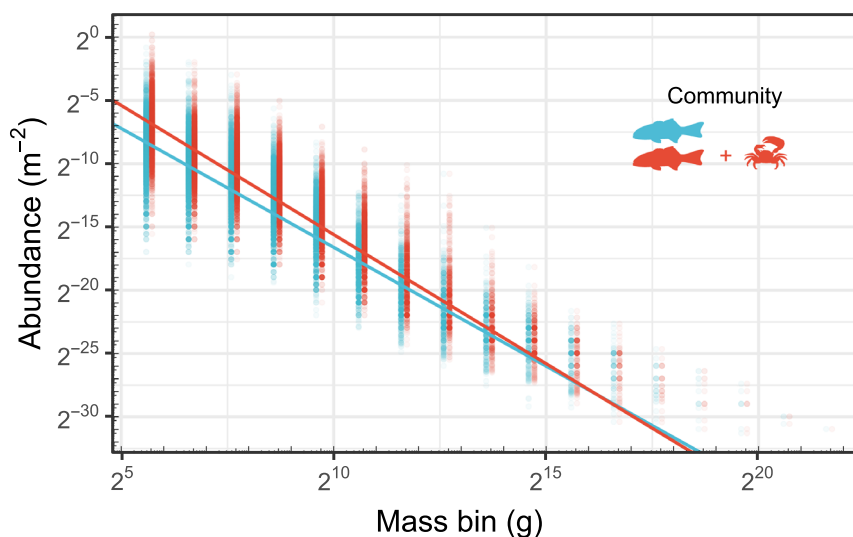


Figure 2 Invertebrates steepen the normalised abundance size spectrum. Separate normalised abundance size spectra are shown for the fish-only and combined (fish and invertebrate) communities, with solid lines representing fits from linear mixed-effects models for the global data (‘Site’ nested within ‘Ecoregion’ as random effects). Fish-only slope = -1.88 ± 0.06 , combined slope = -2.04 ± 0.06 . Points have been offset on the x-axis for clarity.

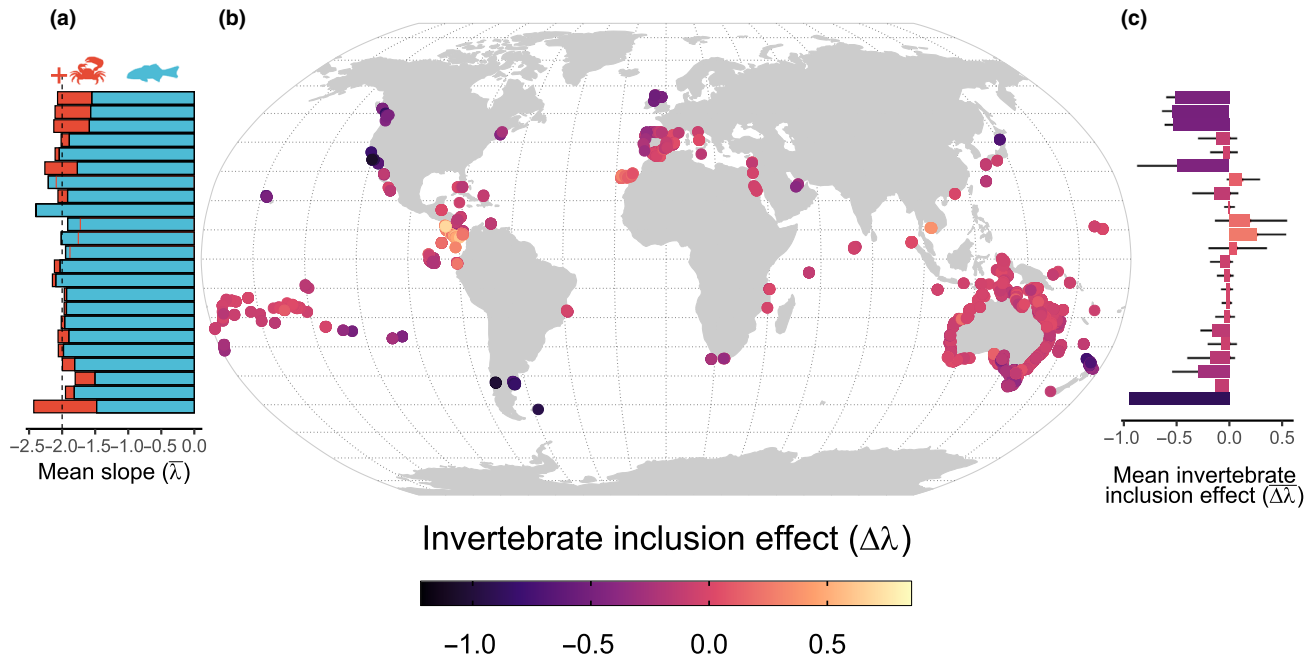


Figure 3 The inclusion of invertebrates results in a consistent community size spectrum slope of ~ -2 . (a) The size spectrum slope for fish-only communities (blue) and when including invertebrates (orange) - orange vertical lines have been used to indicate the top of the orange bar when obscured. (b) A map of the invertebrate inclusion effect ($\Delta\lambda$) across the globe. (c) The latitudinal variation in the 'invertebrate inclusion effect' ($\Delta\lambda$). The steepening effect when including invertebrates is greatest at high latitudes. Each bar in (a) and (c) represents the mean over 5° of latitude. Error bars in C represent the 95% confidence intervals, and missing error bars represent insufficient data.

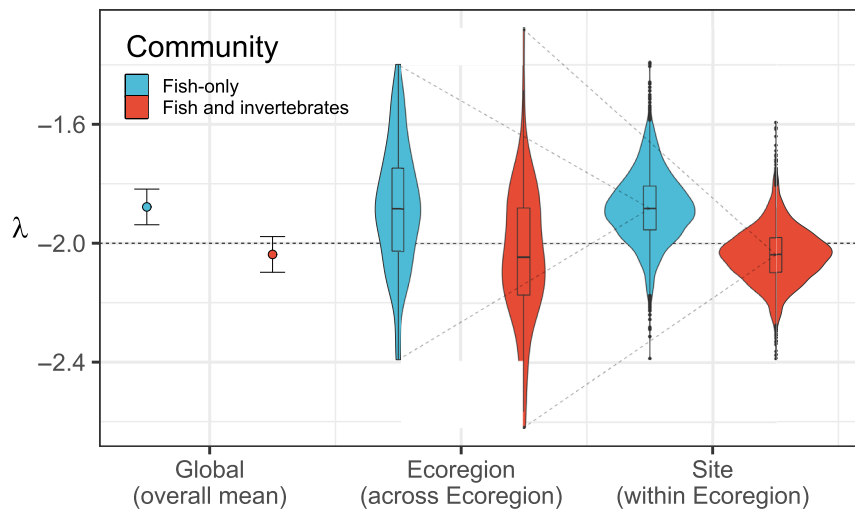


Figure 4 The contribution of spatial scale to abundance size spectra slope estimates. 'Ecoregion' refers to the variation among ecoregions globally in the linear mixed-effects model and 'Site' refers to the variation among individual reef sites within ecoregions. Dotted lines between the violins are added to emphasise that the variation at the site level represents the added variation after accounting for the variation at the ecoregion level. A horizontal dotted line at -2 is added to highlight the slope in previous studies based on pelagic studies.

reefs has been hampered by a lack of knowledge of an appropriate baseline, due to apparent discrepancies between the simplifying assumptions of size spectrum theory and lack of consistency across reef fish size spectra. Previous studies on local reef fish communities have shown slopes shallower than -2 (e.g. -1.13 to 1.95 , Robinson *et al.*, 2017; -1.75 , Ackerman *et al.*, 2004; -1.58 , Robinson *et al.*, 2016), potentially due to energetic subsidies (Trebilco *et al.*, 2013, 2016),

relatively greater levels of herbivory (Steneck *et al.*, 2017), or size-dependent habitat refugia (Rogers *et al.*, 2014), but still within the range of slopes estimated here for fish-only communities. Although not all these studies specifically aimed to test theory related to energy flow, the exclusion of invertebrates in these studies would have likely changed the slopes found. On average globally, we found that the inclusion of invertebrates into the community size spectrum steepened λ

from -1.88 to -2.04 ($\Delta\lambda = -0.16$), closer to the value of -2 that would be expected according to the 'biomass equivalence rule'. All sites in this study are subject to varying levels of human disturbance (e.g. fishing), and therefore we might expect that in the absence of fishing pressure, reef communities would have shallower size spectra than this -2 estimate.

The effect of including invertebrates varied geographically, with a much greater effect at higher latitudes. At the highest latitudes considered here (c. 60° N or S), fish-only size spectra had slopes that were more consistent with an inverted biomass pyramid (Trebilco *et al.*, 2013), where biomass increases with body size and trophic level. The opposite was true for invertebrate-only size spectra, whereby the steepest slopes were observed at the highest latitude (Figure 3a). These two taxonomic groups, however, are not independent food web entities and interact through competition and predation. Combining these two groups into the size spectrum led to consistency in the slope across latitudes. The resultant pattern translates to an even distribution of log-log biomass across all body sizes and across latitudes, supporting previous conjectures of biomass equivalence holding from bacteria to whales and from the tropics to the poles (Sheldon *et al.*, 1977; Kerr and Dickie, 2001). The latitudinal difference of including invertebrates is likely due to their dominance on temperate reefs, compared to more fish-dominated tropical reefs (Edgar *et al.*, 2017). Whilst fishing pressure is non-random across the globe (Anticamara *et al.*, 2011), it is unlikely to be the cause of the observed latitudinal patterns in the invertebrate inclusion effect, as we observe similar latitudinal patterns in sites within the most highly effective marine protected areas (Figure S6.1). Herbivores are also important on tropical reefs, and previous work has suggested that communities with a high biomass of herbivores, which do not feed according to size, should produce shallower size spectra (Robinson *et al.*, 2017), as a result of being able to obtain relatively larger body sizes due to less energy lost through transfer efficiency (Brown and Gillooly, 2003). Larger-bodied herbivores also have the added advantage of reduced predation risk from gape-limited predators (e.g. Mumby, 2006), leading to a relatively greater number of large-bodied individuals and a shallower slope. In this study, across the globe, the slope was steeper than would be expected according to that reasoning. These steeper slopes could be due to a combination of functionally distinct trophic pathways affecting energy availability (Dinmore and Jennings, 2004; Maxwell and Jennings, 2006), greater human impacts affecting tropical reefs (Graham *et al.*, 2005; Robinson *et al.*, 2017) (see also Figure S6.1), or other factors affecting local variation in reef size spectra (Edgar, 1994; Rogers *et al.*, 2014), and require further study.

A better understanding of the mechanisms underlying consistency and variability of slopes needs information on the spatial scales at which variability arises (Polishchuk and Blanchard, 2019). Investigation of different processes acting at local (e.g. sites) and larger spatial scales (e.g. ecoregions, global) should help to inform whether macroecological patterns are scale invariant (Rahbek, 2004; Connolly *et al.*, 2017). A first step is to assess how much variation occurs at each scale. Here, we found that variation from the overall global size spectrum slope was explained about equally at both the

ecoregion and site scales. Despite this scale-invariance of slope, the drivers of this variation still probably differ with scale, and our work opens the door for further studies into the factors shaping the size spectrum slope at different scales. At the ecoregion scale, drivers of variation likely include commercial fishing practices (e.g. Blanchard *et al.*, 2005), large-scale habitat loss (e.g. Morais *et al.*, 2020b), changing climate (e.g. Robinson *et al.*, 2019a,b) and environmental forcing (e.g. Heenan *et al.*, 2020). Potential drivers at the site scale include population processes (e.g. Barneche *et al.*, 2014, Barneche *et al.* 2016), local community interactions, eutrophication (e.g. Turner, 2001), coastal pollution (e.g. Ernesto *et al.*, 2010) and small-scale patchiness in fishing pressure related to human access (e.g. Robinson *et al.*, 2017; Campbell *et al.*, 2020).

Changes in size spectra slopes through time and space, have been used previously to assess changes in community and ecosystem health associated with the intensity of human activities (Dulvy *et al.*, 2004; Shin *et al.*, 2005; Wilson *et al.*, 2010; Graham *et al.*, 2005). Here, we used time-averaged size spectra on fished reefs, but future work on how size spectrum slopes vary with human activities (e.g. fishing and pollution) across time and space is needed. Reefs are also under pressure from the multifaceted effects of climate change (Graham *et al.*, 2007). Integrative modelling, and empirical and mechanistic studies (e.g. Barneche *et al.*, 2014; Morais *et al.*, 2020a), are all needed to disentangle the combined and relative influences of multiple anthropogenic stressors when contrasted with natural ecological variation affecting size spectra. Advancing this research goal would assist the development of predictive modelling tools for mapping changes on reefs, giving us a better idea of baseline reef size spectra and thus helping improve marine biodiversity policy and management (Stuart-Smith *et al.*, 2017).

In order to use the size spectrum slope as an indicator of reef health across systems, we must first understand the theoretical baseline slope (Jennings and Blanchard, 2004), from which environmental, ecological and anthropogenic drivers of the remaining variation in slopes can be estimated. Our study highlights the importance of including invertebrates in reef size spectrum analyses for both the estimate of the baseline and for reducing variability in the slope estimates. When accounting for the invertebrates in the reef community, we show extremely high consistency in the size spectrum slope, supporting the generality of the biomass equivalence rule for reef communities at the global scale.

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AUTHORSHIP

FJH designed the analysis, analysed the data and led the writing of the manuscript. All authors contributed to critical feedback, interpretation and substantial revisions of the paper.

PEER REVIEW

The peer review history for this article is available at <https://publons.com/publon/10.1111/ele.13661>.

DATA AVAILABILITY STATEMENT

Code for the analysis, and to recreate all figures, is available at https://github.com/FreddieJH/inverts_size_spec. Most of the data used in this study are publicly available at <https://reeflife-survey.com/survey-data/>, although body size information is unavailable until public release of the redeveloped database in mid-2021. In the interim, these data can be provided upon request by contacting enquiries@reeflifesurvey.com (<https://doi.org/10.5281/zenodo.4287663>).

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SUPPORTING INFORMATION

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