Species richness and identity both determine the biomass of global reef fish communities

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Changing biodiversity alters ecosystem functioning in nature, but the degree to which this relationship depends on the taxonomic identities rather than the number of species remains untested at broad scales. Here, we partition the effects of declining species richness and changing community composition on fish community biomass across >3000 coral and rocky reef sites globally. We find that high biodiversity is 5.7x more important in maximizing biomass than the remaining influence of other ecological and environmental factors. Differences in fish community biomass across space are equally driven by both reductions in the total number of species and the disproportionate loss of larger-than-average species, which is exacerbated at sites impacted by humans. Our results confirm that sustaining biomass and associated ecosystem functions requires protecting diversity, most importantly of multiple large-bodied species in areas subject to strong human influences.
Extinctions in nature are often biased towards species that are rare and large-bodied\textsuperscript{1,2}, yet experiments designed to predict the ecosystem consequences of such extinctions have typically removed species at random\textsuperscript{3}. The few experiments that have employed more realistic extinction scenarios by removing rare and functionally-unique species have revealed greater changes in ecosystem functioning than from random losses\textsuperscript{4,5}. Observational datasets provide a unique opportunity to examine how the nonrandom loss of species, particularly those that are too large or too rare to feasibly manipulate in experiments, affects ecosystem processes under natural circumstances. Such datasets reflect the cumulative processes—including natural environmental forcing, local losses and gains of species, and anthropogenic impacts—that generate gradients of biodiversity observed in the real world\textsuperscript{6}.

Even as observational evidence supports strong links between species richness and ecosystem functioning, specifically biomass production\textsuperscript{7}, it remains unclear how much of this relationship stems from reducing the number of species (at random) versus the specific traits proxied by the identities of species. This distinction is important if extinctions are biased towards individual species that contribute disproportionately to functioning, such as large-bodied fishes\textsuperscript{8}. Selective removal of such species through overexploitation and sensitivity to human disturbance might be expected to have a larger-than-anticipated influence on community productivity compared to random losses\textsuperscript{9}.

Separating the effects of changing species richness and community composition on ecosystem properties in observational data has proved historically challenging\textsuperscript{10}. One useful approach is to mathematically partition the total change in an aggregate property, such as community biomass, into component factors\textsuperscript{11–13}. Inspired by these earlier approaches, we present a partitioning of the difference in standing stock biomass between pairs of communities—a reference site vs. a comparison site—into five additive components representing the effects of both gains and losses in species richness and composition, and the effects of any remaining influences such as environmental and ecological factors that are not already captured in the elements of community diversity.

We must first describe the background of our decomposition (Fig. 1). Species that are absent or “lost” in the comparison community relative to the reference community contribute to the “richness loss” and the “compositional loss” components of the biomass difference, hereafter \textit{RICH-L} and \textit{COMP-L}. To quantify these components, it is necessary to choose a frame of reference for the “expected” biomass loss per absent species, under the null hypothesis that there is nothing unusual about the species that are absent. Here, we use the average biomass of species present in both communities (“shared species”) as this reference value (for more details, see Methods and Supplementary Materials). \textit{RICH-L} is the amount of biomass lost based on the number of species absent from the comparison community weighted by this “expected” biomass per species. The actual loss in biomass at a community level can differ due to deviations in the biomass of the species that are absent compared to the species that are shared, which is captured by the \textit{COMP-L} term. This term reflects the change in biomass between communities that is attributable to the kinds of species that are absent rather than simply the number of absent species.

The third and fourth terms reflect changes in biomass from species present at the comparison site but absent from the reference site (“gained” species): \textit{RICH-G} is the expected gain in biomass given the number of gained species (again based on the average biomass of species shared among both sites), and \textit{COMP-G} reflects any additional biomass change from the gained species being different in average biomass from shared species. The addition of the previous four terms (\textit{RICH-L + COMP-L + RICH-G + COMP-G}) summarizes the total difference in community biomass between the two sites due to both losses and gains in the number (richness) and identities (composition) of species, which we call the “total diversity effect” or \textit{DIV}.

The fifth and final term aims to estimate all additional factors contributing to the difference in biomass between sites that is not due to diversity, broadly speaking. These include, for example, differences in the number of individuals and their sizes, predator–prey interactions, resource availability, habitat quality and complexity, and/or underlying environmental drivers, collectively called the “context-dependent effect” (or \textit{CDE}). The \textit{CDE} term considers only differences in biomass among species shared by both communities rather than species, which are lost or gained between communities. We note that the \textit{CDE} term cannot mechanistically distinguish among these various influences, as has been done using other experimental and observational datasets\textsuperscript{14,15}, but its relative magnitude can be assessed against other terms to quantify the importance of losses or gains of whole populations (i.e., \textit{RICH} and \textit{COMP}) versus changes in individual biomass or abundance\textsuperscript{16}. By extension, environmental factors can also drive changes in species richness and composition, so \textit{CDE} should not be interpreted as exclusively indicative of environmental forcing. Rather, it can be thought of as the cumulative influence of all extrinsic factors on fish community properties that is independent of their realized effects on the observed differences in richness and composition.

Here, we applied this decomposition to quantitatively assess the relative contributions of species richness, composition, and context-dependent effects to reef fish community biomass, a common proxy for secondary production, derived from a fisheries-independent global dataset of underwater visual censuses collected by the Reef Life Survey program (www.reelfishesurvey.com). We selected reefs with the highest biomass to serve as the reference sites, as high fish biomass is a desirable state and often the goal of conservation and restoration efforts. We then compared these high-biomass sites to all other sites within a 100-km radius (representing a potential upper limit on direct dispersal\textsuperscript{17}) to determine whether variation in elements of biodiversity contributes to other nearby sites not reaching their biomass potential. We conducted 173 comparisons of reference sites (i.e., those with the highest biomasses) against a total of 2867 comparison sites, with each reference site having on average 16.6 (range = 5–136) comparison sites (Fig. 2). Sites were split near equally between tropical realms (mostly coral-dominated; \textit{n} = 1603) and temperate realms (mostly rocky reefs; \textit{n} = 1437). We go on to show that loss of biomass at the comparison site is most associated with biodiversity loss, which can be further attributed to both losses in the total number of species and the absence of uniquely large contributors to biomass. These effects were most pronounced in areas with high human populations, implicating humans as a primary driver of changes in fish biomass across the seascape.

\textbf{Results and discussion}

When we applied the decomposition (Fig. 1) to our data, we found that low reef fish biomass between reference and comparison sites was associated predominantly with changes in richness and composition rather than environmental and other drivers (Fig. 3). The cumulative biodiversity effect \textit{DIV} was 5.7x stronger (95% confidence intervals: [5.1, 6.5]) than the \textit{CDE} effect (Fig. 3). Furthermore, the biodiversity effect on community biomass was driven almost entirely by the loss (\textit{L}) rather than the gain (\textit{G}) of species (Fig. 3). We note that this finding is not an artifact of the partitioning equation: indeed, high-biomass
the spatial extent over which the local comparisons are conducted (15–100 km, Supplementary Fig. 1), with slightly weaker effects for 15-, 25-, 50- and 100-km radii, respectively). The proportion of species shared among any two sites, however, remained remarkably consistent from scales of 15–100 km (49.1–51.5% of the total community), and reference sites are not mathematically constrained to have more species, but apparently do in nature. The majority (69%) of comparisons to the reference site exhibited a net loss rather than a net gain of species (−11.4 ± 0.7% average (± SE) reduction in the number of species across all comparisons). Thus, our analysis indicates that differences in biodiversity, primarily those that reflect the absence of species from site to site, are the primary determinants of fish community biomass across the world’s coral and rocky reefs over the scale of 10 s of kilometers.

The strongest effect of biodiversity loss on community biomass was due to changes in species composition. Loss of biomass attributable to changing species identities (COMP-L) was 3.8x greater (95% CIs: [3.5, 4.1]) than that based on the “expected” or average outcome of removing species at random (RICH-L, Fig. 3). In approximately three-quarters of cases, the COMP-L term was negative, indicating that compositional effects were primarily due to the loss of greater-than-average contributors to biomass (i.e., large-bodied fishes) and/or those with extreme abundances (e.g., schooling fishes). Thus, we demonstrate that species’ identities and their traits (e.g., body size, gregariousness) are key factors in predicting the ecosystem consequences of species’ losses. We note that the strong effect of species identity does not negate the contributions of richness: indeed, reductions in the number of species necessarily change composition, and thus both operate simultaneously to influence community biomass.

Our finding that biodiversity loss is the primary determinant of fish community biomass among reefs within a region is robust to the spatial extent over which the local comparisons are conducted (15–100 km, Supplementary Fig. 1), with slightly weaker effects of composition observed at reduced scales (mean COMP-L = −0.38, −0.38, −0.40, and −0.43 for 15-, 25-, 50- and 100-km radii, respectively). The proportion of species shared among any two sites, however, remained remarkably consistent from scales of 15–100 km (49.1–51.5% of the total community), and...
thus our results do not appear to be biased by an increased overlap in composition between communities at smaller scales or by more rare or unique species at larger scales. Additionally, our results are largely consistent across temperate and tropical realms (Supplementary Fig. 2), with two notable exceptions. First, we observed slightly more extreme declines in biomass due to richness losses in tropical areas, presumably because tropical sites have, on average, 2.8x more species than temperate ones and therefore possess more scope for change. Second, we observed similarly larger declines in biomass due to compositional losses at temperate sites, reflecting that individuals there are, on average, 1.5x larger than at tropical sites and therefore more likely to contribute biomass in excess of the “expected” or average species. Nevertheless, these two differences cancel each other out such that the total diversity effect (DIV) does not differ between the two realms (Supplementary Fig. 2).

To understand the potential drivers of our results, we used random forest analysis to identify the environmental and anthropogenic variables most associated with each component of our decomposition. The top-ranked predictor for biomass loss associated with richness and compositional loss was consistently human population density, particularly for the COMP-L and DIV terms (Supplementary Fig. 3). Sites near even a minimal human population exhibited lower total fish biomass, smaller-bodied fishes, and fewer fish species (P < 0.001 in all cases, from generalized linear mixed-effects models regressing fish community characteristics against human density) (Fig. 4). Our index provides a proxy for a range of human impacts, including direct removal of species and biomass through fishing and the indirect effects of numerous factors such as reduction in suitable habitat and food availability. In contrast, the context-dependent effect (CDE) was most influenced by environmental variables, specifically salinity, nutrients (dissolved phosphorus), and temperature variation (Supplementary Fig. 3). This result would be expected when environmental variation affects the weight and abundance of individuals but does not exclude species entirely, clarifying our interpretation of the CDE term as capturing underlying gradients that might alter the capacity of species to acquire resources and produce biomass. Environmental factors also affected the diversity terms (e.g., phosphorus for RICH-L, Supplementary Fig. 3), albeit to a lesser degree, reflecting that environment can also be responsible for driving local species absences or latent indicate a degraded state which certain species find unsuitable.

The effects of species identity on community biomass recovered by our analysis are most likely related to selective removal of large species, as found in prior studies of fisheries, rather than changes in the number of individuals. To test this hypothesis, we partitioned the contributions of species unique to reference and comparison sites as a function of size class. As expected, the unique contributors at the most productive reference sites were concentrated in the largest size class (>200 cm observed total length) (Fig. 5). Of species unique to the reference sites, 56.4% of the biomass was contributed by fishes >200 cm in length, whereas this size class contributed only 19.0% of the total biomass at the comparison sites. These observations also explain the bimodal distribution of the COMP-L term: the majority of highly negative values reflect losses of larger-than-average species from reference sites, as opposed to smaller-than-average fishes which are
The lack of large species at comparison sites could have resulted from the inclusion of species that are more mobile and have larger home range sizes, and just so happen to have occurred at a particular reference site. To test whether the contributions of such species might have unduly influenced our results, we re-ran our analysis after removing all 156 pelagic/non-site attached species (~5% of all species, classifications from 22). This subsequent analysis yielded nearly identical results to those from the full dataset (Supplementary Fig. 4).

We set the reference site as the highest biomass site to simplify the interpretation of biomass variation across communities because this makes the net change always negative 13. Analyzing the data in this way allows us to ask whether losses in biomass are due primarily to cumulative changes in diversity, broadly defined (DIV), or losses within species (CDE). However, this choice of reference may also bias towards a compositional effect, since larger species that contribute most to biomass are more likely to be found at high-biomass sites. To test for this bias, we repeated our analysis on 1000 simulated versions of our original dataset that randomized composition while holding species numbers and per capita contributions constant. The value of COMP-L in the observed data was always more negative than those returned by our simulations ($P < 0.001$ based on a one-tailed comparison of means; Supplementary Fig. 5). Thus, our results do not appear to be an artifact of how the reference community was defined (see also Supplementary Materials for outcomes when alternatively establishing the reference as the most speciose sites). Instead, our
findings reflect the inherent organization of natural systems: high-biomass sites support large-bodied species that are not often found elsewhere.23

Hundreds of experimental manipulations show that biodiversity loss is among the pre-eminent factors contributing to variation in community biomass in a wide range of ecosystem types and taxa driven by both changes in richness and composition16,24. Our study extends theoretical inferences from this work to highly diverse, natural ecosystems across a range of contexts. A synthesis of >200 terrestrial plant manipulative experiments revealed roughly equivalent contributions by species richness and species identity10, a result which matches our findings for global reef fishes. Here, we also provide a new mathematical approach to address this question. It is important to note that, like other recent ecological decompositions, ours is not a mechanistic model.12 Instead, it integrates across all potential mechanisms driving the aggregate loss and gain of species, including their response to abiotic drivers, and how they affect ecosystem functioning. Therefore, our application permits disentangling richness vs. compositional effects operating in these communities.8,25,26 We now show these effects arise (e.g., through dominance, complementary resource use, indirect interactions such as facilitation, etc.) requires further investigation.26 Similarly, assessing processes such as production (for which we substitute standing stock biomass as a proxy) using this new partitioning is necessary if the goal is to link diversity to more robust measures of ecosystem functioning.

Our results strengthen previous findings that human actions have large impacts on fish biomass: many studies have demonstrated that humans are selectively removing large-bodied fishes from the ocean, largely on the basis of fishing data, resulting in a loss of functional diversity and ecosystem-scale consequence of this phenomenon: reductions in large-bodied species leads to greater divergence between high- and low-biomass sites. Our data, therefore, provide compelling new evidence that selective removal of species with particular traits and reduction of local biodiversity across the seascape through collective human impacts profoundly alter the structure and function of whole reef communities worldwide. This result has important implications for the continued management of reef ecosystems: conservation practices that are tied to management of reef ecosystems: conservation practices that are tied to the restoration of high-performing species30 and those that aim to preserve the diversity of whole communities (e.g., marine protected areas)31,32 are both required to maximize the provision of a critical ecosystem function in an increasingly human-dominated world.33

Decomposition of difference in ecosystem functioning. Our equation was inspired by previous decompositions, principally the Price equation originally derived in the field of evolutionary biology as a means of understanding how genetic and environmental influences on phenotypic change over time.27, Fox and Kerr modified the Price equation to describe how the difference in the ecological function between two communities can be decomposed into components with different ecological interpretations. We follow a similar approach but use a different decomposition whereas the resulting components are similar to, but not the same as, the components proposed by Fox and Kerr.12

We begin by assuming that the ecological function of the community, such as biomass, is a simple additive function of the contributions of its constituent species. We then compare two communities, one of which we refer to as the reference community and the other we refer to as the "comparison" community. The species present in the reference community can be classified into two types: species that are unique to the reference community (i.e., not present in the comparison community) and those that are in common with the comparison community. Let $s_{uR}$ be the number of unique species in the reference community, and $s_{c}$ be the number of species in common between the two communities. Let $z_{uR}$ be the average ecological function contributed per unique species to the reference community, and $z_{c}$ be the average ecological function contributed per shared species in the reference community. The total ecological function $T_{R}$ of the reference community can thus be decomposed as:

$$T_{R} = s_{uR}z_{uR} + s_{c}z_{c}$$  \hspace{1cm} (1)

with a similar interpretation to Eq. (1). Though these are $s_{uR}$ species in common between the two communities, the average per species contribution need not be the same in the two communities (i.e., $z_{uR}$ may differ from $z_{c}$).

The species in common between the two communities can serve as a reference point for comparison between communities. It is useful to define $\delta_{u} = z_{uR} - z_{c}$ and $\delta_{c} = z_{c} - z_{uR}$ as the difference in average ecological function per species of unique versus shared species in reference and comparison communities, respectively. From this perspective, we consider the average ecological function of a species unique to the reference community as being equal to the average ecological function of shared species (as measured in the same community) plus the deviation from this value $z_{uR} = z_{c} + \delta_{c}$. Using this equality and the analogous one for $z_{uR}$, the difference in the ecological function between two communities can be decomposed as:

$$\Delta T = T_{R} - T_{C} = -s_{uR}\delta_{u} + s_{uR}\delta_{c} + s_{c}\delta_{u} + s_{c}\delta_{c}$$  \hspace{1cm} (3)

The first two terms represent the loss in ecological function in the comparison community due to the loss of species that are unique to the reference community. Specifically, the first term represents the loss in ecological function due to the absence of unique species if these species had the same average value of functioning as each of the shared species. In other words, it is the amount by which biomass is expected to decline if species were interchangeable. Therefore, we interpret this term as the "richness loss" or the loss in functioning due strictly to the loss of species: $RICH-L (= -s_{uR}\delta_{u})$. It will always be negative, assuming there is at least one species unique to the reference population. In cases where $z_{uR} \geq z_{c}$, it is possible for RICH-L to exceed the total functioning observed at the reference site, which complicates interpretation of the raw values. In this case, it is useful to consider only the relative quantities (each component is scaled by the sum of the absolute values of all components). We note that this situation arises only 41 times out of 2867 comparisons in our analysis, and removing these cases has no effect on our findings. We advise future applications be aware of this potential issue for its influence.

Methods

Reef life survey. Reef fish communities were censused by a combination of experienced marine scientists and trained recreational SCUBA divers using globally standardized Reef Life Survey methods. All surveys were undertaken on 50 m long transects laid along a contour (at consistent depth) on predominantly hard substrate (usually rocky or coral reef) in shallow waters (depth range of transects 1 to 20 m, average ~7.2 m). Full details of fish census methods, data quality, and training of divers are provided in refs.22,24,34 and in an online methods manual (www.reelfishesurvey.com). Fish abundance counts and size estimates per 500 m² transect area (2 x 250 m² blocks) were converted to biomass using length–weight relationships for each species obtained from Fishbase (www.fishbase.org). In cases where length–weight relationships were provided in Fishbase using standard length or fork length, rather than total length as estimated by divers, length–length relationships provided in Fishbase allowed conversion to the total length. For improved accuracy in biomass assessments, observed sizes were also adjusted to account for the bias in divers’ perception of fish size underwater using an empirical calibration.26 Length–weight coefficients from similar-shaped close relatives were used for those species where length–weight relationships were not available in Fishbase. All transects were collapsed into a single average value of biomass for each species at a location to account for any differences in the total number of transect surveys performed.

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shared species: Rich-G (\(= +_{s}A_{s}\)). It is always positive, assuming the comparison community has at least one unique species. The fourth term, COMP-G (\(= +_{s} A_{s}\)), reflects the difference in composition (with respect to average value) of gained versus shared species. This term can be positive or negative, being positive if the gained species have a higher per species value than the shared species. The final term focuses on the changes in biomass considering only the species that are present in both communities. This can be thought of as holding richness and composition constant and considering changes in the community biomass that are present in both communities. This can be thought of as holding richness constant, because both equations partition the same total quantity. Thus, it is important to note that using either decomposition yields the same inference with respect to comparisons of DIV and CDE.

Our decomposition differs from Fox and Kerr’s because the two approaches differ in the points. We take the perspective that the shared species form the basis for comparison between two communities, so we then evaluate the average value of a unique species with respect to its deviation from an average value of a shared species. In contrast, Fox and Kerr effectively evaluate the average value of a unique species with respect to its deviation from the average value of any species in that community (averaging over both unique and shared species). In both decompositions, the “composition” components only exist if there is some difference in the average value of shared and unique species. We prefer our decomposition for this case because it works with that difference directly rather than indirectly via the difference between unique and all species (which is the average of unique and shared species). Moreover, our composition makes intuitive sense that the function of the “average” species is determined by the ones that are known to exist at both sites. A full comparison of the Fox and Kerr formulation and ours is provided in the Supplementary Materials.

Statistical analysis. A general function to conduct our new decomposition from a site-by-species biomass matrix, and a second function to perform the simulations, can be found here: https://gitlab.com/jlechle/76076c7c52dkoecb87113dcdb6b6b4. We first ordered all sites by decreasing total biomass. Beginning with the highest biomass site of all sites as the first reference site, we identified all other sites within a certain spatial extent (15–25, 30–, or 100-km) to serve as the comparison sites. Setting the reference to be the site with the highest community biomass constrains the sum of the terms to be negative. This choice simplifies the language used to discuss the output and allows us to speak directly to the consequences of real-world activities like overharvesting (and their implications).

We then computed the components for each set of comparisons. We standardized the output to the same scale (\(\ldots, 1\)) by first taking the sum of the absolute value of all components, and then dividing each component by this value. This relativization was done to account for the fact that raw biomass may differ substantially among sites and regions and to make our results comparable across the entire dataset. Once the scaled components were computed, the reference and comparison sites were removed from the ordered list from any further comparisons to prevent any bias that might arise from including the same site multiple times. We then moved onto the next most productive site in the list, identified the comparison sites within 100 km, computed the components, and so on, until all sites were analyzed. From these individual comparisons, we computed the means of all components while omitting any reference sites for which there were fewer than five common species. We also computed the means for each reference site and then took the grand mean of these average values, although this additional level of aggregation did not qualitatively change our results (Supplementary Fig. 6). We have chosen to present the raw values in the main text to demonstrate the level of variability inherent in the individual comparisons, which might otherwise be condensed by showing only the means for each reference site. We repeated the analysis over multiple spatial radii to assess whether the spatial extent and therefore the size and composition of the species pool might influence our results.

We calculated the relative strength of the total diversity effect vs. the context-dependent effect for each comparison after the DIV and CDE, and our compositional vs. richness losses as:

\[ Q = \left( \frac{\mu_{s}A_{s} - \mu_{s}A_{s}}{\mu_{s}A_{s}} \right) \]

(4)

In this case, \(Q = (\text{DIV} + \text{RICH-L})/\text{RICH-L}\), which reduces to the average value of unique species relative to the average value of shared species at the reference site. This quantity reflects the magnitude to which species unique to the reference site contribute to biomass relative to the “expected” contribution per species. To avoid biases associated with averaging ratios, we report the geometric mean of both quantities. Bootstrapped 95% confidence intervals were derived by randomly resampling DIV/CDE and Q for a total of 5000 times. For DIV/CDE, some value was valid if we excluded them in both the original and bootstrap samples. As an alternative approach that focused on the magnitude of effect, we examined the absolute value of DIV | CDE |. In this case, the ratio was 6.9x with bootstrap 95% CIs of [6.2, 7.7].

To explore the drivers of the components of our decomposition, we applied random forest analysis to account for potential collinearity and interactions among the suite of predictors previously selected in ref. 39. Depth was recorded on the surveys while the following predictors were obtained from the combination of remote sensed and in situ measurements compiled in the Bio-ORACLE database: mean, minimum, maximum, and range of sea surface temperature; mean, minimum and maximum for surface chlorophyll-a; mean salinity; mean PAR; mean dissolved oxygen; mean nitrate concentration; mean phosphate concentration. Finally, an index of human population density was calculated by fitting a smoothly tapered surface to each settlement point on the year 2010 world-population density grid using a quadratic kernel function described previously (ref. 31).

Random forests were fit using the default settings in the randomForest package in R version 4.1.0. Variable importance was determined using the percent increase in the mean-square error after randomly permuting the predictor of interest for each tree in the random forest, averaging the error of the models, and then computing the difference relative to the accuracy of the original model.

Null simulations. A key finding of our analysis is that compositional losses are considerably greater than losses due to other aspects of the reef fish community. We wanted to evaluate the possibility of whether such a result could be an artifact of applying our decomposition to a dataset in which we assign the site with the higher total biomass as the “reference” community and the site with lower total biomass as the “focal” community. To do so, we conducted simulations in which we created communities with species richness values matching the observed data, but for community compositions that were random. Following the same procedure we used with the real communities, we applied our decomposition to these simulated communities to generate null distributions for the average values of each of the components when community composition is random. Comparing our observed values to these null distributions tells us if the values of the compositional components (or indeed any component) we observed arose as an artifact of our procedure or, alternatively, because high-biomass sites actually contain more high-biomass species than expected under random community assembly.

Our simulation procedure focused on the site-by-species biomass matrix from each set of comparisons used in the main 100-km analysis. We divided this matrix by the corresponding site-by-species abundance matrix to yield the observed per capita contribution of each species in each community. We then averaged the per capita contributions of each species across all communities where the species was present to yield a single vector representing mean per capita contributions for all S species within that set of comparisons.

We initially constructed each simulated community by populating it with every species in the region (“maximum richness”). To determine the biomass of each species in each community we applied the following procedure. First, we identified the minimum and maximum abundance of each species in all communities where it is present. For a single community, we sampled an integer from the minimum to maximum abundance for each species. Second, we then randomly subsampled a single vector for representing mean per capita contributions for all species within that community. We then repeated this for all n communities in the original site-by-species matrix and bound these vectors together in a new “maximum richness” version of the site-by-species matrix. For the i-th row (community) in the original dataset, we calculated the richness, \(s_{i}\). We then randomly subsampled \(s_{i}\) species at random from the simulated “maximum richness” site-by-species matrix and set the biomass to zero for all remaining species to zero. We repeated this for each community to yield a simulated “observed richness” site-by-species matrix with the same dimensions as the original biomass matrix. This procedure ensures that richness is held at the observed levels and that the biomass contribution of each species is within the observed range.

As predicted, the communities were indeed constrained with respect to composition as our goal was to test whether the observed compositional effects in the raw data are significantly different than under this null hypothesis with respect to composition. Thus, using the simulated “observed richness” site-by-species matrix, we computed the (scaled) components as we had with the real data and then measured across all samples per means expected using a one-tailed t-test to determine whether the observed components were more or less extreme than would be expected by chance.
References

1. Pimm, S. L., Jones, H. L. & Diamond, J. On the risk of extinction. Am. Nat. 132, 757–785 (1988).
2. Payne, J. L., Bush, A. M., Heim, N. A., Knope, M. L. & McAuley, D. J. Ecological selectivity of the emerging mass extinction in the oceans. Science 330, 298–301 (2010).
3. Srivastava, D. S. & Vellend, M. Biodiversity-ecosystem function research: is it relevant to conservation? Annu. Rev. Ecol. Evol. Syst. 36, 267–294 (2005).
4. Solan, M. Extinction and ecosystem function in the marine benthos. Science 306, 1177–1180 (2004).
5. Zavaleta, E. S. & Hulvey, K. B. Realistic species losses disproportionately reduce grassland resistance to biological invaders. Science 306, 1175–1177 (2004).
6. Reacken, M. E. S. & Low, N. H. N. Realistic losses of rare species disproportionately impact higher trophic levels. Ecol. Lett. 15, 461–467 (2012).
7. Duffy, J. E., Godwin, C. M. & Cardinale, B. J. Biodiversity effects in the wild are common and as strong as key drivers of productivity. Nature 549, 261–264 (2017).
8. Estes, J. A., Heithaus, M., McCauley, D. J., Rasher, D. B. & Worm, B. Megafaunal impacts on structure and function of ocean ecosystems. Annu. Rev. Environ. Resour. 41, 83–116 (2016).
9. Mellin, C. et al. Humans and seasonal climate variability threaten large-bodied coral reef fish with small ranges. Nat. Commun. 7, 10491 (2016).
10. Cardinale, B. J. et al. The functional role of producer diversity in ecosystems. Am. J. Bot. 98, 572–592 (2011).
11. Fox, J. W. Using the Price Equation to partition the effects of biodiversity loss on ecosystem function. Ecology 87, 2687–2696 (2006).
12. Fox, J. W. & Kerr, B. Analyzing the effects of species gain and loss on ecosystem function using the extended Price equation partition. Oikos 121, 416–424 (2012).
13. Winfree, R., Fox, J. W., Williams, N. M., Reilly, J. R. & Cariveau, D. P. Abundance of common species, not species richness, drives delivery of a real-world ecosystem service. Ecol. Lett. 18, 626–635 (2015).
14. Genung, M. A. et al. The relative importance of pollinator abundance and species richness for the temporal variance of pollination services. Ecology 98, 1807–1816 (2017).
15. Genung, M. A., Fox, J. & Winfree, R. Species loss drives ecosystem function in experiments, but in nature the importance of species loss depends on dominance. Glob. Ecol. Biogeogr. 29, 1–11 (2020).
16. Hooper, D. U. et al. A global synthesis reveals biodiversity loss as a major driver of ecosystem change. Nature 486, 105–108 (2012).
17. Kinlan, B. P. & Gaines, S. D. Propagation dispersal in marine and terrestrial environments: a community perspective. Ecology 84, 2007–2020 (2003).
18. Duffy, J. E., LeFebvre, J. S., Stuart-Smith, R. D., Navarrete, S. A. & Edgar, G. J. Biodiversity enhances reef fish biomass and resistance to climate change. Proc. Natl Acad. Sci. USA 113, 6230–6235 (2016).
19. Graham, N. A. J., Newell, S. A. & Jennings, S. Abundance of common species, not species richness, drives delivery of a real-world ecosystem service. Ecol. Lett. 18, 626–635 (2015).
20. Jackson, J. B. C. et al. Historical overfishing and the recent collapse of coastal ecosystems. Science 293, 629–637 (2001).
21. Zgliczynski, B. J. & Sandin, S. A. Size-structural shifts reveal intensity of exploitation in coral reef fisheries. Ecol. Indic. 73, 411–421 (2017).
22. Stuart-Smith, R. D. et al. Integrating abundance and functional traits reveals new global hotspots of fish diversity. Nature 501, 539–542 (2013).
23. Stevenson, C. et al. High apex predator biomass on remote Pacific islands. Coral Reefs 26, 47–51 (2007).
24. Cardinale, B. J. et al. Biodiversity loss and its impact on humanity. Nature 489, 326–326 (2012).
25. Loreau, M. & Hector, A. Partitioning selection and complementarity in biodiversity experiments. Nature 412, 72–76 (2001).
26. Clark, A. T. et al. How to estimate complementarity and selection effects from an incomplete sample of species. Methods Ecol. Evol. 10, 2141–2152 (2019).
27. McCauley, D. J. et al. Marine defaunation: animal loss in the global ocean. Science 347, 1255641 (2015).
28. Mora, C. et al. Global human footprint on the linkage between biodiversity and ecosystem functioning in reef fishes. PLoS Biol. 9, e1000606 (2011).
29. Graham, N. A. J. et al. Human disruption of coral reef trophic structure. Curr. Biol. 27, 231–236 (2017).
30. Cinner, J. E. et al. Bright spots among the world’s coral reefs. Nature 535, 416–419 (2016).
31. Palumbi, S. R. et al. Managing for ocean biodiversity to sustain marine ecosystem services. Front. Ecol. Environ. 7, 204–211 (2009).
32. Edgar, G. J. et al. Global conservation outcomes depend on marine protected areas with five key features. Nature 506, 216–220 (2014).
33. Topor, Z. M., Rasher, D. B., Duffy, J. E. & Brandl, S. J. Marine protected areas enhance coral reef functioning by promoting fish biodiversity. Conserv. Lett. 12, 1–9 (2019).
34. Edgar, G. J. & Stuart-Smith, R. D. Systematic global assessment of reef fish communities by the Reef Life Survey program. Sci. Data 1, 1–8 (2014).
35. Edgar, G. J. & Stuart-Smith, R. D. Ecological effects of marine protected areas on rocky reef communities-A continental-Scale analysis. Mar. Ecol. Prog. Ser. 388, 51–62 (2009).
36. Edgar, G. J., Barrett, N. S. & Morton, A. J. Biases associated with the use of underwater visual census techniques to quantify the density and size-structure of fish populations. J. Exp. Mar. Bio. Ecol. 308, 269–290 (2004).
37. Price, G. R., others. Selection and covariance. Nature 227, 520–521 (1970).
38. Fox, J. W. Using the Price Equation to partition the effects of biodiversity loss on ecosystem function. Ecology 87, 2687–2696 (2006).
39. Breiman, L. Random forests. Mach. Learn. 45, 5–32 (2001).
40. Tyberghien, L. et al. Bio-ORACLE: a global environmental dataset for marine species distribution modelling. Glob. Ecol. Biogeogr. 21, 272–281 (2012).
41. Silverman, B. W. Density Estimation for Statistics and Data Analysis (Chapman & Hall, 1986).
42. Law, A. & Wiener, M. Classification and regression by randomForest. R. N. 2, 18–22 (2002).
43. R: a language and environment for statistical computing (R Core Team, 2021).

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G.J.E., R.D.S.-S. led the data collection; J.S.L., A.F.A., G.J.E., R.D.S.-S., A.E.B., C.W., S.J.B., S.K., S.D.L., J.E.D. and D.B.R. developed the concepts; J.S.L. and A.F.A. ran the analyses; G.J.E. and R.D.S.-S. performed the experiments; G.J.E. and R.D.S.-S. wrote the paper with input from all authors.

Competing interests

The authors declare no competing interests.

Additional information

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