**Habitat, fishing and biodiversity control grazing potential on coral reefs**

**Authors**

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**Abstract**

**Introduction**

Herbivory is crucial to ecosystem function and community structure, playing a key role in cycling nutrients (Metcalfe et al. 2014), regulating species diversity and productivity (Royo et al. 2010, Prieditis et al. 2017), and controlling habitat regime shifts (Zimov et al. 1995; Hughes et al. 2007; Keesing and Young 2014). Herbivory processes are generally measured at local scales relevant to individual behaviours and population sizes, which restricts our understanding of how ecosystem functions operate across larger spatial? scales. Furthermore, anthropogenic pressures typically impact ecosystem processes, including herbivory, across much larger areas. Therefore, developing our understanding of both natural and anthropogenic drivers on herbivory at large scales requires us to integrate fine-scale herbivory observations with macroecological datasets. Such analyses are particularly relevant for coral reef ecosystems, which are facing multiple damaging human pressures and where herbivory is a key ecosystem function (Hughes et al. 2007; Ledlie et al. 2007; Cheal et al. 2010).

On tropical coral reefs, top-down control of algae and promotion of calcifying coral taxa are key functions primarily performed by a diverse guild of herbivorous fishes (Bellwood et al. 2004). Within this group, observations of feeding morphology and behaviour have been used to categorise species into two distinct grazing functions: cropping and scraping (Bellwood and Choat 1990; Polunin et al. 1995; Green and Bellwood 2009). Cropping species graze filamentous turf algae, which maintains algae in cropped states preventing transitions to fleshy macroalgae (Arnold et al. 2010). Scraping species graze dead coral substrate to consume microscopic epiphytes and epilithic and endolithic phototrophs (Choat and Clements 2018). By removing detritus and epilithic algal matrix (EAM), scrapers promote coral recruitment by clearing settlement space (Bonaldo & Hoey 2014). Combined, cropping and scraping are considered essential functions which help sustain coral-dominated states (Bellwood et al. 2004).

Mature algae can proliferate in the absence of grazing pressure (Mumby et al. 2006; Burkepile and Hay 2008), and correlative analyses of fished reef ecosystems have provided evidence of grazing biomass thresholds below which reefs become algae dominated (Graham et al. 2015; Jouffray et al. 2015; Robinson et al. 2018). Herbivore populations are overexploited across the tropics (Edwards et al. 2014), which has compromised grazing functions on reefs which fail to maintain herbivore biomass thresholds (Bellwood et al. 2011, Graham et al. 2015, Robinson et al. 2018). However, fishing effects can be compounded by bottom-up influences on herbivore assemblages (Russ et al. 2015), whereby species-specific habitat associations produce spatial structuring of herbivore populations among different habitat types (Doropoulos et al. 2013) and benthic compositions (Hoey & Bellwood 2008; Heenan et al. 2016). Bottom-up control of fish populations may be particularly strong when fish rely on habitat for both structure and food, such as croppers which are generally small and particularly dependent on the reef matrix for shelter (Wilson et al. 2008). Thus, herbivore assemblage structure is mediated by both habitat composition and fishing intensity but links between these drivers and grazing functions are not well resolved, particularly at macroecological scales.   
 Patterns in herbivore biomass are widely used to imply changes in herbivore functioning on coral reefs (Nash et al. 2016; Robinson et al. 2018). However, biomass data overlooks size- and species-specific differences in feeding rates and roles and so measures of grazing impacts have been developed by integrating information on feeding behaviours to estimate grazing rates (Bellwood and Choat 1990; Bellwood et al. 2003). Furthermore, though allometric grazing ~ body size relationships (Lokrantz et al. 2008; Nash et al. 2013) indicate that the functional role provided by larger species is disproportionately greater (Bonaldo and Bellwood 2008), grazing potential may also depend on community size structure. Abundance decreases logarithmically with increasing body size, meaning that an assemblage of many small-bodied fish may be functionally equivalent to an assemblage of several large-bodied individuals (Munday and Jones 1998; Lokrantz et al. 2008). Indeed, size-selective exploitation which removes larger individuals (Robinson et al. 2017) and species (Taylor et al. 2014) has led to shifts in community size structure, but links between size distribution and grazing rate are unexplored.

Irrespective of body size, assemblage-level grazing rates may also depend on species composition, whereby functional impact varies according to species’ relative abundances and to interspecific variation in bite rates (Hoey and Bellwood 2008). Thus, grazing function is influenced by the species composition of the herbivore assemblage, which implies that habitat- or fishing-induced shifts in biodiversity can result in a disproportionate loss of function. Indeed, composition changes appear to underpin diversity ~ ecosystem functioning relationships on coral reefs, whereby grazing intensity is greatest in speciose grazer assemblages (Burkepile & Hay 2008, Lefcheck et al. 2019). Yet in these studies, biodiversity effects are realised at the scale of individual quadrats, for grazing pressure exerted by individual fishes. As a result, analysis of biodiversity ~ function effects at the scale of entire reefs and relevant to the characteristics of resident fish populations are lacking.

Here, we assess the drivers of herbivore functioning on coral reefs across four regions in the Indo-Pacific (Fig. S1). Our macroecological-scale analysis includes 131 reef sites, spanning a benthic gradient from coral to macroalgal dominance and a fishing gradient from open-access fisheries to no-take fishing zones and near-pristine wilderness areas. By integrating feeding observations with UVC data on grazing population sizes, we measure potential grazing levels at the scale of reef sites, which is highly relevant for understanding how benthic and fishing influences may alter ecosystem functioning (Nash et al. 2016). We ask the questions: 1) How does fishing pressure and benthic composition influence the functioning of two major functional feeding groups (croppers and scrapers)? 2) Does grazing function scale consistently with herbivore biomass? 3) Do biodiversity effects cause grazing function to decouple from grazing biomass?

**Methods**

*Survey methods*

Fish surveys were point counts of 7 m radius (Seychelles) or belt transects of 50 m length (Maldives, Chagos, GBR) conducted on hard-bottom reef slope habitat at 3-8 m depth. to minimise diver effects on fish large mobile species were censused before smaller territorial species in both point count and belt transects. In belt transects, large mobile fish were surveyed over a 5 m transect width, and small site-attached species were recorded over a 2 m transect width. For both survey types, all diurnal, non-cryptic (>8 cm) reef-associated fish were counted and sized to the nearest centimetre (total length, TL). TL measurements were calibrated by estimating the length of sections of PVC pipe and comparing it to their known length prior to data collection each day. All fish sizes (total length, cm) were then converted to body mass (grams) using published length ~ weight relationships (Froese and Pauly 2018), and standardised by survey area to give species-level biomass estimates that were comparable across datasets (kg ha-1). The UVC dataset included 101 herbivore species, with 11 species common to all four regions (Table S1). Although we combined two UVC methods to estimate fish biomass, point counts and belt transects give comparable biomass estimates (Samoilys and Carlos 2000). Observation error and bias were minimised because one observer (NAJG) performed all fish surveys.

Following fish surveys, benthic habitat composition was surveyed with eight 10 m line intercept transects (Seychelles), or 50 m point intercept (benthos recorded every 50 cm) transects (Maldives, Chagos Archipelago, and Great Barrier Reef). Taxa were grouped into broad functional groups (e.g. CCA, macroalgae, turf algae) and, if they were hard corals, identified to genus level. The structural complexity of the reef was visually estimated on a six-point scale, ranging from 0 (no vertical relief) to 5 (complex habitat with caves and overhangs) (Polunin and Roberts 1993), which correlates strongly with a range of other methods for capturing the structural complexity of coral reefs (Wilson et al. 2007). Survey methods and site descriptions for each region are described in the Supplementary Material.

*Herbivore feeding observations*

Feeding observations of Indo-Pacific herbivores provided species-level estimates on bite rates and, for scrapers, bite volumes. Surveys were conducted in the Red Sea (AH), Indonesia (AH), and GBR (AH and AGL). We analysed feeding observations for species observed in the UVC dataset (n = 39). For each observed fish, we estimated the average feeding rate (bites per minute) and, for scrapers, we estimated the average bite scar size. We define an individuals' functional impact by its feeding intensity (bite rate). By using *in situ* feeding observations that track fish forays, our approach accounts for variation in feeding frequency (the timing and distance of feeding forays) of individual fish (Nash et al. 2013).

*Ecological variable processing*

Herbivore species were categorised as croppers or scrapers according to published diet observations (Froese & Pauly 2018) and observations of feeding behaviours (Green and Bellwood 2009). Croppers were defined as species feeding primarily on the epilithial algal matrix (EAM) including detritus, turf algae, and scrapers as species feeding primarily on exposed coral substrate (Choat et al. 2002; Choat et al. 2004) (Table S1). By targeting live corals and removing larger portions of reef substrate, large ‘excavating’ scraper species also contribute to coral predation (Doropoulos et al. 2012) and bioerosion (Bellwood et al. 2011) but, because feeding observations did not record scar depth, we include these species in the scraper group and define their scraping function as potential area grazed.

We defined grazing functions separately for each functional group, and used feeding observations to convert UVC biomass estimates into the total grazing potential of croppers and scrapers. We used a Bayesian hierarchical modelling framework that estimates species- and genera-level functional rates, which allowed us to estimate grazing rates for UVC species which were not observed in feeding surveys (n = 63). Cropper function was quantified in terms of potential feeding intensity, measured as the total number of bites per minute and derived from a predictive model which accounted for species- and genera-specific bite rates (Eqs. 1,2)

 Eq. 1

 Eq. 2

We estimated the grazing rate of each cropper observed in UVCs, and used allometric relationships to convert bite rates into grams of carbon removed through EAM consumption (Marshell and Mumby 2015). Following Van Rooij et al. (1998), daily carbon intake was linked to body mass as

 Eq. 3

which we then divided by the predicted number of bites per day to produce an estimate of grams carbon consumed per minute by each cropper fish observed in UVCs. We summed estimates within each UVC replicate (i.e. point count or transect) and averaged across replicates to give site-level estimates of potential cropping function.

For scrapers, we defined the potential scraping function in terms of area of substrate removed per minute. Feeding observations provided estimates of bite rates, which we modelled as a function of body size (TL, cm) according to species- and genera-specific grazing rates, for gamma distributed errors (Eqs. 4, 5).

 Eq. 4

 Eq. 5

Scraping herbivores leave distinctive bite scars which represent the area of substrate removed in each bite. To account for potential differences in scraping action among species and across sizes, we used a second underwater feeding observation dataset of scraper bite areas. Scar area (cm2) was modelled as a function of body size (TL, cm), for Gamma distributed errors (Eqs. 6,7).

 Eq. 6

 Eq. 7

By including size (TL) as an explanatory covariate, our model accounted for scar area increasing with body size (Fig. S2A) and bite rates decreasing with body size (Fig. S2B). For each observed scraper in the UVC dataset, we estimated the expected bite rate and scar size according to its species identity and body size. Species which were not observed in feeding observations were assigned genera-level bite rates. These estimates were converted to area grazed per minute (bite rate \* scar size = area grazed) (m2 minute-1 hectare-1), summed within surveys and averaged to give site-level estimates of potential scraping function.

*Statistical modelling*

We modelled variation in herbivore functioning according to gradients in benthic habitat composition, exploitation pressure, and grazing assemblage biodiversity. Explanatory covariates were derived from fish and benthic surveys. First, to account for fishing effects ranging from the remote and protected Chagos archipelago to heavily-exploited reefs in Seychelles, we estimated total community biomass as a proxy for exploitation pressure. This proxy, hereafter fishable biomass, is highly sensitive to exploitation pressure and, in the Indian Ocean, is predicted by human population size, access to markets, and fisheries management (McClanahan et al. 2016). Reefs were also assigned a categorical fishing pressure covariate to distinguish between protected (i.e. no-take areas), exploited, and remote reefs.

Second, benthic surveys provided site-level estimates of benthic composition. We estimated the site-level cover for four major habitat-forming groups (live hard coral, macroalgae, available substrate, and rubble) and structural complexity by averaging across replicates at each site. To understand the range of benthic habitat types across the dataset, we categorised reefs according to their benthic regime, using a correlation-based PCA and K-means clustering (Jouffray et al. 2015). The optimal number of clusters was found using an elbow method with k=2-15 range, and then applied to the K-means clustering. For reefs in Seychelles which were surveyed in multiple years, we estimated regimes at each site by averaging cover values over time.

Third, we quantified compositional differences in grazing assemblages according to site-level α- and β-diversity. For α-diversity, we estimated rarefied species richness using coverage-based rarefaction curves which set estimates to the lowest sample coverage measured in the dataset (Chao and Jost 2012; Hsieh et al. 2016). For β-diversity, we estimated the local contribution to β diversity (LCBD) of each site, where higher values indicate sites which have unusually dissimilar compositions, relative to every other site (Legendre & De Cáceres 2013). By basing LCBD estimates on the full dataset, we examine how rare or endemic species might cause differences in grazing rates among reef assemblages formed from different regional pools. Prior to statistical modelling, we scaled and centered all continuous covariates to a mean of zero and standard deviation of one, and converted the categorical fishing status covariate into two dummy variables (fished - protected, fished - pristine) (Schielzeth 2010).

We used multimodel inference to assess parameter effect sizes. For each function, we fitted a global linear mixed effects model with five benthic fixed effects (hard coral, macroalgae, sand, rubble and structural complexity) and four exploitation fixed effects (fishable biomass, pristine reef, protected reef and mean size), for gamma distributed errors (). Potential covariance among reefs in the same dataset and year was modelled using nested random intercept terms where, for each observation *i* at each reef *j* in dataset *k*:

[](https://www.codecogs.com/eqnedit.php?latex=grazing_%7Bijk%7D%20%3D%20A%20%2B%20B*hardcoral_%7Bijk%7D%20%2B%20C*macroalgae_%7Bijk%7D%20%2B%20D*sand_%7Bijk%7D%20%2B%20E*rubble_%7Bijk%7D%20%2B%20F*complexity_%7Bijk%7D%20%2B%20G*fishablebiomass_%7Bijk%7D%20%2B%20H*pristine.fished_%7Bijk%7D%20%2B%20I*fished.protected_%7Bijk%7D%20%2B%20%2B%20J*mean.size_%7Bijk%7D%20%2B%20reef_j%20%2B%20dataset_k%20%2B%20%5Cepsilon_%7Bijk%7D%250) Eq. 8

From the global model, we fitted all possible subset models (Bartoń 2013) and assessed their support using Akaike’s Information Criterion (AIC), where the top-ranked model had the lowest AIC score (Burnham and Anderson 2003). We visualised relative covariate effect sizes by extracting standardised t-values for all models within 7 AIC units of the top-ranked model and, for each model, rescaling t-values so that 1 is the strongest predictor in a given model, and weighing that value by the models’ AIC weight (Cade 2015). These scaled t-values represent the relative effect size of each covariate between 0 (unimportant) and 1 (important). Next we generated model predictions to visualise the effect of each covariate with scaled t-value > 0.4, excluding remaining fixed effects and random effects and correcting predictions by each models’ AIC weight, with prediction uncertainty represented by the AIC-weighted sample variance (Robinson et al. 2017). Our multi-model approach accounts for uncertainty in the ‘best’ fitted model when AIC scores indicate several models are equally valid (Burnham and Anderson 2003). We avoid potential biases in model-averaged coefficient sizes by presenting effect sizes as standardised t-values, which are more informative measures of covariate importance than sums of AIC weights (Cade 2015).

We examined the dependency of grazing function on grazing biomass, abundance and assemblage composition. For each function, we fitted a linear mixed effects model between function ~ biomass, with random intercepts of reef nested within dataset and gamma distributed errors. Deviation from 1:1 relationships (i.e. decoupling) was evaluated with R2 values, whereby high R2 indicated a tight correlation between function and biomass/abundance and low R2 indicated decoupling of function from biomass. We further investigated decoupling by fitting a global linear mixed effects model, for each observation *i* at reef (*j*) in dataset (*k*) (nested random intercepts) and gamma distributed errors:

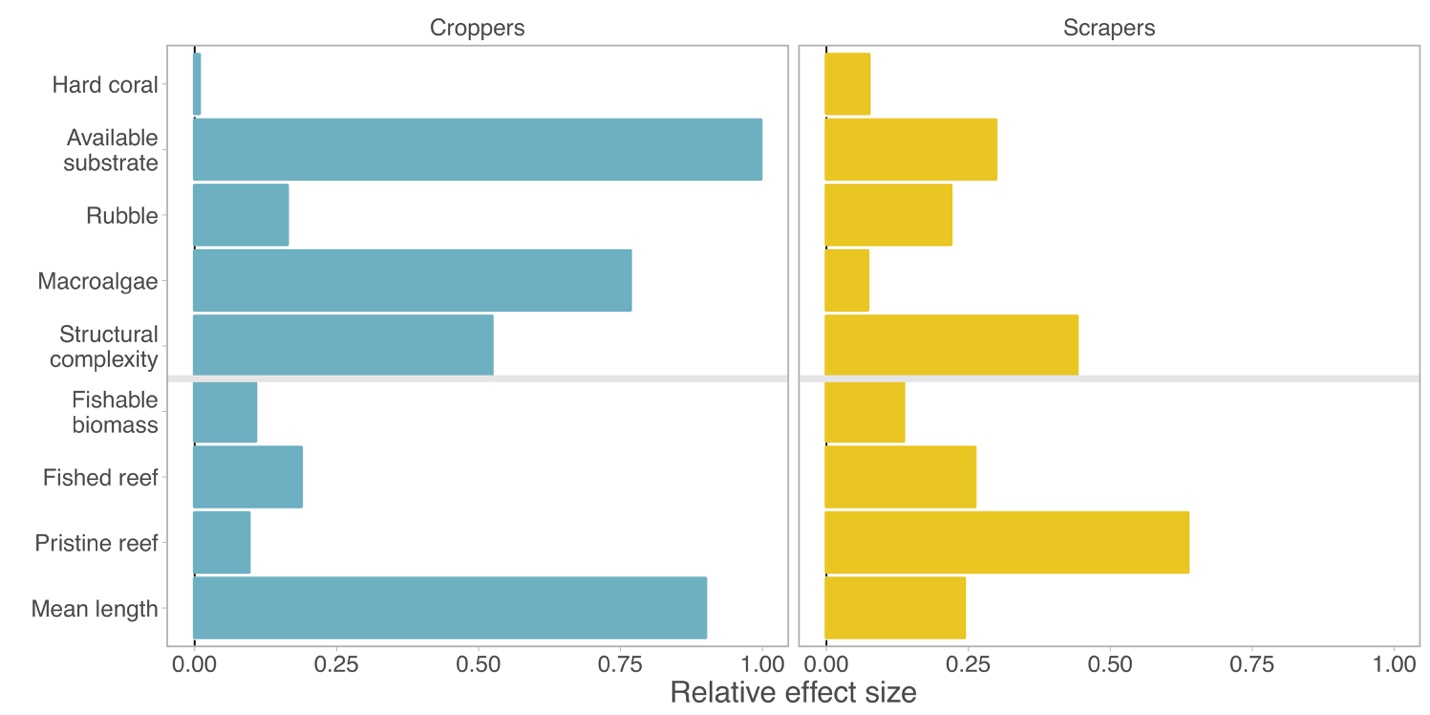
[](https://www.codecogs.com/eqnedit.php?latex=grazing_%7Bijk%7D%20%3D%20A%20%2B%20B*biomass_%7Bijk%7D%20%2B%20C*abundance_%7Bijk%7D%20%2B%20D*rarefied.richness_%7Bijk%7D%20%2B%20E*LCBD_%7Bijk%7D%20%2B%20reef_j%20%2B%20dataset_k%20%2B%20%5Cepsilon_%7Bijk%7D%250) Eq. 9

This model allowed us to assess the influence of assemblage diversity and composition on function, while accounting for biomass and abundance effects. We fitted all subset models and weighed model support with AIC, and in this analysis, the top-ranked model was > 2 AIC units from other models, and thus covariate effect sizes and model predictions were interpreted directly from that model (Burnham and Anderson 2003).

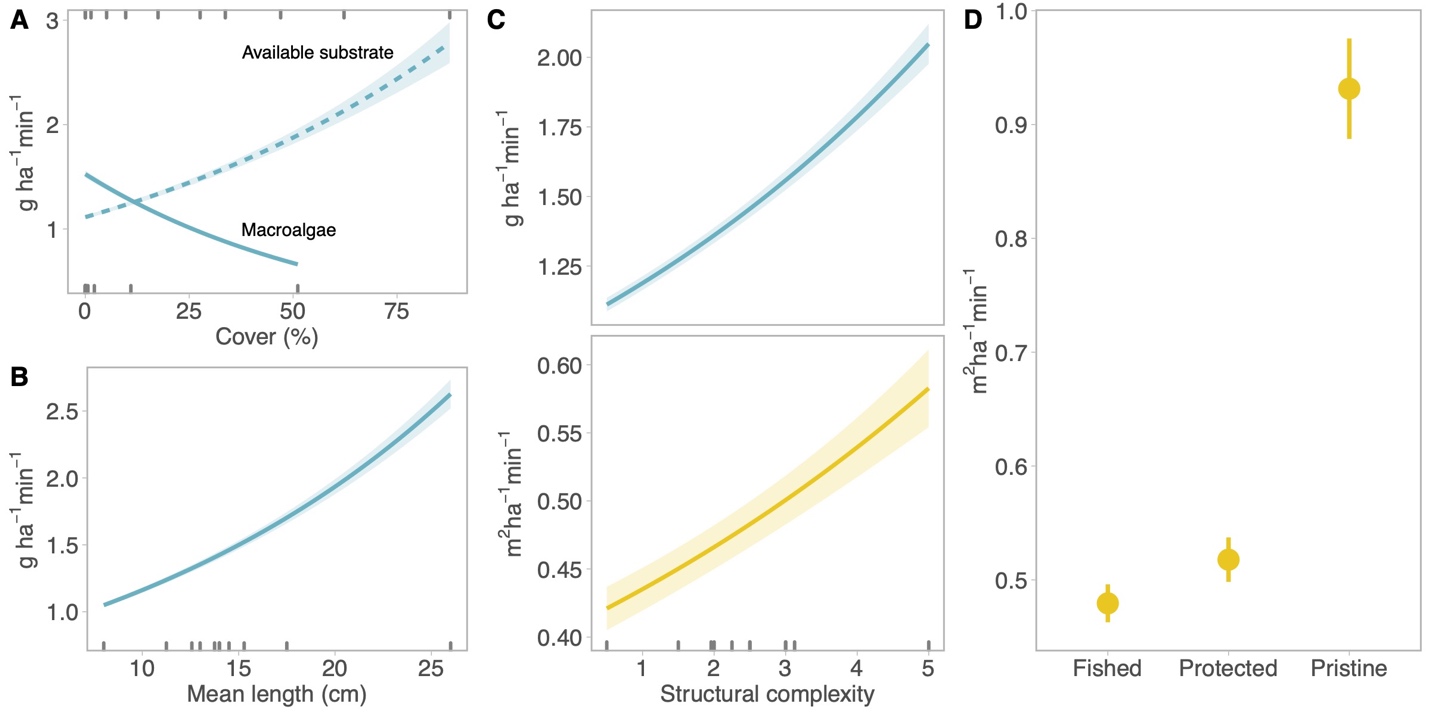
All data were analysed in R (R Core Team 2018), using packages *iNext* (rarefaction; Hsieh et al. 2016), *lme4* (linear mixed effect models; Bates et al. 2015), *MuMIn* (multimodel inference; Bartoń 2013), *rethinking* (Bayesian models; McElreath 2017), and *vegan* (diversity estimates; Oksanen et al. 2017).

**Results**

Visual census data were integrated with *in situ* feeding observations for 131 reefs in four Indo-Pacific archipelagos. For croppers, 9 species were assigned individual bite rates (32.9% of UVC biomass), and remaining species were assigned genera-specific (54.4%) or an average cropper bite rates (12.6%). Combined with biomass, bite rates corresponded with assemblage-level algal consumption rates ranging from 0.04 to 5.52 g ha-1 min-1, with grazing highest on GBR and Chagos reefs (Fig. S3A). Irrespective of region, algal consumption was maximised in complex habitats with high substrate availability and low macroalgal densities, while hard coral or rubble cover were weak influences (Fig. 1, 2A). Algal consumption rates were unaffected by fishing intensity, with remote, protected and fished reefs hosting similar cropping function potential (Fig. 1). Algal consumption did increase with average cropper size, indicating that reefs with cropper assemblages dominated by larger fishes had a higher grazing potential (Fig. 2B).

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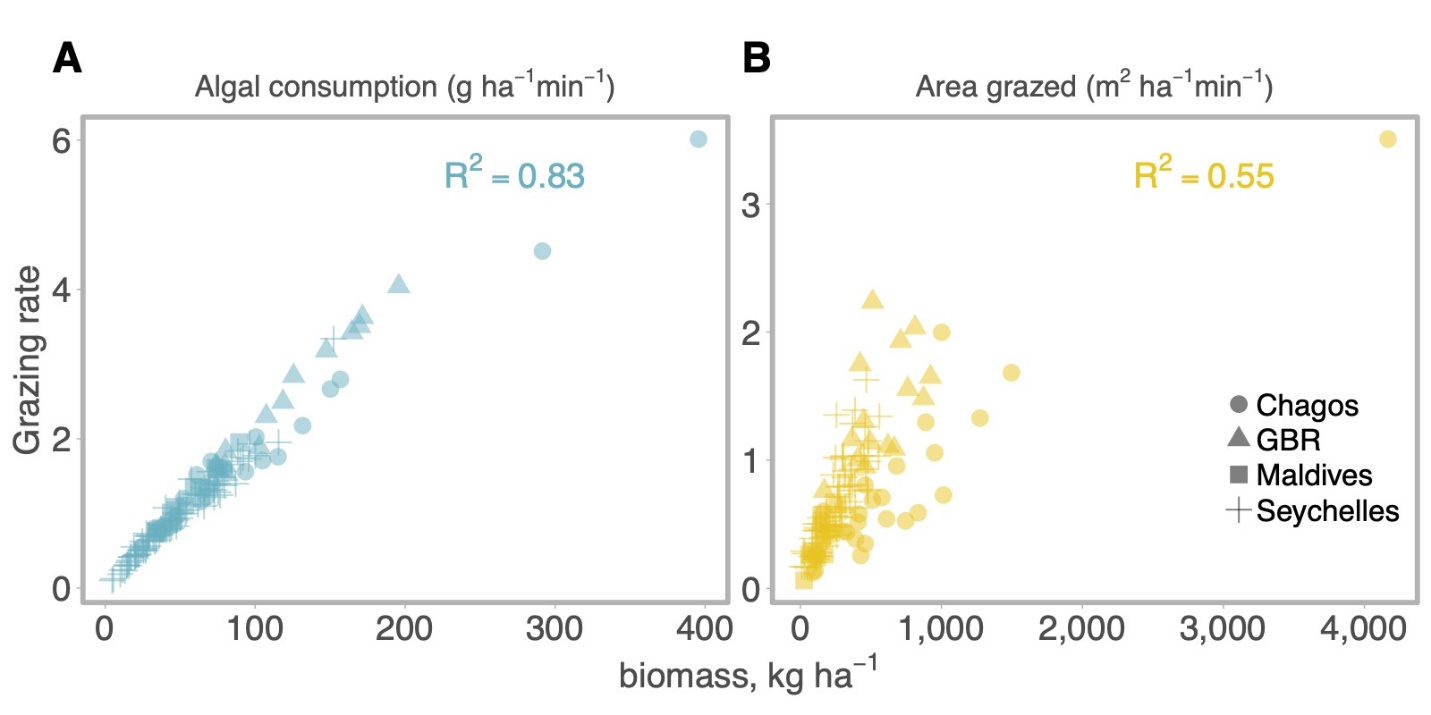
**Figure 1. Relative effect of benthic composition and fishing pressure on cropping and scraping rates.** Bars are relative effect size ratios of each covariate for top-ranking model sets (models ≤ 7 AIC units of top-ranked model), scaled to indicate very weak (0) or very important (1) drivers of grazing rates.



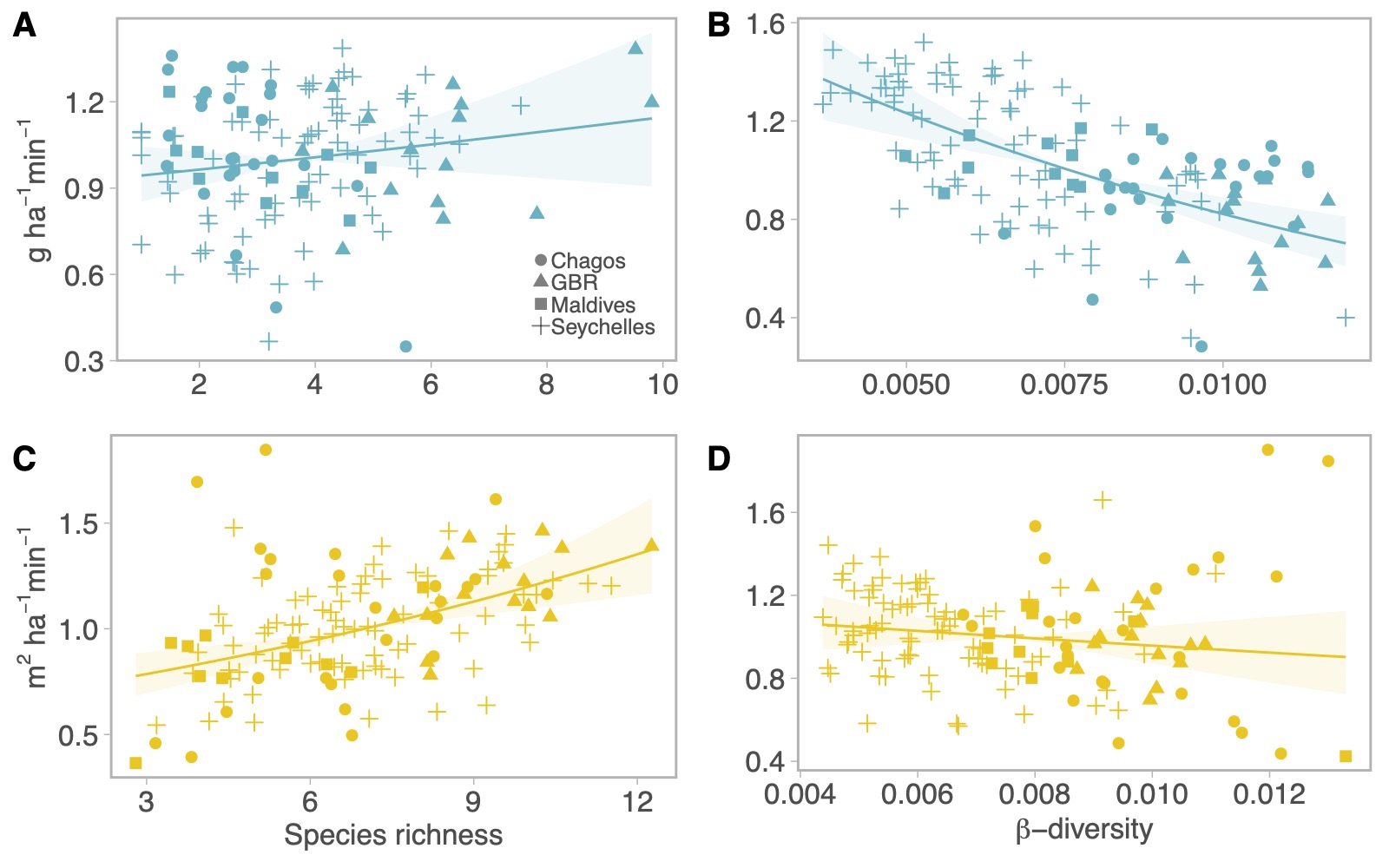
**Figure 2. Predicted effects of benthic and fishing drivers on cropper (A,B,C) and scraper (C,D) function.** Lines and points are herbivore functions as predicted by top model sets (≤ 7 AIC units from top-ranking model) holding other covariates to their means, with each model prediction weighted by its AIC weight and error represented as sample variance. All selected covariates had relative effect size ratios > 0.4 (Fig. 1). Decile rugs indicate the spread of observed data (in A, top rug is available substrate and bottom rug is macroalgae).

Feeding data were more highly resolved for scraping herbivores, with all fishes assigned size-based bite areas, and either species- (27 of 35 species, 80.9% of UVC) or genera-specific bite rates (19.1%). Potential area scraped was greatest on GBR reefs (> 1 m2 min-1 ha-1) and lowest on Maldives reefs (< 0.3 m2 min-1 ha-1) (Figure S4B). Scraping rates increased with structural complexity (Fig. 2C) but, in contrast to croppers, were relatively invariant across benthic cover covariates (Fig. 1). Remote reefs had the greatest scraping rates, which were considerably lower on fished reefs than protected ones (Figs. 1, 2D). After accounting for these coarse protection effects, scraping was only weakly associated with fishable biomass and assemblage size structure (Fig. 1).

Grazing biomass is often used as a proxy for grazing function. Here, cropping function was strongly and positively correlated with cropper biomass (R2 = 0.83, Fig. 3A), indicating that the drivers of biomass variation would match tightly to the modelled drivers of cropper function. Scraping function also increased with scraping biomass, but with greater levels of unexplained variation in area scraped (m2 min-1 ha-1) (R2 = 0.55) which occurred across the full biomass gradient and in all four regions (Fig. 3B). We found that ~~the remaining~~ variation in grazing rates was also explained by biodiversity effects. For both groups, the addition of biodiversity covariates to grazing ~ biomass models improved predictive power (Table S3), indicating that decoupling of function from biomass was partially explained by differences in the identities and relative abundance of species among grazing assemblages. For croppers, grazing rates were moderately higher in speciose assemblages, and considerably lower for assemblages with high compositional dissimilarity (Fig. 4A,B). Scraping diversity relationships followed the same direction but changed in magnitude, with stronger richness effects and weaker composition effects (Fig. 4C,D). From the least to most speciose assemblages, richness effects produced a 21% increase in algal consumption and 76% increase in area scraped. In contrast, grazing was reduced by 95% (croppers) and 17% (scrapers) at sites where compositional dissimilarity was highest.



**Figure 3. Association between grazing function and grazing biomass.** Reef-level estimates of cropper algal consumption (A) and scraper area grazed (B) plotted against UVC biomass, with shapes indicating regions and labels indicating marginal R2 from a linear model of function ~ biomass.

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**Figure 4. Biodiversity effects on decoupling of herbivore function from herbivore biomass.** Lines are predicted effects of rarefied species richness (A,C) and β-diversity (LCBD) (B,D) on the grazing rates of croppers (blue) and scrapers (yellow) after accounting for biomass and abundance, shaded with two standard errors. Points are partial residuals for different regions.

**Discussion**

Herbivore functioning varied substantially across the Indo-Pacific in accordance with top-down (i.e. fishing pressure) and bottom-up (i.e. benthic habitat) drivers which were specific to each functional group. ~~Small-bodied~~ croppers were primarily controlled by bottom-up influences, with function maximised in complex habitats with high substrate availability and low macroalgae cover. Conversely, ~~for larger~~ scraping herbivores, feeding was maximised on remote reefs in the Chagos archipelago which is isolated from fishing pressures, and was weakly associated with benthic habitat. After accounting for the strong dependency of grazing on fish biomass, we also identified biodiversity effects on grazing rates which demonstrate that variation in the number and relative abundance of species can alter grazing functions across large spatial scales.

Cropping rates were primarily mediated by benthic habitat type, in particular structural complexity, macroalgae cover, and substrate availability. Our results emphasize the strong dependence of small-bodied reef fishes on benthic composition (Munday and Jones 1998; Wilson et al. 2010), and demonstrate that top-down fishing effects do not affect cropping function, likely because croppers are small-bodied and not targeted in many reef-associated fisheries (Hicks & McClanahan 2012). Strong relationships between benthic composition and the grazing function of small-bodied reef fish may reflect the importance of food availability, which has been shown to have stronger control on cropping surgeonfishes than fishing pressure (Russ et al. 2018). For example, the decrease in function with increasing macroalgae is likely because turf algae are less accessible to croppers under macroalgal canopies (Roff et al. 2015) whereas, on reefs with high substrate availability and limited macroalgae, expansive and easily accessible turf mats tend to support large grazer populations (Williams & Polunin 2001). Strong benthic effects imply that cropper functioning will respond more strongly to habitat disturbances, such as coral bleaching, habitat destruction or enrichment of algal communities, than to fishing. For example, disturbances which increase substrate availability for turf algal growth, such as coral mortality from heat stress (Gilmour et al. 2013), might therefore be expected to stimulate an increase in cropping function. However, since structural complexity was also shown to be a strong driver, any positive rebound of cropping function may be negated if disturbances also erode structural complexity (Graham et al. 2006).

Scraping function was strongly influenced by fishing pressure, which suppressed grazing rates far below those supported at remote wilderness reefs in the Chagos Archipelago. Our results further indicate that exploitation of large-bodied scrapers has compromised scraping functions on coral reefs (Bellwood et al. 2011). This effect superseded influences of benthic cover and small-scale fishing protection, suggesting that bottom-up control of scraping assemblages on reefs leads to relatively minimal variation in their function, and that small-scale fishing protection does not conserve wilderness levels of scraping function. Movement of fish across reserve boundaries (Green et al. 2014) and low protection compliance (Bergseth et al. 2018) likely limited the effectiveness of these small MPAs, many of which are adjacent to fishing grounds. Despite weak benthic cover effects, scraping rates increased moderately with structural complexity, further underlining the importance of coral reef structure in supporting herbivory (Nash et al. 2016).

Although biomass was the strongest predictor of grazing function for both functional groups, unexplained variation in function ~ biomass relationships was partially attributable to biodiversity effects. Positive effects of species richness on grazing rates, particularly for scrapers, are broadly consistent with recent coral reef studies which have uncovered positive biodiversity effects on herbivory (Lefcheck et al. 2019, Topor et al. 2019). The mechanisms underlying our results are, however, entirely different to those in small-scale experiments where biodiversity leads to feeding complementarity (Burkepile and Hay 2008, 2011) and intensifies grazing of individual fishes (Lefcheck et al. 2019, Topor et al. 2019). By focusing on benthic plots, these studies do not necessarily extrapolate to entire reefscapes where grazing pressure is dependent on the size of the resident fish assemblage, which itself is controlled by the availability of benthic habitat and historic fishing levels. Therefore, our approach of integrating feeding rates with UVC data enabled us to generate reef-level estimates of potential grazing pressure across a gradient of grazing biomass. At this scale, we confirm that more diverse reefs have higher potential grazing pressure. Here, however, β patterns contradict Lefcheck et al.’s (2019) finding that high species turnover raised grazing rates, likely because our LCBD estimates assessed turnover of species among distinct regional pools rather than among connected habitats. Across regional pools, we suggest that assemblages dominated by widely-distributed species have a lower grazing potential than those dominated by endemic species.

Biodiversity effects partially explained why grazing function decoupled from grazing biomass. Decoupling was strongest in scrapers, likely because all fishes were assigned species-, genera- and size-specific bite rates (Lokrantz et al. 2008) and so scraping estimates were more sensitive to changes in species diversity. In contrast, croppers were more tightly coupled to biomass levels, due to a lack of bite size data and to the high proportion of individual fishes which were assigned average grazing rates. Indeed, we note that our definitions of grazing functions were limited by our generalisation across species with similar functions but different feeding modes. This may have been particularly problematic for cropping species which have well-documented differences in morphology, diet (e.g. detritivores or turf), and feeding behaviours (Choat et al. 2002, Brandl et al. 2015, Tebbet et al. 2017).

The modelling framework we used to generate grazing estimates is a significant improvement on the procedure employed by previous macroscale grazing studies (e.g. Bellwood et al. 2011). By modelling genera- and species-specific bite rates from observations collected in several regions, we were able to leverage observational data in a hierarchical framework which predicts grazing rates of new, related species, given uncertainties in species, genera and body sizes. For example, we were able to assign bite rates to species observed in UVC but not observed in feeding surveys, with estimates that were informed by the feeding behaviour of closely related congeners. Such models could be further improved with additional feeding data on other herbivore species in different regions, and could even be developed to account for temperature effects on grazing rates (Bruno et al. 2015) that might confound comparisons of herbivory across temperature regimes.

The random intercepts in our predictive models indicated that regional similarities in grazing rates were unexplained by benthic, fishing and biodiversity covariates, which is likely due to unmeasured processes that control herbivore biomass. For example, herbivore biomass variation (and thus grazing function) has been linked to differences in oceanic productivity (Heenan et al. 2016) while video observations indicate that grazing intensity is constrained by wave exposure (Bejarano et al. 2017). Similarly, long-term studies of frequently perturbed coral reefs indicate that grazing assemblages continually reorganize in response to disturbance (Han et al. 2016), implying that grazing intensity will respond in similar non-linear trajectories. Temporal analyses linking habitat suitability, primary productivity, and herbivory would greatly develop our understanding of how herbivory rates influence long-term changes in reef state.

Our study demonstrates how benthic habitat, fishing pressure and biodiversity influence the functional potential of herbivore assemblages at scales which are relevant for understanding ecosystem-level responses to disturbances such as bleaching and management of the marine environment (Nash et al. 2016). Cropping pressure is likely to increase in response to stressors which clear substrate space for turf growth. Intact reef structure will be critical for maintenance of scraping functions, though reefs in close proximity to human populations are unlikely to return to wilderness levels of grazing pressure, even with protection from fishing. For a given level of biomass, protection of biodiversity will enhance grazing, but differences in regional pools mean that grazing potential of fish assemblages may vary naturally among reefs. We stress that biomass was by far the most important predictor of scraping function, and recovery or protection of fish biomass will help ensure herbivory processes are functionally intact on degraded coral reefs (Williams et al. 2016).

**Acknowledgements**

Data? Grants?

[**Supplementary Material**](https://github.com/jpwrobinson/grazing-gradients/blob/master/writing/ms/supp-material.pdf)

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