

The importance of sponges and mangroves in supporting fish communities on degraded coral reefs in Caribbean Panama

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Fish communities associated with coral reefs worldwide are threatened by habitat degradation and overexploitation. We assessed fish communities on coral reefs, mangroves fringes, and seagrass meadows on the Caribbean coast of Panama to explore the influences of habitat cover, connectivity and environmental characteristics in sustaining biomass, richness and trophic structure in a degraded tropical ecosystem. We found 94 % of all fish across all habitat types were of small body size (≤ 10 cm), with communities dominated by fishes that usually live in habitats of low complexity, such as Pomacentridae (damselfishes) and Gobiidae (gobies). Total fish biomass was very low, with small fishes from low trophic levels over-represented, and top predators under-represented, compared to coral reefs elsewhere in the Caribbean. For example, grazing and omnivorous fishes (trophic level 2 – 2.7) comprised 37 % of total fish biomass, and the small parrotfish *Scarus iseri* comprised 72 % of the parrotfish biomass. For the reef-associated fish community, we found evidence that biogenic habitats provided a mechanism of resilience. In particular, the abundance of sponges on a given reef and proximity of mangroves were found to be important positive drivers of reef fish richness, biomass, abundance and trophic structure. Our study indicates that a diverse fish community can persist on degraded coral reefs, and that the availability and arrangement within the seascape of habitat-forming organisms additional to corals, including sponges and mangroves, is critical to the maintenance of functional processes in such ecosystems.

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12 Abstract

13 Fish communities associated with coral reefs worldwide are threatened by habitat degradation
 14 and overexploitation. We assessed coral reefs, mangroves fringes, and seagrass meadows on the
 15 Caribbean coast of Panama to explore the influences of habitat cover, connectivity and
 16 environmental characteristics in sustaining biomass, richness and trophic structure of fish
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 19 habitats of low complexity, such as Pomacentridae (damselfishes) and Gobiidae (gobies). Total
 20 fish biomass was very low, with small fishes from low trophic levels over-represented, and top
 21 predators under-represented, compared to coral reefs elsewhere in the Caribbean. For example,
 22 herbivorous and omnivorous fishes (trophic level 2 – 2.7) comprised 37 % of total fish biomass,
 23 and the small parrotfish *Scarus iseri* comprised 72 % of the parrotfish biomass. For the reef-
 24 associated fish community, we found evidence that biogenic habitats provided a mechanism of
 25 resilience. In particular, the abundance of sponges on a given reef and proximity of mangroves
 26 were found to be important positive drivers of reef fish richness, biomass, abundance and trophic
 27 structure. Our study indicates that a diverse fish community can persist on degraded coral reefs,
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 30 ecosystems.

31 Introduction

32 Coral reef fishes are useful model communities for exploring drivers of diversity at landscape
 33 and regional scales (Galzin et al. 1994; Fabricius et al. 2005; Knowlton et al. 2010; Wilson et al.
 34 2010). They are sensitive to changes in habitat and anthropogenic impacts – a particular concern
 35 given that fishes play an important role in coral reef ecosystems, and declines of coral reef fishes
 36 threaten people’s livelihoods and food security (Cesar 2000; Cesar et al. 2003; Bellwood et al.
 37 2004; Paddack et al. 2009). A variety of human impacts are responsible for coastal degradation,
 38 including habitat destruction, eutrophication, and sedimentation (Hughes 1994; Jackson et al.
 39 2001; Aronson et al. 2003). Climate change has additionally contributed to ecosystem decline
 40 through coral die-off from bleaching, hypoxia events and storms (Beukers and Jones 1998;
 41 Wilson ; Alvarez-Filip et al. 2009; Wilson et al. 2010; Altieri et al. 2017). The consequence of
 42 these processes is structural collapses and habitat homogenization in coral reefs which have a
 43 variety of potential direct and indirect implications for the organisms that reside there (Bell and
 44 Galzin 1984; Jackson et al. 2001; Kuffner et al. 2007; Wilson et al. 2010; Alevizon and Porter
 45 2015; Mora 2015).

46

47 Additional factors contributing to declining reef fish abundances are unsustainable fisheries and
 48 increasing demand for fish products for a growing human population (Hodgson 1999; Jackson et
 49 al. 2001; Zaneveld et al. 2016). The overexploitation and disproportionate targeting of large size
 50 classes and high trophic levels affects fish population structure, growth, and reproduction, and
 51 contributes to a trophic imbalance and shifts in trait composition in the reef fish community
 52 (Pauly et al. 1998; Mumby et al. 2006). This, in turn has led to further changes in habitat

structure, phase shifts from coral to algal communities, and decreasing ecosystem stability (Saila et al. 1993; Jennings and Lock 1996; White and Jentsch 2001).

Reef fish populations have also been negatively affected by the loss of coastal habitats as nurseries (Nagelkerken et al. 2000). The nursery-role concept suggests that many reef fishes (e.g., families Lutjanidae, snappers; Serranidae, groupers; Haemulidae, grunts) have life cycles that include seagrass meadows and mangroves as nursery and feeding grounds (Beck et al. 2001; Nagelkerken et al. 2002; Unsworth et al. 2008; Ley 2014; Serafy et al. 2015). Seagrass meadows and mangrove forests have high primary and secondary productivity relative to unvegetated substrates, and support high diversity and abundances of reef fishes (Nagelkerken et al. 2000; Beck et al. 2001; Mumby et al. 2004). Many fish species on coral reefs therefore depend on the connectivity to, and integrity of, associated habitats.

Our study region, the Caribbean Sea, has experienced declining reef fish populations as a result of pollution, ecosystem degradation and unsustainable reef fisheries (Hughes 1994; Gardner et al. 2003; Bellwood et al. 2004; Paddock et al. 2009). These problems appear particularly prominent at our focal study area in Bocas del Toro on the Caribbean coast of Panama, where rapid human population growth connected with agriculture (banana industry) and tourism has accelerated the decline of water quality, physical destruction of reefs and fishing pressure (Guzmán and Jiménez 1992; Collin 2005; D’Croz et al. 2005; Cramer 2013; Aronson et al. 2014; Seemann et al. 2014). Bocas del Toro encompasses a coastal coral reef-seagrass-mangrove system in a semi-enclosed lagoon. It is composed of six major islands and the mainland, which surrounds the Almirante Bay, and includes mangroves fringing the mainland and mangrove

islands scattered across the bay (Collin 2005; Guzmán et al. 2005). Reefs are typically dominated by corals with a high stress tolerance, including *Porites furcata* in shallow (1-4 m) and *Agaricia* spp. (>3 m) in the deeper areas (Seemann 2013; Aronson et al. 2014; Seemann et al. 2014). Associated seagrass meadows are dominated by *Thalassia testudinum* (turtlegrass). Mangrove fringes are comprised of *Rhizophora mangle* (red mangrove). Several rivers, creeks and oceanic inlets discharge sediments and nutrients into the bay (Beulig 1999; Collin 2005). Bleaching and low oxygen events occur regularly due to lagoonal characteristics including retention of warm water and depletion of oxygen (Kaufmann and Thompson 2005; Seemann et al. 2014; Altieri et al. 2017). Bocas del Toro reefs potentially represent a model system for improving predictions relevant throughout the region due to their exposure to common stressors, such as high terrigenous run off, nutrient levels, and overfishing, that are afflicting other coral reefs in the Caribbean (Riegl et al. 2009; Sammarco and Strychar 2009; Leinfelder et al. 2012; Aronson et al. 2014).

This study aims to characterize the ecosystem attributes that facilitate the maintenance of essential functions, biodiversity and biomass of coral reef fish communities in a degraded ecosystem. Specifically, we (1) quantify the fish community at 67 sites in 5 bioregions of the Caribbean to assess the status of our focal study system in Bocas del Toro along a gradient of ecosystem degradation and over-fishing, (2) examine the effects of proximity of mangroves and seagrass for fish communities on coral reefs, and (3) identify characteristics of coral reef habitat that are positively related to biomass, abundance and structure of the fish community. Addressing these objectives contributes to a better understanding of how landscape-scale features underlie the resilience of degraded coastal habitats.

99

100 **Methods**

101 **Study system**

102 To place results within the wider regional context, we conducted fish surveys at reefs with
103 different fishery management restrictions in different Caribbean ecoregions.

104 In our focal study areas of Bocas del Toro on the Caribbean coast of Panama (Fig. 1), we also
105 conducted comprehensive surveys of fish communities, benthic surveys, and water quality
106 measurements in adjacent seagrass and mangrove fringe areas. All data from Bocas del Toro
107 were collected from May to July 2015. Data for the other Caribbean regions were collected from
108 2012 to 2015.

109 Research was conducted under a Scientific Permit from the Ministry of the Environment Panama
110 (MiAmbiente) and Autoridad de los Recursos Acuáticos de Panamá (ARAP) with the Number:
111 SE/APO-1-15 & 10b.

112

113 **Caribbean Data Set**

114 **Fish surveys.** We conducted visual fish surveys using the Reef Life Survey (RLS) method 1
115 protocol (Edgar and Stuart-Smith 2014) at reefs at 67 sites in the following five ecoregions
116 (Spalding et al. 2007): Southern Caribbean (14 sites, Bonaire), Southwestern Caribbean (31 sites,
117 Bocas del Toro, Kuna Yala, Archipelago of San Andres), Greater Antilles (1 site, Grand
118 Cayman), Floridian (17 sites, Florida Keys) and Bahamian (4 sites, Turks and Caicos Islands).
119 Surveys involved underwater visual censuses by scuba divers at reef sites (each with 2-6
120 replicate transects) in depths of 1 to 35 m. Divers counted and assigned all fish species observed
121 within binned size-classes along a 50 x 5 m belt transect (250 m²). All fishes sighted on each

transect were recorded on a waterproof datasheet as the diver swam along the transect at approximately 2 m min⁻¹. We identified fish species to the highest taxonomic resolution possible, and estimated body length for each individual. The order of priority for recording accurately was to first ensure all species observed along transects were included, then individuals of larger or rare species were accurately counted, then estimates made of abundances of common species. If an individual could not be identified underwater, a photograph was taken for later identification. Abundance, size and species identity were used to estimate biomass in kg ha⁻¹ using conversion factors provided by Fishbase (www.fishbase.com), as described by Edgar and Stuart-Smith (2014).

Bocas del Toro Data Set

Fish surveys. The RLS method employed on coral reefs in five ecoregions was also applied in seagrass and mangrove habitats located within 250 m of reef sites in Bocas del Toro. Seagrass sites ranged in depth from 1 m to 4 m, whereas mangrove fringe root systems had maximal depth of 2 m. Mangrove surveys were conducted amongst the mangroves prop roots below the upper intertidal fringe with counts and size estimates made for all fishes in a 5 m wide belt within the mangrove root system. Two 50 m transects were laid end-to-end along the mangrove fringe given that side-by-side replicate transects typical of the RLS protocol could not be applied within mangrove root habitats.

Habitat assessment. We conducted benthic surveys to characterize coral reef and seagrass bed habitats. Reef and seagrass benthos were analyzed with 20 photo quadrats (0.5 m²), which were taken every 2.5 m along the 50 m long transects at each site. Photos were analyzed via point counting using the Coralnet annotation tool (coralnet.ucsd.edu). A total of 25 points were

randomly distributed on each photo and categorized. Substratum categories for analyses were: healthy hard coral, bleached hard coral, recently-dead coral, anemones, non-calcifying corals (including hexacorals and octocorals), sponges, sessile worms (tube worms, mostly polychaetes), zoanthids, rubble, sand, rock, calcifying algae, seagrass and macroalgae. If sessile organisms were too small for identification or obscured by dark shadows, then they were excluded from the dataset. In addition, the distance between reef sites surveyed and nearest mangrove was measured using GPS coordinates (table 1).

Water quality measurements. Water quality was assessed by quantifying temperature ($^{\circ}\text{C}$), salinity (psu), water depth (m), total dissolved solids (TDS, mg L^{-1}), dissolved oxygen (mg L^{-1}), pH, turbidity (FNU), chlorophyll ($\mu\text{g L}^{-1}$), blue-green algae concentrations ($\mu\text{g L}^{-1}$), and dissolved organic matter (fDOM, RFU) with an Exo2 multiparameter sonde (YSI, Xylem brand) (Snazelle 2015). The sonde was positioned ~ 10 cm above the bottom in each habitat (reef, seagrass and mangrove fringe). Measurements were recorded at intervals of 1 – 6 min over a time period of at least 30 min during the fish surveys, and constrained to the mid-day hours between, hence measurements were subject to the daily variability of weather conditions or tidal cycles.

Data analyses.

The Caribbean reef fish data set was used to characterize the fish community in relation to the protection status of the sites. All 67 sites from the five different ecoregions were individually classed by management type using the criteria of Edgar et al. (2014): NTZ (no take zones, $n=27$), RZ (restricted zones that allow local fishing within an MPA, $n=19$) and OZ (open zones where

167 fishing is unrestricted, $n=12$). These data were compared to data from Bocas del Toro (OZ, $n=9$).
 168 Replicated surveys from each site were averaged.
 169
 170 Data from the fish surveys were used to calculate fish community metrics, including total
 171 abundance (density), abundance of major fish families within size bins (≤ 10 cm; $>10 - 20$ cm;
 172 >20 cm), total biomass, biomass of fishes ≤ 10 cm, and total species richness. We also calculated
 173 the mean trophic level as an abundance weighted mean of the reef fish community by
 174 multiplying the trophic level of each species by their log abundance, summing these values
 175 across species recorded on a transect, and dividing by the total log abundance of all fishes on the
 176 transect. The classification of the trophic level (2 – 5) for each species was based on feeding
 177 strategy: herbivores and detritivores (2 – 2.1), omnivores (2.2 – 2.7), low-level carnivores (2.8 –
 178 3.4), mid-level carnivores (3.5 – 3.9) and high-level carnivores (4 – 4.5) (classification and
 179 values obtained from Fishbase; www.fishbase.org). We also compared preferred substrate types
 180 and resilience factors (values obtained from Fishbase) of the fish species, the latter estimated
 181 from population doubling time (low, medium, high). Fish community metrics were averaged
 182 across sites within a sites and compared among regions for significant differences using one-way
 183 ANOVA or a Student's *t-test*.
 184
 185 For the Bocas del Toro dataset only, we assessed whether mangroves and seagrasses provided
 186 juvenile or alternative habitat to coral reefs by comparing the abundance (log transformed) and
 187 composition of fishes in the different habitat types. We assumed that higher abundances of fishes
 188 amongst mangroves and seagrasses compared to reefs, and high species similarity, indicates high
 189 migration and exchange rates, indicates higher likelihood and magnitude of migration and

exchange rates. We excluded small-bodied species (maximum total length ≤ 12.5 cm), which are presumably non-migratory fish species (Dahlgren et al. 2006), such as *Apogon townsendi* (belted cardinalfish), *Canthigaster rostrata* (caribbean sharpnose-puffer) and *Coryphopterus* spp. A principal component analysis (PCA) on correlations (fish abundance log transformed, only fish > 12.5 cm) was used to compare differences and to define distances in the fish communities between reefs at different distances to mangroves and mangroves and seagrass.

We also tested for correlations between environmental factors and the reef fish community metrics across all sites. Environmental factors included reef cover, cover of the seagrass benthos, distance to mangroves and water quality parameters. Fish metrics included biodiversity, fish traits, biomass, size structure and the abundances of individual fish species. Data were characterized using a scatterplot matrix (see appendix) and nonparametric Spearman's tests for pairwise correlation probabilities. For all statistical analyses, fish abundance data were log-transformed to down-weight the extremely high abundance of a few fish species (Edgar et al. 2014). All statistical analyses were conducted using JMP Software 13.01.

Results

Characteristics of the fish community.

We recorded a total of 77 fish species across all habitats in Bocas del Toro, of which 61 species were found on coral reefs. The average mean richness per transect was 29 ± 7 (SD). This value was low compared to our other Caribbean survey sites which had a mean richness per transect of 52 ± 4 species (with a cumulative total of 196 species recorded in the whole Caribbean) (Stuart-Smith et al. 2013; Edgar and Stuart-Smith 2014). Fish biomass on Bocas del Toro reefs (71 ± 63

kg ha⁻¹) was also lower than on other Caribbean reefs, in both no-take zones and MPAs with restricted fishing (ANOVA, $P=0.02$ and 0.001 , respectively), although the difference was not significantly lower for open zones ($P>0.05$) (Fig. 2a). Moreover, the range of total observed fish biomass in Bocas del Toro ($30 - 1350$ kg ha⁻¹) represents the lowest numbers found amongst fish surveys conducted in the Caribbean, which were $140 - 5930$ kg ha⁻¹ elsewhere.

The biomass of herbivorous, omnivorous and detritivorous fishes in Bocas del Toro (trophic level $2 - 2.7$) was 37% of the total biomass and 76% of all individual fishes counted (Fig. 2b). Herbivores alone comprised $27\% \pm 3.5\%$ (SD) of biomass versus $10\% \pm 4\%$ across the wider Caribbean. Pomacentridae (damselfishes) and Scarinae (parrotfishes) were the predominant taxa in terms of biomass. *Scarus iseri* (striped parrotfish) contributed 72% of the parrotfish biomass. High-level carnivores contributed $22\% \pm 3.5\%$ of total fish biomass, versus $31 \pm 4\%$ elsewhere in the Caribbean. Dominant high-level carnivores in Bocas del Toro were *Carangoides ruber* (bar jack), *Cephalopholis cruentata* (graysby), *Hyplopectrus nigricans* (black hamlet) and *Scomberomorus regalis* (cero). There was a trend for fish communities in Bocas del Toro to exhibit a greater proportion of total biomass comprised of herbivores, omnivores and detritivores (trophic level: $2 - 2.7$), and a lower proportion comprised of high-level carnivores (trophic level: $4 - 4.5$), relative to other Caribbean reefs, although the difference was not significantly different for either group (Fig. 2b).

A total of 94% of all fishes observed across all habitat types (reef, seagrass, mangrove) in Bocas del Toro were in the smallest size class (≤ 10 cm length). Fishes ≤ 10 cm represented 59% of the total biomass within the reefs. The abundance of fishes within the smallest size class (≤ 10 cm)

was significantly higher in Bocas del Toro than other Caribbean reefs (ANOVA, $P < 0.0001$), whereas the abundances of medium- ($>10 - 20$ cm) and large- (≥ 20 cm) sized fishes were significantly lower (ANOVA, $P < 0.0001$) (Fig. 2c). This pattern was also evident when comparing reef fish families in Bocas del Toro with other Caribbean reefs (table 2).

Relationships between environmental factors and fish community composition

Some environmental parameter and habitat factors were associated with reef fish community metrics in Bocas del Toro. Multiple environmental factors were not independent, as sponge cover was negatively correlated with the distance to mangroves and also positively to chl *a* ($R^2 = 0.60$ and $R^2 = 0.70$, respectively, $P < 0.01$). The other water parameters were not found to correlate with any fish community or species metrics. Sponge cover was the strongest positive correlate among all environmental parameters for species richness ($R^2 = 0.5$, $P < 0.01$), small fish ≤ 10 cm biomass ($R^2 = 0.85$, $P < 0.01$), and trophic level of the fish community ($R^2 = 0.89$, $P < 0.01$). The abundance of *Abudefduf saxatilis* (sergeant major) was significantly correlated with sponge cover ($R^2 = 0.62$, $p = 0.0027$). Survey sites characterized by high sponge cover and low distance to mangroves were characterized by fishes such as *Abudefduf saxatilis* (sergeant major), *Hypoplectrus nigricans* (black hamlet), *Coryphopterus personatus* (masked goby) and *Coryphopterus glaucofraenum* (bridled goby). *Scarus iseri* (striped parrotfish), *Stegastes partitus* (bicolor damselfish) and *Cephalopholis cruentatus* (graysby) had a positive association with recently dead corals, however, the cover of dead corals was negatively correlated with the abundance of most fish species.

Fish richness on a given reef was positively correlated with fish richness in nearby mangroves ($R^2=0.76$, $P<0.05$). The three reef sites with low sponge cover and without mangroves in close proximity (Salt Creek, Popa, Hospital Point) showed lower biomass, abundances and species richness of fishes (table 1, Fig.3 and Fig. 4). The site without either mangroves or seagrass nearby (Hospital Point) showed the lowest species richness.

Also, distance to mangroves was identified as a factor to influence reef fish communities (Fig. 5), suggesting that mangrove distance has a strong influence on reef community types, likely by mangroves functioning as effective nursery grounds and as alternative complex habitats. The proportion of carnivorous fishes was significantly higher at the sites closer to the mangroves than those sites that were further away (ANOVA, $P<0.01$, Fig. 4). However, a more detailed look at the carnivorous fishes revealed that the sites at an intermediate distance from mangroves (STRI, Juan Point, Coral Cay) possessed a significantly higher proportion of top-level carnivores than sites that were closer or further away (Fig. 4, ANOVA, $P<0.01$).

The highest abundances of all fish observed were recorded for families Pomacentridae (damselfishes) and Gobiidae (gobies) (table 2). However, Gobiidae were only abundant at sites close to mangroves. *Coryphopterus personatus* (masked goby) dominated these sites, with abundances up to 13 individuals m^{-2} . RLS surveys conducted elsewhere in the Caribbean (e.g. San Andres Archipelago, 350 km distant) revealed much lower densities for the same species (0.2 individuals m^{-2}).

Generally, fishes with life cycles closely associated with hard corals (Lewis 1997), such as Pomacanthidae (angelfishes), were present in very low numbers on the reefs of Bocas del Toro (<1 per transect). Other reef fishes typically associated with hard substrates with a high complexity such as Balistidae (triggerfishes), Apogonidae (cardinalfishes), Muraenidae (moray eels), Sciaenidae (drums), Pseudochromidae (dottybacks) and Serranidae (grouper) were scarce within the bay (<1 per transect). Many fish species of low or very low resilience factors, including those at higher trophic levels, such as *Diodon hystrix* (porcupinefish), *Ginglymostoma cirratum* (nurse shark), *Gymnothorax funebris* (moray eel), *Lutjanus jocu* (dog snapper), *Ocyurus chrysurus* (yellowtail snapper), *Pomacanthus arcuatus* (gray angelfish) were only observed on reefs with mangroves in close proximity (≤ 250 m distance).

Discussion

Our surveys revealed that the fish fauna in Bocas del Toro is depauperate in richness and biomass by Caribbean standards. We found evidence that the fish community is representative of a degraded and overexploited ecosystem, characterized by numerical dominance of fishes that are small bodied and also typical of habitats of low complexity, such as Pomacentridae and Gobiidae, with few representatives of fish families that achieve body sizes targeted by fisheries or that are commonly associated with high-relief coral reefs. Nevertheless, sponge cover and proximity to mangroves were found to be positively correlated with fish species richness, biomass, abundance and trophic level. This suggests that sponges as habitat-forming reef organisms, and mangroves as nursery grounds and alternative habitats, continue to provide critical habitats for the reef fish communities in a degraded ecosystem, and may be an important mechanism of resilience where they are able to counteract some effects of degradation.

303

304 Some fishes appeared to be an indicator species for the overall trends observed at our study site.

305 One example is the goby *Coryphopterus personatus*, which forms schools that hover in a

306 vulnerable position above the bottom in extremely high abundances (65-fold higher abundances

307 than in the San Andres Archipelago). Moreover, fish surveys in our Bocas del Toro study area in

308 2002 revealed densities an order of magnitude lower at 1.2 individuals m⁻² (Dominici-

309 Arosemena and Wolff 2005). We suggest this species represents an indicator species for

310 overfished reefs that benefits from loss of predatory fishes that historically kept limited their

311 densities. Another example of an indicator is *Scarus iseri*, which is perhaps an important habitat

312 interactor, given its role as the predominant herbivorous fish in Bocas del Toro (Kuempel and

313 Altieri 2017). It likely plays an important role supporting the growth of sponges and corals by

314 cropping competing macroalgae. Third, *Abudefduf saxatilis* was identified as an indicator for

315 sponge cover which in turn is a factor positively correlated to fish richness, biomass, abundance

316 and relatively high mean community trophic levels.

317

318 A degraded reef fish community in Bocas del Toro is evidenced by low total biomass, under-

319 representation of biomass at high trophic levels, and high abundance of small fishes; all classic

320 symptoms of over-fishing (Pauly et al. 1998; Myers and Worm 2003). Moreover, the range of

321 total observed fish biomass represents the lowest numbers found amongst fish surveys conducted

322 in the Caribbean. High level carnivores and large fishes are depleted in intense fisheries (Cinner

323 and McClanahan 2006; Wilson et al. 2010), causing a skewing of the trophic food web and

324 community size structures. As described in Wilson et al. (2010), the loss of individuals within

325 the largest size classes, which have the highest per capita reproductive output and produce the

majority of juveniles, impacts the recruitment of small size classes containing juveniles in the reef fish population. Accordingly, we observed that small Haemulidae were rare on Bocas del Toro reefs. Exploitation thus appears to have contributed substantially to the distorted fish community patterns observed at Bocas del Toro (Guzmán et al. 2005; Cramer 2013).

Another plausible hypothesis for the low total fish biomass and trophic shifts within the fish community in Bocas del Toro relative to other Caribbean sites is the loss of hard corals (Turner et al. 1999; Wilson et al. 2010). This in turn results in the loss of shelter and feeding grounds (Turner et al. 1999; Alevizon and Porter 2015). This hypothesis was supported by significant negative correlations between the proportions of recently-dead corals and the biomass of fishes, as well as the finding that fish species that are known to associate with hard corals or hard substrate were rare. Instead, fishes known to live on habitats of low complexity (particularly Pomacentridae and Gobiidae) and grazers (particularly Scaridae and Pomacentridae) occurred in very high abundances (Booth and Baretta 1994; Bruggemann et al. 1994).

Herbivores, detritivores and omnivores were overrepresented in the Bocas del Toro fish community compared to elsewhere in the Caribbean. Herbivorous species alone comprised nearly a third of the total fish biomass, which could be explained by a decreased number of predators in the system. Even though most herbivorous fishes were in the smallest size category (≤ 10 cm), this group has the potential to control the growth of macroalgae and prevent algal phase shifts, particularly in combination with invertebrate herbivores, such as sea urchins, which are abundant in this system (Kuempel and Altieri 2017). However, if the reduction of live coral cover continues, herbivorous fishes may reach their limits for grazing control (Williams and

Polunin 2001; Williams et al. 2001). Also, the lack of redundant species within the herbivore functional group is likely to result in low resilience, since a system with a single dominant herbivorous species *Scarus iseri* (72%) is vulnerable to stressors affecting that species (Hughes 1994; White and Jentsch 2001). The reason for the dominance of one herbivore species is probably attributable to the small body size of *S.iseri*, which matures at ~ 65 mm. It is therefore not a targeted fishery species, and escapes most fishing pressure (Kuempel and Altieri 2017).

Sponges cover up to 20% of substrata, and thus provide considerable physical structure on the Bocas del Toro reefs (Diaz and Rützler 2001; Loh and Pawlik 2014; Loh et al. 2015). In the absence of high cover of hard corals, sponges likely play an important role in supporting richness, biomass and expanded trophic levels of the depauperate fish community in our study system. Results furthermore suggest a positive effect and increased abundance of reef fishes with increased sponge cover. Sponges are major determinants of the rugosity and height of the reef (Diaz and Rützler 2001), thus could be an important driver for fish abundance and species richness in Bocas del Toro as in other Caribbean reef systems (Gratwicke and Speight 2005). Sponges also comprise an important food source for spongivorous reef fishes, such as some members of Pomacentridae and Scarinae (Sammarco et al. 1987; Dunlap and Pawlik 1996; Pawlik 1998; Souza et al. 2011). The pomacentrid *A. saxatilis* has been identified to have a functional dependency on sponges, through either shelter or other aspects of habitat complexity that sponges provide (Gratwicke and Speight 2005).

Proximity to mangroves was another important positive factor associated with fish communities, as the biomass and richness of fishes were greater on coral reefs in close proximity to

mangroves. Mangroves are widely recognized for their functions of providing nursery grounds, shelter and food sources for reef fishes (Laegdsgaard and Johnson 2001; Mumby et al. 2004). Our study suggests that the positive effect of mangroves as nursery and alternative habitats is an important factor maintaining diversity and biomass of the reef fish communities, and that this function remains particularly important in a system as degraded as Bocas del Toro. However, we did not find such evidence for seagrass meadows. The closer mangroves are located to reefs, the more effective is their role as nursery or alternative habitat. Lowest fish biodiversity, biomass and trophic level was found on reefs without mangroves in close proximity, presumably because many reef fish species depend on interconnectivity between habitat types (Ley 2014).

Bocas del Toro arguably represents a good model system for reef fish communities that are associated with high levels of anthropogenic stress. Trends suggest that stressed systems are increasingly moving to low diversity, low mean trophic level, and a size distribution skewed to small body size (Pauly et al. 1998). To maintain reef fish communities, resource managers can direct local fisheries and take factors such as sponge cover and connectivity to other habitats including mangroves into consideration to prioritize protection efforts. Our results suggest that reef sponges and mangroves together can maintain physical structure, act as nurseries, and provide alternative habitats and there by compensate for particular functional losses during coral mortality events. Much more information is nevertheless needed on the role of habitat connectivity if fisheries management is to be optimized and diversity hotspots safeguarded through effective marine protected areas (Linton and Warner 2003; Unsworth et al. 2008).

Acknowledgements

We thank divers who helped with the fish surveys and fish identification or benthic surveys, especially Scott Jones, Zachary Foltz, Ross Whippo, Justin Campbell, Jan Vincente and Seamus Harrison. We thank the people from the Bocas Research Station team for logistical help and for assistance with all aspects of the work, particularly Plinio Gondola. This is contribution number X from the Smithsonian's MarineGEO network.

References

- Alevizon W, Porter J (2015) Coral loss and fish guild stability on a Caribbean coral reef: 1974–2000. *Environmental Biology of Fishes* 98:1035–1045
- Altieri AH, Harrison SB, Seemann J, Collin R, Diaz RJ, Knowlton N (2017) Tropical dead zones and mass mortalities on coral reefs. *Proceedings of the National Academy of Sciences*
- Alvarez-Filip L, Dulvy NK, Gill JA, Côté IM, Watkinson AR (2009) Flattening of Caribbean coral reefs: region-wide declines in architectural complexity. *Proceedings of the Royal Society of London Series B-Biological Sciences* 276:3019–3025
- Aronson RB, Hilbun NL, Bianchi TS, Filley TR, Mckee BA (2014) Land use, water quality, and the history of coral assemblages at Bocas del Toro, Panamá. *Marine Ecology Progress Series* 504:159–170
- Aronson RB, Bruno JF, Precht WF, Glynn PW, Harvell CD, Kaufman L, Rogers CS, Shinn EA, Valentine JF (2003) Causes of coral reef degradation. *Science* 302:1502–1504
- Beck MW, Heck Jr KL, Able KW, Childers DL, Eggleston DB, Gillanders BM, Halpern B, Hays CG, Hoshino K, Minello TJ (2001) The identification, conservation, and management of estuarine and marine nurseries for fish and invertebrates: a better understanding of the habitats that serve as nurseries for marine species and the factors that create site-specific variability in nursery quality will improve conservation and management of these areas. *Bioscience* 51:633–641
- Bell J, Galzin R (1984) Influence of live coral cover on coral-reef fish communities. *Marine Ecology Progress Series* 15:265–274
- Bellwood DR, Hughes TP, Folke C, Nystrom M (2004) Confronting the coral reef crisis. *Nature* 429:827
- Beukers JS, Jones GP (1998) Habitat complexity modifies the impact of piscivores on a coral reef fish population. *Oecologia* 114:50–59
- Beulig A (1999) Relative influence of terrigenous vs. reef carbonate silt on turbidity and coral distribution at Bocas del Toro, Panama. *International Conference on Scientific Aspects of Coral Reef Assessment, Monitoring and Restoration*
- Booth DJ, Beretta GA (1994) Seasonal recruitment, habitat associations and survival of pomacentrid reef fish in the US Virgin Islands. *Coral Reefs* 13:81–89
- Bruggemann JH, Kuyper MWM, Breeman AM (1994) Comparative analysis of foraging and habitat use by the sympatric Caribbean parrotfish *Scarus vetula* and *Sparisoma viride* (Scaridae). *Marine Ecology Progress Series* 112:51–66
- Cesar H, Burke L, Pet-Soede L (2003) The economics of worldwide coral reef degradation. *Cesar environmental economics consulting (CEEC)*
- Cesar HS (2000) Coral reefs: their functions, threats and economic value. *Collected essays on the economics of coral reefs*:14–39
- Cinner JE, McClanahan TR (2006) Socioeconomic factors that lead to overfishing in small-scale coral reef fisheries of Papua New Guinea. *Environmental Conservation* 33:73–80
- Collin R (2005) Ecological monitoring and biodiversity surveys at the Smithsonian Tropical Research Institute's Bocas del Toro Research Station. *Caribbean Journal of Science* 41:367–373

- Cramer KL (2013) History of Human Occupation and Environmental Change in Western and Central Caribbean Panama. *Bulletin of Marine Science* 89:955-982
- D'Croz L, Del Rosario JB, Gondola P (2005) The effect of fresh water runoff on the distribution of dissolved inorganic nutrients and plankton in the Bocas del Toro Archipelago, Caribbean Panama. *Caribbean Journal of Science* 41:414-429
- Dahlgren CP, Kellison GT, Adams AJ, Gillanders BM, Kendall MS, Layman CA, Ley JA, Nagelkerken I, Serafy JE (2006) Marine nurseries and effective juvenile habitats concepts and applications. *Marine Ecology Progress Series* 312:291-295
- Diaz MC, Rützler K (2001) Sponges: An essential component of Caribbean coral reefs. *Bulletin of Marine Science* 69:535-546
- Dominici-Arosemena A, Wolff M (2005) Reef fish community structure in Bocas del Toro (Caribbean, Panama): gradients in habitat complexity and exposure. *Caribbean Journal of Science* 41:613-637
- Dunlap M, Pawlik JR (1996) Video-monitored predation by Caribbean reef fishes on an array of mangrove and reef sponges. *Marine Biology* 126:117-123
- Fabrizius K, De'ath G, McCook L, Turak E, Williams DM (2005) Changes in algal, coral and fish assemblages along water quality gradients on the inshore Great Barrier Reef. *Marine Pollution Bulletin* 51:384-398
- Galzin R, Planes S, Dufour V, Salvat B (1994) Variation in diversity of coral reef fish between French Polynesian atolls. *Coral Reefs* 13:175-180
- Gardner TA, Côté IM, Gill JA, Grant A, Watkinson AR (2003) Long-term region-wide declines in Caribbean corals. *Science* 301:958-960
- Granek EF, Compton JE, Phillips DL (2009) Mangrove-Exported Nutrient Incorporation by Sessile Coral Reef Invertebrates. *Ecosystems* 12:462-472
- Gratwicke B, Speight M (2005) The relationship between fish species richness, abundance and habitat complexity in a range of shallow tropical marine habitats. *Journal of Fish Biology* 66:650-667
- Guzmán HM, Jiménez CE (1992) Contamination of coral reefs by heavy metals along the Caribbean coast of Central America (Costa Rica and Panama). *Marine Pollution Bulletin* 24:554-561
- Guzmán HM, Barnes PAG, Lovelock CE, Feller IC (2005) A site description of the CARICOMP mangrove, seagrass and coral reef sites in Bocas del Toro, Panama *Caribbean Journal of Science* 41:430-440
- Hodgson G (1999) A global assessment of human effects on coral reefs. *Marine Pollution Bulletin* 38:345-355
- Hughes TP (1994) Catastrophes, phase shifts, and large-scale degradation of a Caribbean coral reef. *Science* 265:1547-1551
- Jackson JBC, Kirby MX, Berger WH, Bjorndal KA, Botsford LW, Bourque BJ, Bradbury RH, Cooke R, Erlandson J, Estes JA, Hughes TP, Kidwell S, Lange CB, Lenihan HS, Pandolfi JM, Peterson CH, Steneck RS, Tegner MJ, Warner RR (2001) Historical Overfishing and the Recent Collapse of Coastal Ecosystems. *Science* 293:629-637
- Jennings S, Lock JM (1996) Population and ecosystem effects of reef fishing. In: Polunin NVC, Roberts CM (eds) *Reef Fisheries*. Springer Netherlands, Dordrecht, pp193-218
- Kaufmann KW, Thompson RC (2005) Water temperature variation and the meteorological and hydrographic environment of Bocas del Toro, Panama. *Caribbean Journal of Science* 41:392-413
- Knowlton N, Brainard RE, Fisher R, Moews M, Plaisance L, Caley MJ (2010) Coral reef biodiversity. *Life in the World's Oceans: Diversity Distribution and Abundance*:65-74
- Kuempel CD, Altieri AH (2017) The emergent role of small-bodied herbivores in pre-empting phase shifts on degraded coral reefs. *Scientific Reports* 7:39670
- Kuffner IB, Brock JC, Grober-Dunsmore R, Bonito VE, Hickey TD, Wright CW (2007) Relationships Between Reef Fish Communities and Remotely Sensed Rugosity Measurements in Biscayne National Park, Florida, USA. *Environmental Biology of Fishes* 78:71-82
- Laegdsgaard P, Johnson C (2001) Why do juvenile fish utilise mangrove habitats? *Journal of experimental marine biology and ecology* 257:229-253
- Lee S (1995) Mangrove outwelling: a review. *Hydrobiologia* 295:203-212
- Leinfelder RR, Seemann J, Heiss GA, Struck U (2012) Could 'ecosystem atavisms' help reefs to adapt to the Anthropocene? *Proceedings of the 12th International Coral Reef Symposium ICRS2012_2B_2:5*
- Ley JA (2014) Mangrove connectivity helps sustain coral reef fisheries under global climate change. *Interrelationships Between Corals and Fisheries*:171
- Linton DM, Warner GF (2003) Biological indicators in the Caribbean coastal zone and their role in integrated coastal management. *Ocean & Coastal Management* 46:261-276

- 494 Loh T-L, Pawlik JR (2014) Chemical defenses and resource trade-offs structure sponge communities on
495 Caribbean coral reefs. *Proceedings of the National Academy of Sciences* 111:4151-4156
- 496 Loh T-L, McMurray SE, Henkel TP, Vicente J, Pawlik JR (2015) Indirect effects of overfishing on Caribbean reefs:
497 sponges overgrow reef-building corals. *PeerJ* 3:e901
- 498 Mora C (2015) *Ecology of Fishes on Coral Reefs*. Cambridge University Press
- 499 Mumby PJ, Edwards AJ, Arias-González JE, Lindeman KC, Blackwell PG, Gall A, Gorczynska MI, Harborne AR,
500 Pescod CL, Renken H (2004) Mangroves enhance the biomass of coral reef fish communities in the
501 Caribbean. *Nature* 427:533-536
- 502 Mumby PJ, Dahlgren CP, Harborne AR, Kappel CV, Micheli F, Brumbaugh DR, Holmes KE, Mendes JM, Broad K,
503 Sanchirico JN (2006) Fishing, trophic cascades, and the process of grazing on coral reefs. *science*
504 311:98-101
- 505 Myers RA, Worm B (2003) Rapid worldwide depletion of predatory fish communities. *Nature* 423:280-283
- 506 Nagelkerken I, van der Velde G, Gorissen MW, Meijer GJ, Van't Hof T, den Hartog C (2000) Importance of
507 Mangroves, Seagrass Beds and the Shallow Coral Reef as a Nursery for Important Coral Reef Fishes,
508 Using a Visual Census Technique. *Estuarine, Coastal and Shelf Science* 51:31-44
- 509 Nagelkerken I, Roberts C, Van Der Velde G, Dorenbosch M, Van Riel M, De La Moriniere EC, Nienhuis P (2002)
510 How important are mangroves and seagrass beds for coral-reef fish? The nursery hypothesis tested on
511 an island scale. *Marine ecology progress series* 244:299-305
- 512 Paddock MJ, Reynolds JD, Aguilar C, Appeldoorn RS, Beets J, Burkett EW, Chittaro PM, Clarke K, Esteves R,
513 Fonseca AC (2009) Recent region-wide declines in Caribbean reef fish abundance. *Current Biology*
514 19:590-595
- 515 Pauly D, Christensen V, Dalsgaard J, Froese R, Torres F (1998) Fishing down marine food webs. *Science*
516 279:860-863
- 517 Pawlik JR (1998) Coral reef sponges: Do predatory fishes affect their distribution? *Limnology and*
518 *Oceanography* 43:1396-1399
- 519 Riegl B, Purkis SJ, Keck J, Rowlands GP (2009) Monitored and modeled coral population dynamics and the
520 refuge concept. *Marine Pollution Bulletin* 58:24-38
- 521 Saila S, Kocic VL, McManus J (1993) Modelling the effects of destructive fishing practices on tropical coral reefs.
522 *Marine Ecology Progress Series* 94:51-60
- 523 Sammarco PW, Strychar KB (2009) Effects of Climate Change/Global Warming on Coral Reefs:
524 Adaptation/Exaptation in Corals, Evolution in Zooxanthellae, and Biogeographic Shifts. *Environmental*
525 *Bioindicators* 4:9 - 45
- 526 Sammarco PW, Risk MJ, Rose C (1987) Effects of grazing and damselfish territoriality on internal bioerosion of
527 dead corals : indirect effects. *Journal of Experimental Marine Biology and Ecology* 112:185-199
- 528 Seemann J (2013) The use of 13C and 15N isotope labeling techniques to assess heterotrophy of corals. *Journal*
529 *of Experimental Marine Biology and Ecology* 442:88-95
- 530 Seemann J, Gonzalez CT, Carballo-Bolaños R, Berry K, Heiss GA, Struck U, Leinfelder RR (2014) Assessing the
531 ecological effects of human impacts on coral reefs in Bocas del Toro, Panama. *Environmental*
532 *Monitoring and Assessment* 186:1747-1763
- 533 Serafy JE, Shideler GS, Araújo RJ, Nagelkerken I (2015) Mangroves Enhance Reef Fish Abundance at the
534 Caribbean Regional Scale. *PloS one* 10:e0142022
- 535 Souza A, Ilarri M, Rosa I (2011) Habitat use, feeding and territorial behavior of a Brazilian endemic damselfish
536 *Stegastes rocasensis* (Actinopterygii: Pomacentridae). *Environmental Biology of Fishes* 91:133-144
- 537 Spalding MD, Fox HE, Allen GR, Davidson N, Ferdaña ZA, Finlayson M, Halpern BS, Jorge MA, Lombana A, Lourie
538 SA (2007) Marine ecoregions of the world: a bioregionalization of coastal and shelf areas. *BioScience*
539 57:573-583
- 540 Turner SJ, Thrush SF, Hewitt JE, Cummings VJ, Funnell G (1999) Fishing impacts and the degradation or loss of
541 habitat structure. *Fisheries Management and Ecology* 6:401-420
- 542 Unsworth RK, Salinas De Leon P, Garrard SL, Jompa J, Smith DJ, Bell JJ (2008) High connectivity of Indo-Pacific
543 seagrass fish assemblages with mangrove and coral reef habitats. *MARINE ECOLOGY-PROGRESS*
544 *SERIES-* 353:213
- 545 White PS, Jentsch A (2001) The search for generality in studies of disturbance and ecosystem dynamics
546 *Progress in botany*. Springer, pp399-450
- 547 Williams I, Polunin N (2001) Large-scale associations between macroalgal cover and grazer biomass on mid-
548 depth reefs in the Caribbean. *Coral reefs* 19:358-366

549 Williams ID, Polunin NVC, Hendrick VJ (2001) Limits to grazing by herbivorous fishes and the impact of low
 550 coral cover on macroalgal abundance on a coral reef in Belize. *Marine Ecology Progress Series*
 551 222:187-196
 552 Wilson SK (2006) Multiple disturbances and the global degradation of coral reefs: are reef fishes at risk or
 553 resilient? *Global change biology* 12:2220-2234
 554 Wilson SK, Fisher R, Pratchett MS, Graham NAJ, Dulvy NK, Turner RA, Cakacaka A, Polunin NVC (2010) Habitat
 555 degradation and fishing effects on the size structure of coral reef fish communities. *Ecological*
 556 *Applications* 20:442-451
 557 Zaneveld JR, Burkepile DE, Shantz AA, Pritchard CE, McMinds R, Payet JP, Welsh R, Correa AMS, Lemoine NP,
 558 Rosales S, Fuchs C, Maynard JA, Thurber RV (2016) Overfishing and nutrient pollution interact with
 559 temperature to disrupt coral reefs down to microbial scales. *Nature Communications* 7:11833
 560

Figure 1

Sampling sites in Bocas del Toro

Three reef sites (Punta Caracol, Casa Blanca, Almirante) possess close connectivity with mangrove habitat (within 100 m), three sites (STRI, Juan Point, Coral Cay) represent reef sites further away from mangroves (100 - 250 m), and three reef sites (Popa, Salt Creek, Hospital Point) are not closely connected to mangroves (> 750 m). Yellow areas are reefs and green areas are mangroves islands, gray is land, white is ocean, blue is river and blue polygon is a poorly enforced MPA.

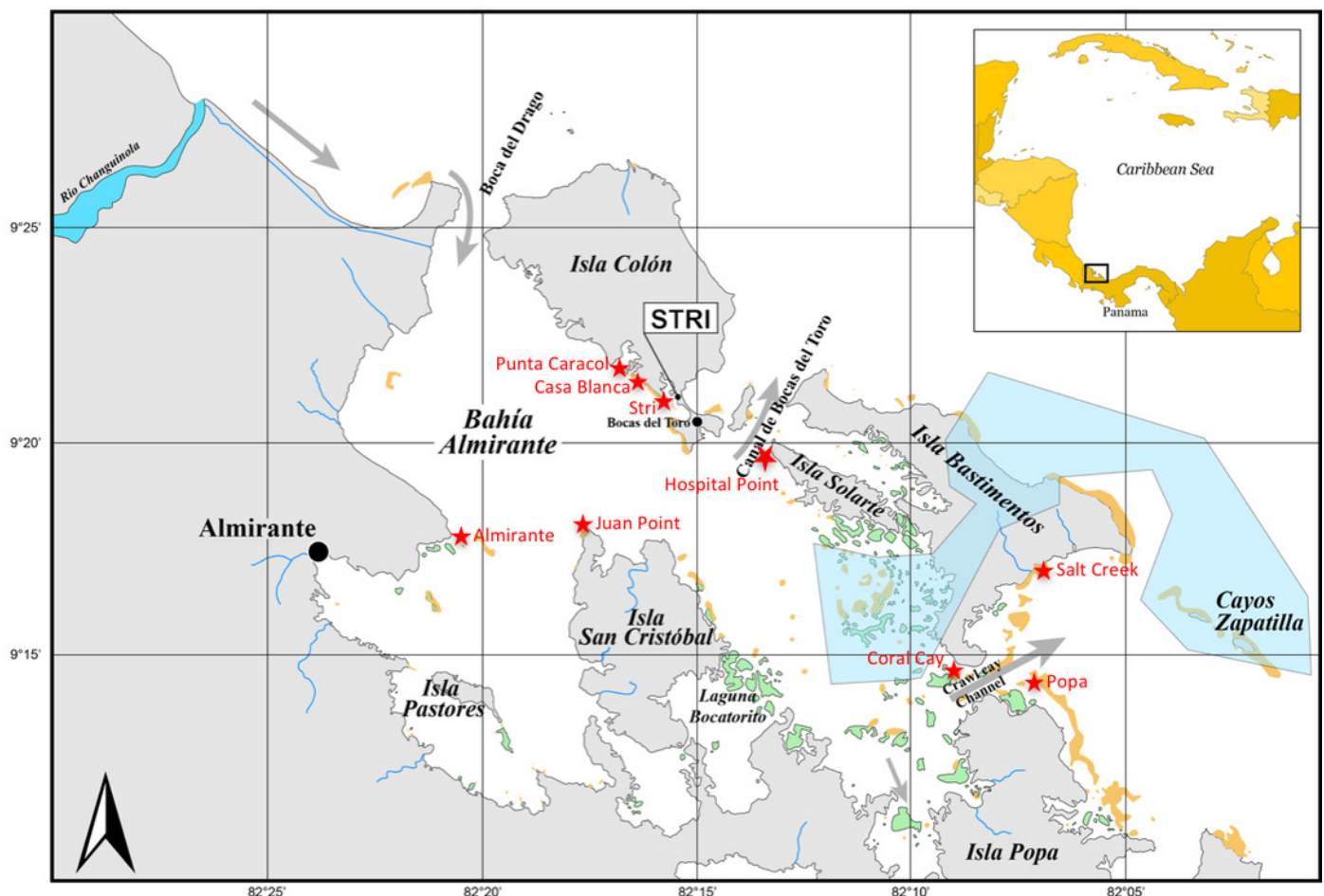


Figure 2

Biomass and composition of the fish community in the Caribbean and Bocas del Toro.

A) The comparison of the total biomass ($AVR \pm SD$) from RLS conducted across the Caribbean, divided in no take zones, restricted zone and open zones, and open zones in Bocas del Toro, groups with different letters are significantly different. B) Distribution of trophic guilds based on total biomass: high-level carnivores (trophic level 4 – 4.5), low and mid-level carnivore (trophic level 2.8 – 3.9), herbivores, omnivores and detritivores (trophic level 2 – 2.7). C) The abundance of fish subdivided in size classes ($AVR \pm SD$), which are indicative of fishing pressure (skew towards smaller body size implies fishing). Asterisk represents significant differences between size abundance data from Bocas and the Caribbean.

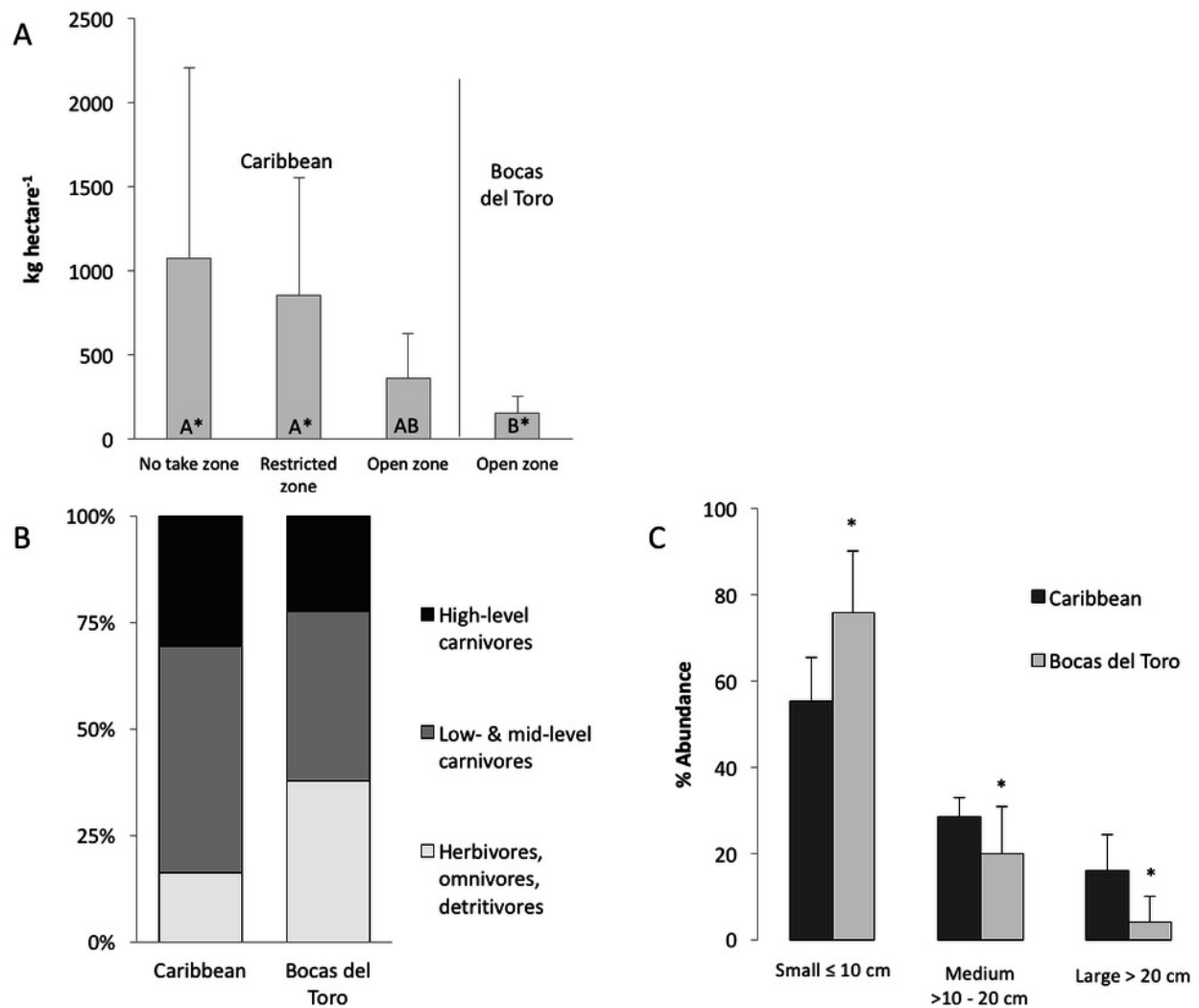


Figure 3

Biomass of trophic guilds of reef fish.

Data were pooled by sites with a similar distance to mangroves: <100 m (Punta Caracol, Casa Blanca, Almirante), 100 – 250 m (STRI, Juan Point, Coral Cay) and >700 m (Popa, Salt Creek, Hospital Point) (see Table 1 for detail).

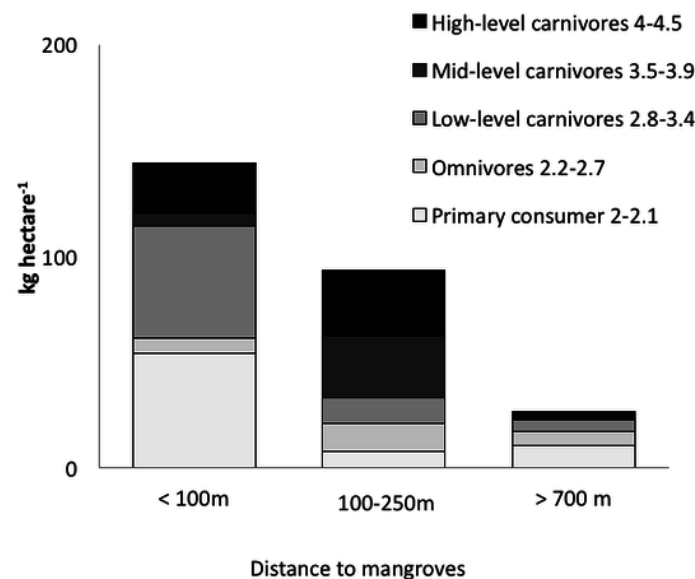


Figure 4

Correlation of reef fish abundance (w/o small-bodied fish) and (A) distance to mangroves or (B) sponge percent cover.

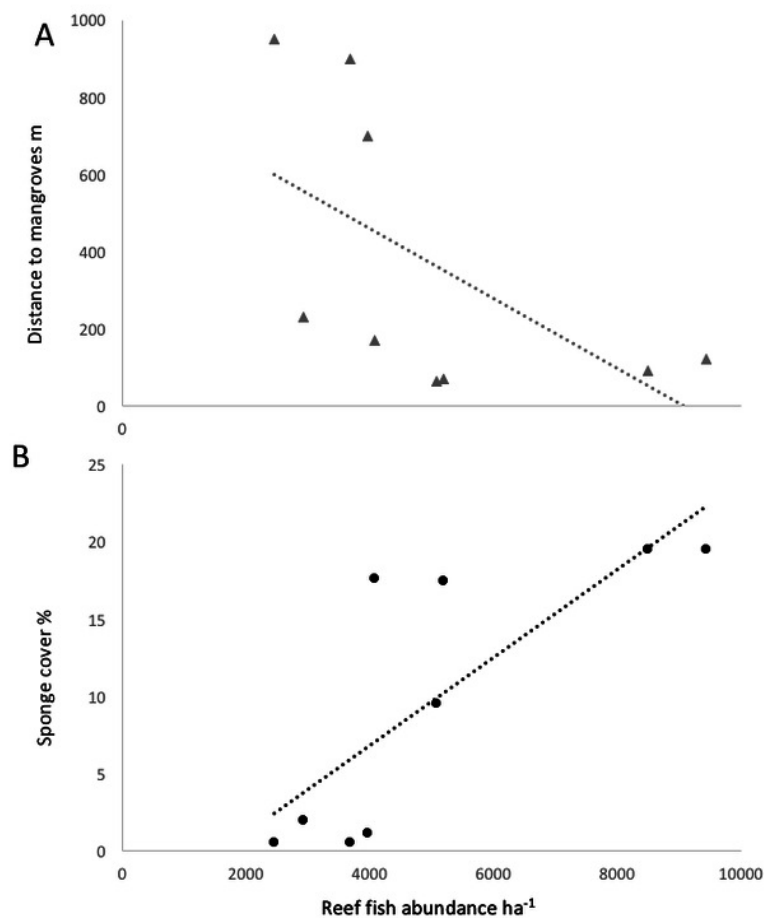


Figure 5

Principal component analyses

This PCA shows the clustering of fish communities, considering species composition and abundance (log transformed). Small bodied fish ≤ 12.5 cm was excluded. Reefs >700 m from mangroves cluster together as a group, separate from the reefs that are closer to mangroves. This indicates that mangrove distance has a strong influence on reef community types. The numbers refer to the site numbers in Table 1.

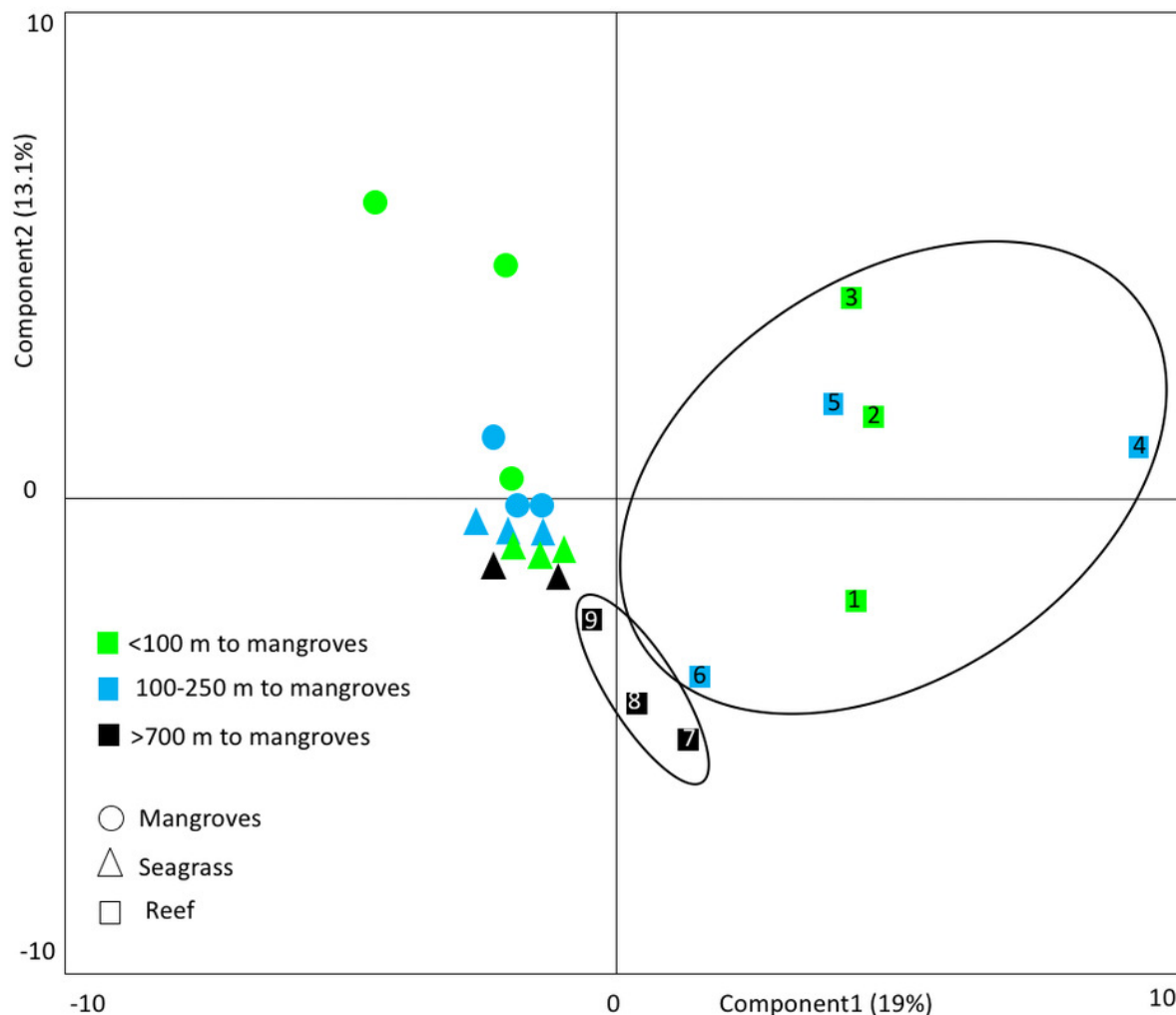


Table 1(on next page)

Major habitat characteristics and location of monitoring sites.

Sites 7, 8 and 9 did not have mangroves in close proximity (≤ 250 m); site 9 also did not have a seagrass meadow close to the reef, effective juvenile habitats were calculated from the fish abundance in mangroves (w/o small bodied fish) compared to reef fish abundance.

1

	Site	Coordinates Lat	Coordinates Long	Depth Reef (m)	Depth Seagrass (m)	Distance Reef- Mangrove (m)	Sponge Cover %	Live Hard Coral Cover %	Hard Substrate %	Reef Fish Biomass kg ha ⁻¹	Seagrass Fish Biomass kg ha ⁻¹	Mangrove Fish Biomass kg ha ⁻¹	Seagrass as Effective Juvenile Habitats %	Mangroves as Effective Juvenile Habitats %	Reef Fish Abundance ha ⁻¹	Seagrass Fish Abundance ha ⁻¹	Mangrove Fish Abundance ha ⁻¹	Reef Fish Richness	Seagrass Fish Richness	Mangrove Fish Richness
1	Punta Caracol	9.3757°	-82.2997°	3	2	65	9.5	41.5	33	201	25	111	109	131	12929	2820	17423	38	12	21
2	Casa Blanca	9.3588°	-82.2737°	3	1	70	17.5	2.5	71	67	32	47	102	73	18741	67660	17570	30	9	16
3	Almirante	9.2900°	-82.3429°	3	2	90	19.5	36.5	71	206	2	202	13	59	11105	2560	1202510	28	6	15
4	STRI Point	9.3483°	-82.2625°	3	4	120	19.5	3.0	57	257	24	15	49	29	71076	73153	42390	35	15	19
5	Juan Point	9.3003°	-82.2921°	4	1	170	17.6	46.4	69	94	14	32	86	29	24045	31760	200660	30	11	10
6	Coral Cay	9.2435°	-82.1478°	5	2	230	2.0	16.0	51	25	12	11	54	139	1717	50850	42060	25	7	9
7	Popa	9.2336°	-82.1120°	3	1	700	1.1	26.9	61	60	2		23		2608	560	17423	24	9	
8	Salt Creek	9.2815°	-82.1012°	6	2	950	0.5	24.8	99	13	0		105		1688	1290		15	12	
9	Hospital Point	9.3326°	-82.2220°	5.5		900	0.5	96.0	33	12					1946			16		

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Table 2 (on next page)

Major fish families (only considering >10 counts ha^{-1} in average in one of the size bins).

1

Caribbean				Bocas del Toro								
Reef				Reef	Seagrass	Mangrove	Reef	Seagrass	Mangrove	Reef	Seagrass	Mangrove
≤10 cm	>10-20 cm	>20 cm		≤10 cm	≤10 cm	≤10 cm	>10-20 cm	>10-20 cm	>10-20 cm	>20 cm	>20 cm	>20 cm
Acanthuridae	317	351	127	113	127	120	233	0	20	0	0	0
Balistidae	100	380	145	0	0	0	0	0	0	0	0	0
Carangidae	296	1078	145	321	20	2593	330	80	100	40	0	0
Clupeidae	11500	0	0	0	46000	278080	0	0	0	0	0	0
Ephippidae	0	80	280	0	0	0	30	0	0	0	0	0
Gerreidae	0	30	20	0	600	155	0	0	20	0	0	0
Gobiidae	6239	0	0	18182	30	80	0	0	0	0	0	0
Grammatidae	434	0	0	0	0	0	0	0	0	0	0	0
Haemulidae	1959	1395	160	379	752	823	457	300	70	20	0	0
Holocentridae	253	441	50	0	0	0	20	0	0	0	0	0
Inermiidae	300	3444	0	0	0	0	0	0	0	0	0	0
Kyphosidae	463	733	160	0	0	0	0	0	0	0	0	0
Labridae	1749	659	96	254	568	580	252	180	80	0	0	0
Loliginidae	0	240	0	0	0	0	0	0	0	0	0	0
Lutjanidae	263	800	279	80	137	559	80	20	350	0	0	20
Mullidae	245	429	229	50	20	0	20	0	0	0	0	0
Pomacentridae	2145	414	20	618	325	123	110	0	0	0	0	0
Scaridae	741	252	196	494	753	979	333	20	173	80	0	0
Sciaenidae	532	176	40	60	0	0	60	0	0	0	0	0
Serranidae	855	107	208	297	72	40	93	0	0	0	0	0
Sphyraenidae	120	2100	180	0	0	40	0	20	40	0	0	100
Tetraodontidae	247	0	0	193	100	20	0	0	0	0	0	0

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