Importance of differentiating Orbicella reefs from gorgonian plains for ecological assessments of Caribbean reefs

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Importance of differentiating *Orbicella* reefs from gorgonian plains for ecological assessments of Caribbean reefs

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ABSTRACT: Region-wide assessments of coral cover typically rely on meta-analyses of small-scale ecological studies which have combined different coral reef habitats. This is particularly problematic on forereefs where at least 2 habitats can be found; coral-based bioherms and colonized hardgrounds (hereafter *Orbicella* reefs and gorgonian plains), each with very different structure and scleractinian coral cover. Here, we quantify the degree to which the failure to differentiate forereef zones dominated by framework building corals, mainly *Orbicella* spp. (hereafter *Orbicella* reefs) from gorgonian plains can lead to biased assessments of coral cover. We also provide a baseline of an extensive sample of Caribbean coral reefs in 2010−2012 for the 2 habitats within the forereef. Mean scleractinian coral cover (±SE) at *Orbicella* reefs was 24 ± 1.3%, more than double the coral cover found on the gorgonian plains (10 ± 1.6%). The difference in coral cover between habitats within the same geomorphological zone is consistent with those calculated from an independent dataset for the basin (Atlantic and Gulf Rapid Reef Assessment). Furthermore, the average coral cover calculated for Caribbean *Orbicella* reefs was more than double the values previously reported for entire reefs in the region a decade ago (10%), which integrated data from different habitats, depths, time periods and surveyors. Differentiating between forereef habitats has provided a meaningful baseline of coral state, which allows for realistic targets for management in the Caribbean basin.

KEY WORDS: Coral · Forereef habitats · *Orbicella* reefs · Gorgonian plains · Caribbean · AGRRA

INTRODUCTION

Caribbean coral reefs have changed dramatically in the last 5 decades (Jackson et al. 2014); the abundance of reef-associated organisms, especially corals and fishes, have suffered massive declines through hurricanes, disease outbreaks, bleaching, pollution, sedimentation and overfishing (Hughes 1994, Aronsen & Precht 2001, Kramer 2003, Weil et al. 2006, Paddack et al. 2009). A number of studies have quantified the state of Caribbean reefs at a regional scale and, in some cases, their trajectory (Gardner et al. 2003, Côté et al. 2005, Bruno et al. 2009, Schutte et al. 2010, Jackson et al. 2014). Regional assessments allow a benchmark to be set against which individual reefs can be compared in space and time, quantifying regional trends (Jackson et al. 2014) and comparing different regions to identify the drivers of large-scale differences (Roff & Mumby 2012). Most regional assessments have used meta-analysis to integrate
the results of multiple studies carried out by different surveyors, usually with different objectives in mind. An exception has been the analysis of regionally standardized survey methodologies, such as the Atlantic and Gulf Rapid Reef Assessment (AGGRA) (Kramer 2003), although this too suffers from the issue of multiple surveyors and habitat types, as well as some methodological inconsistencies that increase the variability in the data (e.g. variable sampling effort, haphazardly located transects) (Kramer 2003).

Regional meta-analyses have been shown to be robust to common inconsistencies among data, such as variable survey methodology (Côté et al. 2005), however they cannot account for inconsistencies that have not been recorded. In particular, it is typically assumed that reefs can be grouped by major reef zone (e.g. patch reef, backreef, forereef) and depth. Yet, forereef environments, which can be identified by their geomorphology (Andréfouët et al. 2006), comprise at least 2 contrasting habitats even at the same depth. The first are true coral habitats that were generally built by the major framework-building corals *Orcbigella* spp. and some species of acroporids (Goreau 1959, Geister 1977, Mumby & Harborne 1999). These habitats, hereafter *Orcbigella* reefs, can be structurally complex and usually possess high diversity and density of species, while also supporting the most significant ecosystem services, which include cultural, educational and fishery services (Mumby et al. 2008, Harborne 2009). The second are ‘gorgonian plains’ (Mumby & Harborne 1999, Mumby 2014), also known as hardgrounds, which are relatively featureless, flat pavement that typically possesses very low relief and is dominated by octocorals. Gorgonian plains are usually located in more exposed environments (Burke 1982, Torres et al. 2001), and have very little scleractinian coral, likely because higher exposure resuspends sediments near the benthos, providing an unstable substrate for recruitment. However, octocorals, the predominant benthic invertebrates on this habitat, are more resilient to fluctuating habitats (Opresko 1973) and commonly recruit and survive in areas with water movement (Barham & Davies 1968, Yoshioka & Yoshioka 1989). Few empirical studies identify habitat type explicitly and, with the exception of CARICOMP (Caribbean Coastal Marine Productivity Program) (CARICOMP 2002), most regional survey methodologies, including AGRRA, combine all forereef habitats together during analyses. Regional assessments of coral reef condition have therefore potentially combined hardbottom habitats that are naturally characterized by low coral cover with coral-based bioherms. Such confounding would tend to reduce the average coral cover by including sites that lack and have always lacked high abundance of coral, in an unbalanced design through time and space (i.e. proportion of sites per habitat is unlikely to be the same), which would therefore add noise to the signal and obscure spatial and temporal patterns within the basin.

Recent advances with remote sensing have now made it possible to discriminate *Orcbigella* reefs from gorgonian plains at a large scale. Several techniques are available, including the use of acoustic sensors that detect the roughness and hardness of the seabed (Bejarano et al. 2010, 2011), as well as measures of wave exposure, which have been shown to accurately (81%) discriminate *Orcbigella* reefs from gorgonian plains within the forereef, given that gorgonian plains thrive in heavily exposed environments (Chollett & Mumby 2012). Wave exposure can be obtained using only information on the configuration of the coastline and prevailing wind speed and direction, representing a cheap alternative for the large-scale mapping of forereef habitats (Chollett & Mumby 2012). In the present study, one of our major goals was to assess the impact of confounding forereef habitats on average scleractinian coral cover using a dataset gathered by the FORCE (Future of Reefs in a Changing Environment) project, collected with the same methods and surveyors in 12 countries in the Caribbean. We validated our results with an independent dataset available for the region, AGRRA, and put our results into context by comparing our estimates of current scleractinian coral cover to other regional studies (Gardner et al. 2003, Jackson et al. 2014).

**MATERIALS AND METHODS**

**Data collection (FORCE)**

Coral reef benthic communities were surveyed from 2010 to 2012 in 12 countries: Honduras, Belize, Curaçao, Bonaire, Jamaica, Barbados, Dominican Republic, Antigua, St. Lucia, St. Vincent and the Grenadines, Costa Rica and Panamá (Fig. 1). Sites within the forereef according to geomorphological maps produced by the Millennium Coral Reef Mapping project (Andréfouët et al. 2006), were selected using the ‘generate random points’ tool of the Hawth Tools program in ArcGIS 9.2 (ESRI 2009). Sites were assessed *in situ* and defined as having a homogeneous habitat within a spatial extent of at least 200 × 200 m. Originally, we set our experimental design to assess the current state...
of forereefs across the Caribbean, and at the time of sampling we lacked the recently developed tools to identify forereef habitats based on wave exposure (Chollett & Mumby 2012). Consequently, the number of sites per habitat is not balanced among countries (Table 1). This shortcoming has been taken into account in the statistical analyses (see following subsection). Surveys were conducted at 92 *Orbicella* reefs (Fig. 2a) and at 28 gorgonian plains (Fig. 2b) at 10–15 m depths with the exception of Honduras and Belize where surveys were conducted at 5–10 m and 15–20 m (Table 1). Coral cover did not vary significantly between depths in Honduras and Belize, therefore transects from the different depth zones were pooled for further statistical analyses. At each site, benthic communities were assessed on 6 randomly placed 10 m transect lines which were set parallel to the coastline. Scleractinian coral cover was measured using the point intercept method and corals were identified to species level and recorded every 10 cm. There are some shortcomings to using point intercept technique, in that coral cover estimate may be lower in habitats that are characterized by small coral colonies, common on gorgonian plains.

To assess the sufficiency of our sampling given the sample size, the margin of error in coral cover was calculated for each forereef habitat. The margins of error were moderate, with a median of 4% for gorgonian habitat. The margins of error were moderate, with a median of 4% for gorgonian habitat and 6% for *Orbicella* reefs.

### Statistical analysis

#### Habitat differences (FORCE data)

To test whether combining forereef habitat types skewed coral cover estimate in the Caribbean basin and within individual countries, we used generalized linear mixed models (GLMM) (Bolker et al. 2009) using the lme4 package (Bates et al. 2014) in R. We identified the effects of forereef habitat on the cover of coral using the entire FORCE dataset. Coral cover was the response variable, while habitat was treated as the fixed factor and sites and countries were random, with sites nested within countries. To select the optimal structure of the random component, we ran 2 models, the first with no random term and the second with the random component using site nested in country. The optimal model was chosen based on the lowest Akaike’s information criteria (AIC) value. We used Poisson distributions, given that the response variable was a proportion (percentage of coral), and estimated the parameters with Laplace approximations because there were fewer than 5 random variables (Breslow & Clayton 1993) for

<table>
<thead>
<tr>
<th>Country</th>
<th><em>Orbicella</em></th>
<th>Gorgonian</th>
<th>Depth (m)</th>
<th>Sampling date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antigua (ANT)</td>
<td>3</td>
<td>5</td>
<td>10−15</td>
<td>Jul 2011</td>
</tr>
<tr>
<td>Barbados (BAR)</td>
<td>8</td>
<td>0</td>
<td>10−15</td>
<td>Mar 2011</td>
</tr>
<tr>
<td>Belize (BEL)</td>
<td>5</td>
<td>10</td>
<td>5−10, 15−20</td>
<td>Nov 2010</td>
</tr>
<tr>
<td>Bonaire (BON)</td>
<td>7</td>
<td>0</td>
<td>10−15</td>
<td>Jan 2011</td>
</tr>
<tr>
<td>Costa Rica (CR)</td>
<td>0</td>
<td>8</td>
<td>10−15</td>
<td>Apr 2012</td>
</tr>
<tr>
<td>Curaçao (CUR)</td>
<td>8</td>
<td>0</td>
<td>10−15</td>
<td>Jan 2011</td>
</tr>
<tr>
<td>Dominican Republic (DR)</td>
<td>7</td>
<td>7</td>
<td>10−15</td>
<td>Jun 2011</td>
</tr>
<tr>
<td>Honduras (HON)</td>
<td>11</td>
<td>3</td>
<td>5−10, 15−20</td>
<td>Oct 2010</td>
</tr>
<tr>
<td>Jamaica (JAM)</td>
<td>11</td>
<td>0</td>
<td>10−15</td>
<td>Feb 2011</td>
</tr>
<tr>
<td>Panamá (PAN)</td>
<td>8</td>
<td>0</td>
<td>10−15</td>
<td>Apr 2012</td>
</tr>
<tr>
<td>St. Lucia (STL)</td>
<td>7</td>
<td>1</td>
<td>10−15</td>
<td>Jul 2011</td>
</tr>
<tr>
<td>St. Vincent and Grenadines (SVG)</td>
<td>12</td>
<td>3</td>
<td>10−15</td>
<td>Aug 2011</td>
</tr>
</tbody>
</table>
GLMMs. The optimal model was validated by plotting the residuals against the fitted values and using a quantile-quantile plot. Significance levels (p-values) were obtained from Z tests (Zuur et al. 2009).

Consistency of results (AGRRA data)

To confirm the differences in coral cover between habitats observed in the FORCE results, we repeated the analysis using the AGRRA (www.agrra.org) dataset. As a conservative measure we included pre-bleaching data (2005), therefore focusing on benthic surveys carried out between 1998 and 2011. Many sites in the AGRRA database have repeated measurements; therefore, we used the most current year of data for each site.

The AGRRA dataset includes a variety of habitat types; therefore, data were extracted from forereef habitats and further classified as *Orbicella* reefs and gorgonian plains using the method described by Chollett & Mumby (2012). We calculated the probability of occurrence of *Orbicella* reefs within forereef zones given a predictive binomial model that uses wave exposure as a sole predictor of habitat (Chollett & Mumby 2012); wave exposure values were extracted for each AGRRA site at a 50 m spatial resolution for the Mesoamerican Barrier Reef (Chollett & Mumby 2012), and a 1 km spatial resolution for the rest of the Caribbean (Chollett et al. 2012). Predicting habitat type from wave exposure has shown to be accurate (81 %) at a spatial resolution of 50 m (Chollett & Mumby 2012); however, the accuracy of the method is unknown when data at a lower spatial resolution are used. Therefore, we tested the accuracy of the 1 km² spatial resolution data on predicting habitat type using the FORCE dataset and permutation analysis. *Orbicella* reefs and gorgonian plains were misclassified 23 and 12 % (respectively) of the time with FORCE surveys, indicating that the accuracy of the predictive model is adequate at the spatial resolution for the assignment of habitat to the AGRRA data (1 km²). To acknowledge the uncertainty in habitat assignment in further analyses, we resampled the AGRRA dataset, and the mean coral cover reported is based on the propagated errors, i.e. the mean and standard errors were calculated from 100 permutations of samples, whereby 12 % of the samples (randomly selected from gorgonian plains data) were replaced with data from randomly selected *Orbicella* reef coral cover. This procedure was repeated for *Orbicella* reefs, replacing 23 % of the samples.

Within- and between-country comparisons (FORCE data)

Given the unbalanced design of the dataset, we assessed the variation in coral cover between the forereef habitats in countries where data existed for both habitats: Antigua (ANT), Belize (BEL), Dominican Republic (DR), Honduras (HON), and St. Vincent and Grenadines (SVG). We used a GLMM test to identify the effects of forereef habitat on coral cover within a particular country. Coral cover was the response variable, whilst country and habitat were fixed variables, and site was the random variable. Subsequently, we
tested the interaction between country and habitat, with site as a random variable. The relevance of the random component was assessed by comparing the 2 models described above. Model distributions and procedures for model selection, validation, and obtaining p-values were identical to those used in GLMMs for testing habitat differences (see ‘Habitat differences [FORCE data]’).

RESULTS

Habitat differences (FORCE data)

Coral cover was significantly greater (GLMM; \( p < 0.0001 \)) on *Orbicella* reefs (24 ± 1.3%, mean ± SE) than on gorgonian plains (10 ± 1.6%) (Table 2). We observed the random effects (site nested in country, country) to be significant (\( p < 0.0001 \)); standard errors of the random effects were 0.41 and 0.22, while variances were 0.17 and 0.05 for site nested in country and country, respectively. The frequency distribution of coral cover was skewed for both the habitats, whereby the median scleractinian coral cover on *Orbicella* reefs was 23%, more than double that of gorgonian plains (9%). Because *Orbicella* sites dominated the dataset, the overall average cover on both habitats was modestly high, at 20 ± 0.5%. Maximum recorded coral cover was 22.3% on gorgonian plains and 44.1% on *Orbicella* reefs (Fig. 3). Sixty-seven percent of gorgonian plain sites exhibited a coral cover ≤10%, while 55% of *Orbicella* reefs comprised a coral cover >20%.

Table 2. Effects of forereef habitat (Habitat) on coral cover in the Caribbean region observed in the FORCE dataset using a generalized linear mixed effect model (GLMM). The model presented was the optimal model, which included random effects. Therefore, habitat was categorized as a fixed factor, and country and site were random variables, with site nested within country.

<table>
<thead>
<tr>
<th>FORCE: Coral cover</th>
<th>Estimate</th>
<th>SE</th>
<th>Z-value</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fixed effects</td>
<td>Intercept</td>
<td>3.08</td>
<td>0.08</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>Habitat</td>
<td>-0.81</td>
<td>0.12</td>
<td>-6.94</td>
</tr>
<tr>
<td>Random effects</td>
<td>Site (Country)</td>
<td>0.17</td>
<td>0.41</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>Country</td>
<td>0.05</td>
<td>0.22</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>AIC: 5967.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Cross-validation of the FORCE generalized linear mixed effect model (GLMM) using the entire AGRRA dataset (1998–2011). The structure of the model was the same as Table 2; coral cover was the response variable, habitat was the fixed variable and random effects were site nested in country, and country.

<table>
<thead>
<tr>
<th>AGRRA: Coral cover</th>
<th>Estimate</th>
<th>SE</th>
<th>Z-value</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fixed effects</td>
<td>Intercept</td>
<td>5.02</td>
<td>0.07</td>
<td>66.97</td>
</tr>
<tr>
<td></td>
<td>Habitat</td>
<td>0.19</td>
<td>0.05</td>
<td>3.81</td>
</tr>
<tr>
<td>Random effects</td>
<td>Site (Country)</td>
<td>0.29</td>
<td>0.54</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>Country</td>
<td>0.08</td>
<td>0.29</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>AIC: 7835.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Consistency of results (AGGRA data)

We used AGRRA data in order to assess the consistency of the results observed in the FORCE GLMM model, therefore testing the significance of the fixed factor (habitat) and random factors (site nested in country). The entire AGRRA dataset includes a variety of habitat types, with 22% of the sites (673 in total) characterized as *Orbicella* reefs and 78% as gorgonian plains. As observed with the FORCE data (see Table 2), coral cover significantly differed between forereef habitats within the AGRRA data (GLMM; \( p = 0.0001 \)) (Table 3), with mean coral cover (±SE) on AGRRA *Orbicella* sites being 19.2 ± 0.04%, while the average at gorgonian plains was 14.8 ± 0.05%. We observed the random effects (site nested in country, country) to be significant (\( p = 0.0001 \)) in the model; standard errors of the
random effects were 0.54 and 0.29, while variances were 0.29 and 0.08 for site nested in country and country, respectively.

**Within- and between-country comparisons (FORCE data)**

The highest mean coral cover (± SE) on *Orcicella* reefs was recorded in Bonaire (33.9 ± 1.7%), Panamá (31.7 ± 1.4%), and Barbados (29.8 ± 1.7%), while the lowest was observed at Antigua (11.7 ± 1.5%), Jamaica (16.7 ± 0.9%), and St. Lucia (17.1 ± 1.4%) (Fig. 4). For gorgonian plains, the highest mean coral cover was recorded in Belize (14.6 ± 1.2%) and Dominican Republic (14.3 ± 1.8%) and the lowest cover was in St. Vincent and Grenadines (6.2 ± 1.3%), Honduras (7.0 ± 1.2%) and Costa Rica (7.7 ± 1.0%) (Fig. 4). We then assessed the variation in coral cover between the forereef habitats in individual countries where data were available (ANT, BEL, DR, HON, SVG). The random variable, site, was significant (p = 0.0001) in both the main and interaction model (Table 4). As seen in the interaction model, the difference in coral cover between the 2 forereef habitats was insignificant in Antigua and Belize (Table 4). In contrast, coral cover was significantly greater on *Orcicella* reefs when compared to gorgonian plains in Dominican Republic, Honduras and St. Vincent and the Grenadines (Table 4). Coral cover on *Orcicella* reefs in Honduras and St. Vincent and Grenadines was 19.9 and 15.9% greater, respectively, than cover on gorgonian plains (Fig. 4).

**Table 4. Effects of forereef habitat (Habitat), country (see Table 1 for abbreviations) and interaction of habitat (Gorg = gorgonian plains, Orb = *Orcicella* reefs) and country on coral cover in individual countries where data were available using a generalized linear mixed effect model (GLMM) and as observed in the FORCE dataset. Country and habitat were categorized as fixed factors and site as a random variable. Bold indicates significant interaction terms.**

<table>
<thead>
<tr>
<th>FORE: Coral cover</th>
<th>Estimate</th>
<th>SE</th>
<th>Z-value</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fixed effects</td>
<td>Intercept</td>
<td>2.64</td>
<td>0.18</td>
<td>14.65</td>
</tr>
<tr>
<td></td>
<td>BEL</td>
<td>0.53</td>
<td>0.20</td>
<td>2.60</td>
</tr>
<tr>
<td></td>
<td>DR</td>
<td>0.54</td>
<td>0.21</td>
<td>2.58</td>
</tr>
<tr>
<td></td>
<td>HON</td>
<td>0.51</td>
<td>0.20</td>
<td>2.47</td>
</tr>
<tr>
<td></td>
<td>SVG</td>
<td>0.28</td>
<td>0.20</td>
<td>1.38</td>
</tr>
<tr>
<td></td>
<td>Habitat</td>
<td>−0.72</td>
<td>0.12</td>
<td>−5.92</td>
</tr>
</tbody>
</table>

**Random effects**

AIC=2979.3

| Fixed effects     | Intercept| 2.39| 0.20    | 11.68   | <0.0001 |
|                   | Gorg × ANT| −0.21| 0.29 | −0.73   | 0.464   |
|                   | Orb × BEL| 0.49| 0.26    | 1.93    | 0.054   |
|                   | Gorg × BEL| 0.25| 0.24    | 1.03    | 0.303   |
|                   | Orb × DR| 0.73| 0.23    | 3.12    | 0.002   |
|                   | Gorg × DR| 0.24| 0.29    | 0.85    | 0.394   |
|                   | Orb × HON| 0.89| 0.23    | 3.89    | 0.0001  |
|                   | Gorg × HON| −0.51| 0.30 | −1.74   | 0.082   |
|                   | Orb × SVG| 0.63| 0.23    | 2.77    | 0.006   |
|                   | Gorg × SVG| −0.62| 0.30 | −2.09   | 0.037   |

**Random effects**

AIC=2967.4

**DISCUSSION**

Our Caribbean-wide analysis shows that combining habitats that are characterized by low coral cover (gorgonian plain) with those habitats that have significant reef development (*Orcicella* reef), can produce biased estimates of coral cover. In both the FORCE and AGRRA datasets, coral cover at *Orcicella* reefs was significantly different and greater than at gorgonian plains. Combining habitats resulted in an underestimation of mean coral cover by 12%, when using consistent methods and surveyors. Since *Orcicella* sites dominated the FORCE dataset, the overall cover (±SE) on both habitats was modestly high at 20% (±0.5). Therefore, when combining results from multiple studies, the exact value...
would depend on the relative abundance of *Orbicella* reefs vs. gorgonian plains in the particular study. Our overall estimates of coral cover on *Orbicella* reefs were based on extensive sampling from multiple countries in the Caribbean. However, we did not sample all physical environments in the Caribbean (Chollett et al. 2012) and therefore the regional representativeness of the dataset is impossible to assess.

Gardner et al. (2003) estimated 10% coral cover for 263 reefs in the Caribbean in 2001, but less than 6% of the *Orbicella* sites surveyed in the FORCE study fell at or below this value. The Gardner et al. (2003) meta-analysis included data that was collected by different observers using contrasting monitoring methods and at different reef zones/habitats, depths, and collection times. After identifying the habitats included in Gardner et al. (2003), we found that 77% of sites were forereef habitats, and based on wave exposure regime (Chollett & Mumby 2012), only about 27% of the forereef sites were probably located on *Orbicella* reefs (Fig. 5), suggesting the inclusion of multiple habitats as a possible cause of low coral cover reported in this region-wide study. In addition, some of the sites included in the meta-analysis could have been collected in *Acropora palmata* zones which have suffered large declines from white band disease. The differences in coral cover observed cannot be explained by coral recovery because such trends are scarce in the region (Connell 1997, Roff & Mumby 2012). An alternative explanation of the discrepancy could be geographic bias; however, this seems unlikely to be the main cause of the disparity because our dataset included countries with some of the lowest coral cover in the Caribbean, such as Jamaica.

The most recent regional assessment of coral reef conditions was carried out by Jackson et al. (2014). In support of our findings, their regional coral cover assessment was also considerably higher than the coral cover reported in the Gardner et al. (2003) study. Overall, our coral cover values do not vary far from the values reported in 2014. There are a few exceptions: for example, the coral cover on Barbados and Panamá in our study was double the value reported by Jackson et al. (2014). Disparities between studies can be expected given the differences of locations, time, depth and reef habitat of sites surveyed. However, both studies agree that the reefs in Bonaire are in a better condition compared to those of other countries with highly degraded reefs, such as Antigua, Jamaica, St. Lucia and Belize.

Many reefs in the Caribbean are degraded. However, on 94% of the *Orbicella* sites, mean coral cover was well above 10%, a putative level below which reef accretion might fail to outweigh erosion (Perry et al. 2013), while others suggest that positive carbonate budgets are still feasible throughout this century provided that local and global stressors are managed and where coral cover is higher than ~20% (Kennedy et al. 2013). Taking a 20% coral cover threshold, 64% of the *Orbicella* reef sites surveyed meet this target, implying that there is considerable scope for continued ecosystem functioning in the Caribbean. How-

![Fig. 5. (a) Distribution of sites assessed by Gardner et al. (2003) per habitat type, (b) frequency distribution of the probability of occurrence of *Orbicella* reefs within forereef habitats, according to a probability threshold of wave exposure of 0.4 identified by Chollett & Mumby (2012)](image)
ever, there are challenges in minimizing both local and global stressors, while also accommodating the resource needs of coastal human populations.

Distinguishing between forereef habitats before any analysis is necessary to report a baseline for future research or monitoring, or to provide data for comparison with earlier surveys. To reduce the likelihood of habitat confounding effects in the future, habitat descriptors provided by in situ assessments or new mapping techniques could be incorporated into monitoring programs such as AGRRA and future regional analyses of reef condition in the basin.

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