Influence of dietary specialization and resource availability on geographical variation in abundance of butterflyfish

Rebecca J. Lawton & Morgan S. Pratchett

ARC Centre of Excellence for Coral Reef Studies, James Cook University, Townsville QLD, 4811, Australia

Keywords
Chaetodontidae, coral reef, distribution, local abundance, niche breadth, vulnerability.

Correspondence
Rebecca J. Lawton, ARC Centre of Excellence for Coral Reef Studies, James Cook University, Townsville QLD, 4811, Australia.
Tel: +61 7 47816058;
Fax: +61 7 47816722;
E-mail: Rebecca.Lawton@my.jcu.edu.au

Submitted in part by a Queensland Government Smart State Ph.D. Scholarship and grants from the Wildlife Preservation Society of Australia, Project Aware, and the Australian Coral Reef Society to R. J. L.

Received: 29 January 2012; Revised: 5 March 2012; Accepted: 7 March 2012

Ecology and Evolution 2012; 2(7): 1347–1361
doi: 10.1002/ece3.253

Abstract

Empirical evidence indicates that both niche breadth and resource availability are key drivers of a species' local abundance patterns. However, most studies have considered the influence of either niche breadth or resource availability in isolation, while it is the interactive effects that are likely to influence local abundance. We examined geographic variation in the feeding ecology and distribution of coralfish feeding butterflyfish to determine the influence of dietary specialization and dietary resource availability on their local abundance. Dietary composition and abundance of five butterflyfish and coral dietary resource availability were determined at 45 sites across five locations (Lizard Island and Heron Island, Great Barrier Reef; Kimbe Bay, Papua New Guinea; Noumea, New Caledonia; and Moorea, French Polynesia). Multiple regression models using variables representative of total dietary resource availability, availability of specific dietary resources, and interspecific competition were used to determine the best predictors of local abundance across all sites and locations for each species. Factors influencing local abundance varied between butterflyfish with specialized and generalized diets. Dietary resource availability had the strongest influence on the abundance of Chaetodon trifascialis—the most specialized species. Local abundance of C. trifascialis was best predicted by availability of the Acropora corals that it preferentially feeds on. In contrast, abundance of generalist butterflyfish was poorly described by variation in availability of specific resources. Rather, indices of total dietary resource availability best predicted their abundance. Overall, multiple regression models only explained a small proportion of the variation in local abundance for all five species. Despite their relatively specialized diets, dietary resource availability has limited influence on the local abundance of butterflyfish. Only the most specialized species appear to be consistently limited by prey availability. Local and total abundance of species are influenced by a wide range of different factors and there is definite need to conduct independent species assessments.

Introduction

Many macroecological studies have sought to understand what determines a species' patterns of abundance (Brown 1984; Gregory and Gaston 2000). Niche breadth is often invoked as a key determinant following Brown's (1984) niche breadth hypothesis, which predicts that generalist species will be locally abundant as they are able to exploit a wide range of environmental conditions and resources, while specialist species will be locally rare (Brown 1984; Brown et al. 1995). However, the link between niche breadth and abundance remains unclear, with studies both supporting (Pyron 1999; Harcourt et al. 2002) and failing to find (Gaston et al. 1997; Gregory and Gaston 2000; Brandle et al. 2002) a relationship. These contrasting results may in part be due to resource availability—specialist species may attain high local abundance if their preferred resources are also locally abundant. Resource availability has been identified as a key determinant of local abundance patterns in multiple studies across a range of taxa (e.g., fish: Holbrook et al. 2000; birds: Tellería and Pérez-Tris 2003; mammals: Womble and Sigler 2006; bees: Roulston and Goodell 2011). But despite the recognized importance of both niche breadth and resource availability in determining local abundance (e.g., Munday 2002), most
Studies have considered these factors in isolation and the interactive effects between them remain largely unstudied. Local abundance is likely to be influenced by both niche breadth and resource availability, such that at a given location, local abundance will be determined not only by the availability of resources but also by an organism's ability to utilize those resources. As generalist species are able to utilize a wide range of resources, their local abundance should be largely unaffected by differences in the availability of specific resources, as long as total resource availability remains constant. In contrast, specialist species are expected to be limited by the availability of critical resources (Brown 1984). Accordingly, their local abundance is more likely to be correlated with the abundance of the specific resources they specialize on, rather than total resource availability (Munday 2002; Pratchett and Berumen 2008), and will therefore be more variable than that of generalist species. Indirect evidence for these predictions comes from several sources. Specialist species are more sensitive to changes in resource availability (e.g., Harcourt et al. 2002; Kotze and O’Hara 2003; Swihart et al. 2003; Julliard et al. 2004; Aitken and Martin 2008) and can also have more restricted distributions across habitats with varying resource availability (Ostergard et al. 2009) compared to generalist counterparts. Furthermore, the abundance of some specialist species can vary significantly between sites with similar levels of total resource availability, but differing availability of specific resources (Graham 2007). These findings suggest that overall indices of resource availability will not provide an accurate reflection of resource availability for specialist species, and as such, these indices may not be the best predictors of their local abundance. However, the relative importance of total resource availability versus the availability of specific resources as determinants of local abundance for specialist species has rarely been examined.

Butterflyfish (genus *Chaetodon*) provide an ideal model group to investigate the influence of resource availability on the local abundance of specialist versus generalist species. The dietary composition and level of dietary specialization of species within this genus vary considerably, ranging from generalist species such as *Chaetodon citrinellus*, which feed on a broad range of hard coral species as well as soft corals and other reef macroinvertebrates, to highly specialized species such as *C. trifascialis*, which feed on only a small number of hard coral species (Pratchett 2005; Lawton et al. 2012, in press). The composition and abundance of hard coral species, the key dietary resource of these fish, also vary markedly at both a local (e.g., between reefs within a single location) and geographical scale (Edmunds and Bruno 1996; Veron 2000; Berumen et al. 2005). There is also evidence that the local abundance of coral-feeding butterflyfish is related to the availability of coral dietary resources. Numerous studies have found a strong link between total hard coral cover and butterflyfish abundance (Bell and Galzin 1984; Bouchon-Navaro et al. 1985; Cadoret et al. 1999; Bozec et al. 2005; Pratchett and Berumen 2008; Emslie et al. 2010, but see Bell et al. 1985; Fowler 1990) and many butterflyfish have also been observed to decline in abundance following coral loss (e.g., Sano et al. 1987; McClanahan et al. 2002; Halford et al. 2004; Pratchett et al. 2006; Graham 2007). Furthermore, predation on coral-feeding butterflyfish is generally thought to be very low (Cole et al. 2008) and their local abundance tends to be fairly stable in the absence of changes in coral availability (Halford et al. 2004), implying that availability of coral dietary resources, rather than predation or recruitment-driven processes, is likely to be a primary driver of local abundance patterns for these fish. Empirical data indicate that butterflyfish with specialized diets are more susceptible to coral loss than generalist feeders (Pratchett et al. 2006; 2008b; Graham 2007), suggesting that different factors are likely to determine the local abundance of specialist and generalist butterflyfish species. These characteristics make butterflyfish ideal candidates to explore the links between ecological versatility, resource availability, and local abundance.

Coral-feeding butterflyfish have been identified as one of the most vulnerable groups of reef fish to the combined effects of ongoing global coral loss and degradation of coral reef habitats (Wilson et al. 2006; Pratchett et al. 2008b). Identifying key drivers of their local abundance has important implications for understanding how butterflyfish are likely to be impacted by climate change. Many species preferentially feed on corals from the genus *Acropora* (Pratchett 2005; Cole et al. 2008), which are themselves vulnerable to a range of disturbances on reefs (Marshall and Baird 2000; Madin and Connolly 2006; Pratchett 2010). If local abundance of butterflyfish, particularly specialist species, is also linked to these corals, then their vulnerability to global climate change is likely to be greatly increased. Although variation in butterflyfish abundance and the role of hard coral cover as a driver of local abundance patterns have been previously investigated at a number of spatial scales (e.g., geographic regions: Findley and Findley 2001; reefs: Bozec et al. 2005; physiographic reef zones: Pratchett and Berumen 2008), the influence of specific dietary resources on abundance patterns is yet to be examined. Comparisons of local abundance and coral resource availability at replicate sites within different locations will help identify the spatial scale at which these factors are influencing populations (Underwood and Chapman 1996; Hughes et al. 1999; Munday 2002).

Although a wide range of factors can potentially influence the abundance of reef fish, we focus here on the influence of coral dietary resources due to the strong reliance of many butterflyfish on corals, which are highly vulnerable to the impacts of global climate change. Therefore, the objective of this study was to investigate the influence of dietary specialization and coral dietary resource availability on the local abundance of butterflyfish. Our specific aims were to...
(1) determine the dietary composition, level of dietary specialization, and local abundance of five butterflyfish across five geographically separated locations; (2) compare the local abundance of each species to the availability of dietary resources at each location; and (3) determine the best predictors of local abundance for each species across all locations. We expected that local abundance of dietary specialists would be best predicted by the availability of their preferred coral resources, whereas the local abundance of dietary generalists would be best predicted by total dietary resource availability. Local abundance of butterflyfish may also be modified by interspecific competition and the presence of other butterflyfish species. Butterflyfish are known to aggressively defend territories and dominant competitors have been observed to restrict the access of subordinates to habitats containing preferred corals (Crosby and Reese 2005; Berumen and Pratchett 2006a). Therefore, the influence of interspecific competition was also considered as a possible determinant of local abundance.

Methods
Study sites and species
This study investigated local abundance patterns of five common and widespread species of butterflyfish—C. auriga, C. vagabundus, C. citrinellus, C. lunulatus, and C. trifascialis. Chaetodon auriga, C. vagabundus, and C. citrinellus are all facultative corallivores—consuming hard corals as well as other small motile invertebrates and soft corals; while C. lunulatus and C. trifascialis are both obligate corallivores, feeding almost exclusively on hard corals (Pratchett 2005; Cole et al. 2008; Lawton et al. In press). Sampling was conducted at five geographically separated locations throughout the Pacific: (1) Lizard Island, Northern Great Barrier Reef; (2) Heron Island, Southern Great Barrier Reef; (3) Kimbe Bay, Papua New Guinea; (4) Noumea, New Caledonia; (5) Moorea, French Polynesia (Fig. 1). These sites are separated by 1100–6600 km and distributed along known diversity gradients (Bellwood and Hughes 2001). At each location,
Dietary composition

Field observations of feeding behavior at three sites in each location were conducted to determine the dietary composition of each of the five focal species. Individual adult butterflyfish were randomly selected and followed at a distance of 2–5 m for 3 min, following Pratchett (2005). Every effort was made to ensure individual fish were not observed more than once. At least 20 observations for each species were conducted throughout the day at each site, giving a total of at least 60 observations at each location for each species. During each observation, the total number of bites taken on each of six coral taxa groupings (Acropora, Montipora, Pocillopora, Porites, Favidae, other hard corals), noncoral substrates (e.g. sand, rubble, pavement) and any other items (e.g. algae, noncoral macroinvertebrates) was recorded, following Pratchett (2005). Smith’s measure of niche breadth ($FT$) was used to determine the relative degree of dietary specialization for each species at each site (Smith 1982). This measure takes into account resource availability and is less sensitive to the use of rare resources compared to other niche breadth measures (Krebs 1999). $FT$ is a standardized measure, ranging from 0 (most specialized) to 1 (least specialized), therefore allowing comparison of the level of specialization between sites, locations, and species.

Statistical analyses

Variation in the abundance of the five focal butterflyfish species and coral assemblage composition among locations and sites was assessed using multivariate analyses of variance (MANOVAs) comparing the mean abundance of butterflyfish and mean percent cover of corals (grouped into six taxa, plus noncoral substrates and other items) among nine sites nested within each of the five geographically separated locations. Abundance data were log$_{10}$ transformed and coral cover data were arc-sine transformed to satisfy assumptions of multivariate homogeneity and normality. Pillai’s trace statistic was used to determine the significance of MANOVA results. Where significant, variation in butterflyfish abundance and coral composition at each site were displayed using canonical discriminant analyses (CDA). To assist with interpretation of the CDA, structural coefficients of the butterflyfish species and coral taxa were plotted as vectors to indicate the predominant species and taxa at each site. To explore interspecific differences in the relative importance of different spatial scales of comparison, variance in the abundance of each focal species was partitioned among locations, sites, and transects, using the mean squares ratios of univariate $F$ values from independent nested ANOVAs. To explore the relationship between niche breadth and local abundance, the average niche breadth of each species was calculated for each location and plotted against the average abundance of each species in each location. Due to the small number of datapoints, a formal quantitative analysis of this relationship was not undertaken.

Separate multiple linear regressions using a forward stepwise method were run for each of the five focal species to identify factors significantly contributing to their local abundance. Data collected on each transect were treated as an individual replicate. An initial multiple regression analysis was run for each species with their abundance as the dependent variable and a standard set of predictor variables entered into the model simultaneously. Predictor variables with nonsignificant beta coefficients in this initial model were discarded. The model was then rerun using the remaining predictor variables that were entered hierarchically in order of their decreasing contribution to the initial model. Only predictor variables resulting in a significant change in the $R^2$ value on this second model (based on an $F$-ratio test conducted in SPSS) were retained and used in the final model. Residual plots, homogeneity tests, and the Durbin–Watson
The abundance of the five focal species was highest at Lizard Island (8.2 fish per transect ± 0.4 SE) and was lowest in Papua New Guinea (2.8 fish per transect ± 0.4 SE). Abundance of the five focal species varied significantly among locations (nested MANOVA, Pillai’s trace = 1.019, df = 20,716, P < 0.001) and sites nested within locations (nested MANOVA, Pillai’s trace = 1.8768, df = 200,900, P < 0.001). There was some partitioning of site centroids in the CDA; however, there was considerable overlap of site centroids from different locations, inferring that the butterflyfish assemblages were generally similar between sites and locations (Fig. 3a). Structural coefficients indicated that sites in French Polynesia and Lizard Island were characterized by a high abundance of C. vagabundus and C. lunulatus (Fig. 3a). Variance components indicated that for all species except C. trifascialis, most of the variation in abundance was attributable to variation among locations, rather than sites or transects. In contrast, variation in abundance was similar at both sites and locations for C. trifascialis (Fig. 4).

Coral composition

Both total hard coral cover and hard coral species diversity (measured by the Shannon–Weiner J’ index) were highest in New Caledonia (coral cover: 53.2% ± 2.9 SE, species diversity: 0.26 ± 0.01 SE), but the total number of different hard coral species recorded on each transect was highest in Papua New Guinea (18.1 species ± 0.8 SE) (Fig. 2c and d). Total hard coral cover, hard coral species diversity, and the number of hard coral species detected on each transect were all lowest in French Polynesia (coral cover: 25.5% ± 2.0 SE, species diversity: 0.16 ± 0.01 SE, coral species detected: 4.7 ± 0.3 SE). Coral assemblage composition varied significantly between locations (nested MANOVA, Pillai’s trace = 2.634, df = 32,704, P < 0.001) and sites nested within locations (nested MANOVA, Pillai’s trace = 4.538, df = 320,1440, P < 0.001). The groupings of site centroids in the CDA

### Table 1. Predictor variables used in multiple regression analyses for five species of butterflyfish.

| Species          | Dietary categories                     | Significant variables                          | Final model                                      |
|------------------|----------------------------------------|------------------------------------------------|------------------------------------------------|
| C. auriga        | Noncoral substrate, Acropora, other    | Acropora                                        | –1                                              |
|                  | hard corals, other                      |                                                |                                                 |
| C. vagabundus    | Noncoral substrate, Acropora,          | Pocillopora, hard coral species diversity      | Pocillopora, hard coral species diversity       |
|                  | Pocillopora, Montipora, Porites, other |                                                |                                                 |
| C. citrinellus   | Noncoral substrate, Acropora,          | Number of hard coral species, hard              | Number of hard coral species, total coral       |
|                  | Pocillopora, Montipora, Porites,       | coral species diversity, total coral            | coral cover/abundance congenerics               |
|                  | Favidae, other hard corals             |                                                |                                                 |
| C. lunulatus     | Noncoral substrate, Acropora,          | Noncoral substrate, total coral cover/abundance| Noncoral substrate, total coral cover/abundance |
|                  | Pocillopora, Montipora, Porites,       | congenerics, number coral species              | congenerics, number coral species               |
|                  | Favidae, other hard corals             |                                                |                                                 |
| C. trifascialis  | Acropora, Pocillopora, Montipora       | Acropora, total coral cover/abundance congenerics, | Acropora, total coral cover/abundance congenerics |

*Final model was not significant.*

Data displayed are any dietary items comprising more than 1% of total diet across all locations (dietary categories), significant predictor variables in the initial model (significant variables), and predictor variables included in the final model (final model). Significant variables are listed in order of their decreasing contribution to the initial model. See Methods section for more details.
indicated that coral assemblages at each location were more distinct than butterflyfish assemblages (Fig. 3b). Fewer site centroids from different locations overlapped with each other in the coral CDA and, in contrast to the butterflyfish CDA, centroids from Lizard Island and French Polynesia sites were completely separated from each other. Structural coefficients indicated that sites in French Polynesia were characterized by a high abundance of noncoral substrates, reflecting the low total hard coral cover at this location. Coral assemblages at Heron Island, New Caledonia, and Lizard Island were dominated by Acropora corals, with this taxon comprising more than 74%, 63%, and 56%, respectively, of the total hard coral cover at these locations. French Polynesia sites were dominated by Porites and Montipora corals—these two taxa accounting for >80% of the total coral cover in this location. In contrast, cover of different coral taxa was variable at sites in Papua New Guinea, with no one taxa dominating assemblages in this location.
Dietary composition and specialization

Feeding observations were completed for a total of 1506 individual fish (see Table A1). All butterflyfish studied fed from the surface of live corals, but the proportional feeding on corals versus noncoral substrates varied greatly. Chaetodon auriga took at least 85% of all bites on noncoral substrates at each location (see Table A1). Chaetodon vagabundus also fed predominately on noncoral substrates, taking at least 90% of all bites on this category at Lizard Island, Heron Island, and French Polynesia, and more than 60% of all bites on this category in New Caledonia and Papua New Guinea, respectively (see Table A1). Hard corals contributed significantly to the diet of C. vagabundus in Papua New Guinea, accounting for 32.0% (±5.2 SE) of all bites, most of which were taken on Montipora, Pocillopora, and Porites corals. Both C. auriga and C. vagabundus had relatively high niche breaths (Fig. 5), indicating that they were both generalist feeders. Chaetodon citrinellus fed across a broad range of hard coral taxa, noncoral substrates, and other dietary items, and diet was variable among locations (see Table A1). Between 11% and 81% of all bites were taken on noncoral substrates at each site, with the majority of bites at New Caledonia and French Polynesia taken on this category (53.3 ± 9% S.E. and 60.4 ± 18% SE, respectively). Diet was dominated by Acropora corals at Heron Island, with roughly half of all bites on hard corals taken on this taxon, while at Lizard Island Pocillopora corals accounted for nearly 40% of all bites on hard corals on average. In contrast, in both New Caledonia and Papua New Guinea, bites were spread reasonably evenly between all hard coral resource categories. Chaetodon citrinellus was the most generalized of all species, with a high-dietary niche breadth in...
Determinants of Abundance in Butterflyfish

R. J. Lawton & M. S. Pratchett

Corals as shown previously in Lawton is highly con-

C. vagabundus corals, its C. lunulatus C. auriga corals (at least 80% of C. trifascialis was positively correlated with the availability of R. Acropora C. trifascialis C. citrinellus C. auriga corals, while the abundance of (Table 2). Overall regression models and in-

C. trifascialis was the most specialized of all species, with a low to moderate niche breadth in each location (Fig. 5).

Niche breadth and local abundance

Plots of the average niche breadth and local abundance of each species at each of the five study locations were variable in pattern. Overall, there was no strong trend toward a positive or negative relationship between these factors (Fig. 6).

Regression models

Multiple linear regression analyses indicated that the local abundance of each of the five focal species was explained by a different set of variables. Availability of specific resource categories were significant predictors of abundance for three of the study species (Tables 1 and 2), but not for C. auriga or C. citrinellus. In particular, the abundance of the dietary specialist C. trifascialis was best predicted by Acropora corals, its preferred coral prey (Table 2). The abundance of C. vagabundus was positively correlated with the availability of Pocillopora corals, while the abundance of C. lunulatus was negatively correlated with the availability of noncoral substrates (Table 2). Our index of interspecific competition (the ratio of total hard coral cover to total abundance of all congeneric butterflyfish) was negatively correlated with the abundance of C. citrinellus, C. lunulatus, and C. trifascialis, while indices of coral species richness and diversity were a significant predictor of abundance for C. vagabundus, C. citrinellus, and C. lunulatus (Table 2). Overall regression models and individual predictor variables were highly significant for all species except C. auriga, however, the models only explained a low proportion of the variation in abundance for each species, ranging from 11.5% (adjusted $R^2$) for C. vagabundus to 20.8% for C. trifascialis (Table 3).

Discussion

This study revealed that the factors influencing local abundance varied among butterflyfish, with pronounced differences between specialist versus generalist species. Dietary resource availability had the strongest influence on abundance patterns for the most specialized species, C. trifascialis, with the final regression model explaining the highest proportion (20.8%) of variation in abundance for this species. The variance components analysis indicated that variation in the abundance of C. trifascialis among sites was as high as variation among locations, providing further support for the influence of dietary resource availability. Feeding observations conducted across five different geographical locations revealed that the that the diet of C. trifascialis is highly consistent among locations, whereby this species feeds predominantly on Acropora corals as shown previously in Lawton et al. (In press). Accordingly, the abundance of C. trifascialis was best predicted by a model that included the availability of Acropora corals rather than total dietary resource availability. In contrast, feeding observations indicated that C. auriga, C. vagabundus, C. citrinellus, and C. lunulatus were all dietary generalists, and also altered their diets among locations in response to differing availability of certain prey (Lawton et al. In press), across the five study locations. In contrast to C. trifascialis, the abundance of three of these species was best predicted by regression models that included indices of total dietary resource availability (coral species diversity/richness) and variation in their abundance among sites was much lower than variation among locations. However, the low proportion of variance explained by the multiple regression models for all five species indicates that the relationship between dietary resource availability and local abundance is not strong and suggests that other factors are likely to have an important influence on the local abundance of these butterflyfish.

Contrary to numerous studies that have found a strong positive relationship between total hard coral cover and butterflyfish abundance (e.g., Bell and Galzin 1984; Bouchon-Navaro et al. 1985; Cadoret et al. 1999; Bozec et al. 2005; Pratchett and Berumen 2008; Emslie et al. 2010), our results indicate that total coral cover is not an important predictor of abundance for individual butterflyfish species. There are several possible reasons why we did not find the same relationship between abundance and coral cover as these previous studies. In contrast to the current study, most previous studies have only considered the influence of coral cover on the abundance of the entire butterflyfish assemblage or specific feeding guilds (e.g., obligate corallivores). However, relationships between the abundance of individual species and coral cover are likely to vary from that of the butterflyfish assemblage due to community-level interactions. Biogeographical studies have shown that the abundance of individual butterflyfish species is negatively related to the abundance of the total butterflyfish community (Findley and Findley 2001), suggesting that factors such as interspecific competition and density compensation can strongly influence the abundance of individual species independently of total coral cover. The relationship between total coral cover and the abundance of
individual butterflyfish species has only been investigated by Pratchett and Berumen (2008), who found a strong positive correlation for all obligate corallivore species at the scale of a single reef, Lizard Island. However, the total coral cover at sites sampled in their study ranged from roughly 2% to 30%. In comparison, total coral cover at sites in the current study ranged from 6% to 80%. The absence of total coral cover as a significant variable in our multiple regression models suggests that although total coral cover and abundance of individual butterflyfish appear to be linearly related at low to moderate levels of coral cover (e.g., Pratchett and Berumen 2008), the overall relationship is more likely to be asymptotic, such that further increases in coral cover above a certain threshold (e.g., 40% coral cover) have limited influence on the abundance of individual butterflyfish.

The relatively weak effect of dietary resource availability on variation in abundance of all five coral-feeding butterflyfish is also in contrast to previous studies of other coral reef fish, which have shown that that resource availability (specifically, coral cover) is a major determinant of abundance patterns for individual species at geographic spatial scales (e.g., gobies: Munday 2002; damselfish: Holbrook et al. 2000). Our
Determinants of Abundance in Butterflyfish

R. J. Lawton & M. S. Pratchett

Table 2. Coefficients of multiple regression models for four species of butterflyfish.

| Species            | Final predictors          | B     | SE B  | β     |
|--------------------|---------------------------|-------|-------|-------|
| C. vagabundus      | Pocillopora               | 2.354 | 0.437 | 0.344***|
|                    | Number of coral species   | −0.442| 0.181 | −0.156* |
| C. citrinellus     | Total coral cover/abundance congenerics | −1.201| 0.292 | −0.256***|
|                    | Total coral cover/abundance congenerics | −0.897| 0.252 | −0.226***|
| C. lunulatus       | Noncoral substrate        | −0.394| 0.086 | −0.306***|
|                    | Total coral cover/abundance congenerics | −0.897| 0.252 | −0.226***|
| C. trifascialis    | Acropora                  | 0.590 | 0.077 | 0.485***|
|                    | Total coral cover/abundance congenerics | −0.769| 0.201 | −0.242***|

The unstandardized beta coefficients (B), their standard errors (SE B), and the standardized beta coefficients (β) for the predictor variables included in the final regression model for each species are presented. *P < 0.05; **P < 0.01; ***P < 0.001.

Table 3. Final multiple regression results for abundance of five species of butterflyfish.

| Species       | Adjusted R² | Sum of Squares | df | Mean square | F      | Significance |
|---------------|-------------|----------------|----|-------------|--------|--------------|
| C. auriga     | −0.004      | 0.00           | 1,224 | 0.000 | 0.020 | <0.001     |
| C. vagabundus | 0.115       | 1.546          | 2,224 | 0.773 | 15.550 | <0.001     |
| C. citrinellus| 0.163       | 4.194          | 2,224 | 2.097 | 22.751 | <0.001     |
| C. lunulatus  | 0.159       | 3.491          | 3,224 | 1.164 | 15.114 | <0.001     |
| C. trifascialis| 0.208      | 3.212          | 2,224 | 1.606 | 30.340 | <0.001     |

Contrasting findings could be a result of several factors. For some of these butterflyfish, it is possible that the resource categories we used were too poorly resolved to effectively assess variations in the abundance of key dietary components. This is likely to be the case for species such as *C. auriga*, for which small motile invertebrates comprise a significant proportion of diet (Anderson et al., 1981; Pratchett 2005). It was assumed that bites on non-coral substrates not obviously occupied by any macro-invertebrates were targeting these organisms as has been shown previously (Anderson et al. 1981). As their availability is extremely difficult to quantify in the field, our resource category of “non-coral substrates” was used as a proxy. However, the availability of non-coral substrates may not capture the true availability of small motile invertebrates. Obligate coral-feeding species, such as *C. trifascialis* and *C. lunulatus*, exhibit strong preferences for specific coral species (Pratchett 2005). Availability of these individual coral species may vary significantly from the availability of coral taxa groupings that were used in this study. Consequently, categorization of dietary resources at a finer taxonomic resolution may be necessary to reflect true dietary resource availability for these butterflyfish.

A further possibility is that availability of dietary resources is not the primary driver of local abundance at a geographic scale for these butterflyfish. The inclusion of our variable representing interspecific competition (the ratio of total coral cover to the total abundance of all congeneric butterflyfish) in final regression models indicates that competitive interactions are likely to influence local abundance. Competitive interactions may influence the range of habitats used by a species and asymmetric competition between species can lead to the exclusion of subordinate competitors from mutually preferred habitats (Connell 1983; Abramsky et al. 1990; Youn 2004; Bonin et al. 2009). Although the overall predictive power of our final regression models was low, the highly significant negative correlation between our interspecific competition variable and abundance for three of the five focal species is supported by previous studies that have shown that abundance of individual species may increase in the absence of interspecific competitors (Schmitt and Holbrook 1990; Robertson 1996). Competitive interactions are likely to have the strongest influence on the local abundance of obligate coral-feeding butterflyfish as this variable was included in regression models for *C. citrinellus*, *C. lunulatus*, and *C. trifascialis—the three species with the highest proportional consumption of hard corals. In agreement with previous observational studies of competitive interactions between butterflyfish (Crosby and Reese 2005; Berumen and Pratchett 2006a), our results suggest that defense of preferred coral resources by dominant competitors may be negatively influencing local abundance patterns of coral-feeding butterflyfish.

Larval supply and recruitment may also be a key determinant of butterflyfish abundance at large spatial scales. Both of these factors are widely acknowledged to be highly variable.
across both space and time in marine populations (Doherty 1991, 2002; Caley et al. 1996). Differences in adult abundance have been shown to be strongly related to prior levels of recruitment for some reef fish (Victor 1983, 1986; Holbrook et al. 2000) and variable larval supply may be contributing to the local patterns of abundance for the butterflyfish considered here (Bell et al. 1985; Pratchett and Berumen 2008). A further possibility is that both larval supply and resource availability may be constraining local abundance, as appears to be occurring in some other reef fish populations (e.g., Forrester 1995; Schmitt and Holbrook 1999; Holbrook et al. 2000), such that recruit abundance is initially determined by larval supply then consequently regulated by resource availability. Juvenile obligate coral-feeding butterflyfish are consistently found in close association with hard coral colonies in the field (Cole and Pratchett 2011) and the distribution of several species corresponds closely to the distribution of preferred coral microhabitats (Pratchett et al. 2008a). These observations suggest that the availability of coral resources for both settlement habitat and food strongly influences the abundance of juvenile coral-feeding butterflyfish and may override initial abundance patterns established at settlement (e.g., Booth 2002). Further research is necessary to determine both the influence of resource availability on the abundance and distribution of juvenile butterflyfish, and the relationship between juvenile and adult abundance.

Theory predicts that the abundance of specialist species should be lower than that of generalist counterparts, due to increased limits imposed by a restricted set of critical resources (Brown 1984). Our findings confirmed that dietary resource availability has a greater influence on the abundance of the most specialized species (C. trifascialis), but this species was generally more abundant, not less abundant, than generalist counterparts. This is probably due to the high abundance of Acropora corals at most locations. Consequently, it should not be assumed that specialists will always be rarer than generalists. Chaetodon trifascialis is frequently one of most abundant butterflyfish throughout its geographic range (Jones et al. 2002), and is generally only rare where there has been systematic depletion of Acropora corals (Berumen and Pratchett 2006b). Our results highlight the need to be wary of assigning specialized species high vulnerability status due to an assumed low abundance and resource dependence without conducting independent assessments.

Supporting this, recent research has demonstrated that C. trifascialis has a greater level of feeding plasticity and therefore a greater capacity to respond to coral loss than previously assumed (Lawton et al. In press), while genetic evidence suggests there is a high potential for C. trifascialis to recover from population declines (Lawton et al. 2011). These findings indicate that the vulnerability of C. trifascialis to coral loss on reefs is likely to be lower than has been previously assumed.

Acknowledgments

We thank A. Cole, M. Berumen, D. McCowan, K. Chong-Seng, M. Trapon, V. Messmer, and D. Coker for assistance with field work. We also thank the staff of Heron Island Research Station, Lizard Island Research Station, Mahonia na Dari and Richard B. Gump Research Station for logistical support. This research was funded in part by a Queensland Government Smart State Ph.D. Scholarship and grants from the Wildlife Preservation Society of Australia, Project Aware, and the Australian Coral Reef Society to R. J. L.

References

Abramsky, Z., M. L. Rosenzweig, B. Pinshow, J. S. Brown, B. Kotler, and W. A. Mitchell. 1990. Habitat selection: an experimental field test with two gerbil species. Ecology 71:2358–2369.
Aitken, K. E. H., and K. Martin. 2008. Resource selection plasticity and community responses to experimental reduction of a critical resource. Ecology 89:971–980.
Anderson, G. R. V., A. H. Ehrlich, P. R. Ehrlich, J. D. Roughgarden, B. C. Russell, and F. H. Talbot. 1981. The community structure of coral reef fishes. Am. Nat. 117:476–495.
Bell, J. D., and R. Galzin. 1984. Influence of live coral cover on coral-reef fish communities. Mar. Ecol. Prog. Ser. 15:265–274.
Bell, J. D., M. Harmelin-Vivien, and R. Galzin. 1985. Large scale spatial variation in abundance of butterflyfishes (Chaetodontidae) on Polynesian reefs. Pp. 421–426 in Proceedings of the Fifth International Coral Reef Congress. 27 May–1 June 1985, Tahiti.
Bellwood, D. R., and T. P. Hughes. 2001. Regional-scale assembly rules and biodiversity of coral reefs. Science 292:1532–1535.
Berumen, M. L., and M. S. Pratchett. 2006a. Effects of resource availability on the competitive behaviour of butterflyfishes (Chaetodontidae). In Proceedings of the 10th International Coral Reef Symposium, Okinawa, Japan.
Berumen, M. L., and M. S. Pratchett. 2006b. Recovery without resilience: persistent disturbance and long-term shifts in the structure of fish and coral communities at Tiahura Reef, Moorea. Coral Reefs 25:647–653.
Berumen, M. L., M. S. Pratchett, and M. I. McCormick. 2005. Within-reef differences in diet and body condition of coral-feeding butterflyfishes (Chaetodontidae). Mar. Ecol. Prog. Ser. 287:217–227.
Bonin, M. C., M. Srinivasan, G. R. Almany, and G. P. Jones. 2009. Interactive effects of interspecific competition and microhabitat on early post-settlement survival in a coral reef fish. Coral Reefs 28:265–274.
Booth, D. J. 2002. Distribution changes after settlement in six species of damselfish (Pomacentridae) in One Tree Island lagoon, Great Barrier Reef. Mar. Ecol. Prog. Ser. 226:157–164.
Determinants of Abundance in Butterflyfish

R. J. Lawton & M. S. Pratchett

Bouchon-Navaro, Y., C. Bouchon, and M. L. Harmelin-Vivien. 1985. Impact of coral degradation on a Chaetodontid fish assemblage (Moorea, French Polynesia). Proc. 5th Int. Coral Reef Symp. 5:427–432.

Bozec, Y. M., S. Doledec, and M. Kulbicki. 2005. An analysis of fish-habitat associations on disturbed coral reefs: chaetodontid fishes in New Caledonia. J. Fish Biol. 66:966–982.

Brandley, M., A. Prinzing, R. Pfeifer, and R. Brandl. 2002. Dietary niche breadth for Central European birds: correlations with species-specific traits. Evol. Ecol. Res. 4:643–657.

Brown, J. H. 1984. On the relationship between abundance and distribution of species. Am. Nat. 124:255–279.

Brown, J. H., D. W. Mehlman, and G. C. Stevens. 1995. Spatial variation in abundance. Ecology 76:2028–2043.

Cadoret, L., M. Adjeroud, and M. Tsuchiya. 1999. Spatial distribution of chaetodontid fish in coral reefs of the Ryukyu Islands, southern Japan. J. Mar. Biol. Assoc. UK 79:725–735.

Caley, M. J., M. H. Carr, M. A. Hixon, T. P. Hughes, G. P. Jones, and B. A. Menge. 1996. Recruitment and the local dynamics of open marine populations. Ann. Rev. Ecol. Syst. 27:477–500.

Cole, A., and M. Pratchett. 2011. Effects of juvenile coral-feeding butterflyfishes on host corals. Coral Reefs 30:623–630.

Cole, A. J., M. S. Pratchett, and G. P. Jones. 2008. Diversity and functional importance of coral-feeding fishes on tropical coral reefs. Fish Fish. 9:286–307.

Connell, J. H. 1983. On the prevalence and relative importance of interspecific competition—evidence from field experiments. Am. Nat. 122:661–696.

Crosby, M. P., and E. S. Reese. 2005. Relationship of habitat stability and intra-specific population dynamics of an obligate corallivore butterflyfish. Aquat. Conserv. Mar. Freshwat. Ecosyst. 15:S13–S25.

Doherty, P. J. 1991. Spatial and temporal patterns in recruitment. Pp. 261–293 in P. F. Sale, ed. The ecology of fishes on coral reefs. Academic Press, San Diego, CA.

Doherty, P. J. 2011. Variable replenishment and the dynamics of reef fish populations. Pp. 327–358 in P. F. Sale, ed. Coral reef fishes: dynamics and diversity in a complex ecosystem. Academic Press, San Diego, CA.

Edmunds, P. J., and J. F. Bruno. 1996. The importance of sampling scale in ecology: kilometer-wide variation in coral reef communities. Mar. Ecol. Prog. Ser. 143:165–171.

Emslie, M., M. Pratchett, A. Cheal, and K. Osborne. 2010. Great Barrier Reef butterflyfish community structure: the role of shelf position and benthic community type. Coral Reefs 29:705–715.

Findley, J. S., and M. T. Findley. 2001. Global, regional, and local patterns in species richness and abundance of butterflyfishes. Ecol. Monogr. 71:69–91.

Forrester, G. E. 1995. Strong density-dependent survival and recruitment regulate the abundance of a coral reef fish. Oecologia 103:275–282.

Fowler, A. J. 1990. Spatial and temporal patterns of distribution and abundance of chaetodontid fishes at One Tree Reef, southern GBR. Mar. Ecol. Prog. Ser. 64:39–53.

Gaston, K. J., T. M. Blackburn, and J. H. Lawton. 1997. Interspecific abundance range size relationships: an appraisal of mechanisms. J. Anim. Ecol. 66:579–601.

Graham, N. A. J. 2007. Ecological versatility and the decline of coral feeding fishes following climate driven coral mortality. Mar. Biol. 153:119–127.

Gregory, R. D., and K. J. Gaston. 2000. Explanations of commonness and rarity in British breeding birds: separating resource use and resource availability. Oiks 88:515–526.

Halford, A., A. J. Cheal, D. Ryan, and D. M. Williams. 2004. Resilience to large-scale disturbance in coral and fish assemblages on the Great Barrier Reef. Ecology 85:1892–1905.

Harcourt, A. H., S. A. Coppeto, and S. A. Parks. 2002. Rarity, specialization and extinction in primates. J. Biogeogr. 29:445–456.

Holbrook, S. J., G. E. Forrester, and R. J. Schmitt. 2000. Spatial patterns in abundance of a damselfish reflect availability of suitable habitat. Oecologia 122:109–120.

Hughes, T. P., A. H. Baird, E. A. Dinsdale, N. A. Moltschaniwskyj, M. S. Pratchett, J. E. Tanner, and B. L. Willis. 1999. Patterns of recruitment and abundance of corals along the Great Barrier Reef. Nature 397:59–63.

Jones, G. P., P. L. Munday, and J. M. Caley. 2002. Rarity in coral reef fish communities. Pp. 549 in P. F. Sale, ed. Coral reef fishes: dynamics and diversity in a complex ecosystem. Academic Press, Sydney.

Julliard, R., F. Jiguet, and D. Couvet. 2004. Common birds facing global changes: what makes a species at risk? Glob. Change Biol. 10:148–154.

Kotze, D. J., and R. B. O’Hara. 2003. Species decline—but why? Explanations of carabid beetle (Coleoptera, Carabidae) declines in Europe. Oecologia 135:138–148.

Krebs, C. J. 1999. Ecological methodology, 2nd ed. Benjamin/Cummings, Menlo Park, CA.

Lawton, R. J., V. Messmer, M. Pratchett, and L. K. Bay. 2011. High gene flow across large geographic scales reduces extinction risk for a highly specialised coral feeding butterflyfish. Mol. Ecol. 20:3584–3598.

Lawton, R. J., M. S. Pratchett, and M. L. Berumen. 2012. The use of specialisation indices to predict vulnerability of coral-feeding butterflyfishes to environmental change. Oikos 121:191–200.

Lawton, R. J., A. J. Cole, M. L. Berumen, and M. S. Pratchett. Geographic variation in resource use by specialist versus generalist butterflyfishes. Ecography. doi: 10.1111/j.1600-0587.2011.07326.x. In press.

Madin, J. S., and S. R. Connolly. 2006. Ecological consequences of major hydrodynamic disturbances on coral reefs. Nature 444:477–480.

Marshall, P. A., and A. H. Baird. 2000. Bleaching of corals on the Great Barrier Reef: differential susceptibilities among taxa. Coral Reefs 19:155–163.
McClanahan, T., J. Maina, and L. Pet-Soede. 2002. Effects of the 1998 coral morality event on Kenyan coral reefs and fisheries. Ambio 31:543–550.

Munday, P. L. 2002. Does habitat availability determine geographical-scale abundances of coral-dwelling fishes? Coral Reefs 21:105–116.

Ostergard, H., P. A. Hamback, and J. Ehrlen. 2009. Responses of a specialist and a generalist seed predator to variation in their common resource. Oikos 118:1471–1476.

Pratchett, M. 2010. Changes in coral assemblages during an outbreak of Acanthaster planci; at Lizard Island, northern Great Barrier Reef (1995–1999). Coral Reefs 29: 717–725.

Pratchett, M., M. Berumen, M. Marnane, J. Eagle, and D. Pratchett. 2008a. Habitat associations of juvenile versus adult butterflyfishes. Coral Reefs 27:541–551.

Pratchett, M. S. 2005. Dietary overlap among coral-feeding butterflyfishes (Chaetodontidae) at Lizard Island, northern Great Barrier Reef. Mar. Biol. 148:373–382.

Pratchett, M. S., and M. L. Berumen. 2008. Interspecific variation in distributions and diets of coral reef butterflyfishes (Teleostei: Chaetodontidae). J. Fish Biol. 73:1730–1747.

Pratchett, M. S., S. K. Wilson, and A. H. Baird. 2006. Declines in the abundance of Chaetodon butterflyfishes following extensive coral depletion. J. Fish Biol. 69:1269–1280.

Pratchett, M. S., P. L. Munday, S. K. Wilson, N. A. J. Graham, J. E. Cinneri, D. R. Bellwood, G. P. Jones, N. V. C. Polunin, and T. R. McClanahan. 2008b. Effects of climate-induced coral bleaching on coral-reef fishes—ecological and economic consequences. Oceanography and marine biology: an annual review, Vol. 46, pp. 251–296. Crc Press-Taylor & Francis Group, Boca Raton.

Pyron, M. 1999. Relationships between geographical range size, body size, local abundance, and habitat breadth in North American suckers and sunfishes. J. Biogeogr. 26:549–558.

Robertson, D. R. 1996. Interspecific competition controls abundance and habitat use of territorial caribbean damselfishes. Ecology 77:885–899.

Roulston, T. A. H., and K. Goodell. 2011. The role of resources and risks in regulating wild bee populations. Ann. Rev. Entomol. 56:293–312.

Sano, M., M. Shimizu, and Y. Nose. 1987. Long-term effects of destruction of hermatypic corals by Acanthaster planci infestation on reef fish communities at Iriomote Island, Japan. Mar. Ecol. Prog. Ser. 37:191–199.

Schmitt, R. J., and S. J. Holbrook. 1990. Population responses of surferch released from competition. Ecology 71:1653–1665.

Schmitt, R. J., and S. J. Holbrook. 1999. Mortality of juvenile damselfish: implications for assessing processes that determine abundance. Ecology 80:35–50.

Smith, E. P. 1982. Niche breadth, resource availability, and inference. Ecology 63:1675–1681.

Swihart, R. K., T. M. Gehring, M. B. Kolozsvary, and T. E. Nupp. 2003. Responses of ‘resistant’ vertebrates to habitat loss and fragmentation: the importance of niche breadth and range boundaries. Divers. Distrib. 9:1–18.

Tellera, J. L., and J. Pérez-Tris. 2003. Seasonal distribution of a migratory bird: effects of local and regional resource tracking. J. Biogeogr. 30:1583–1591.

Underwood, A. J., and M. G. Chapman. 1996. Scales of spatial patterns of distribution of intertidal invertebrates. Oecologia 107:212–224.

Veron, J. E. N. 2000. Corals of the world. Australian Institute of Marine Science, Townsville.

Victor, B. C. 1983. Recruitment and population dynamics of a coral reef fish. Science 219:419–420.

Victor, B. C. 1986. Larval settlement and juvenile mortality in a recruitment-limited coral reef fish population. Ecol. Monogr. 56:145–160.

Wilson, S. K., N. A. J. Graham, M. S. Pratchett, G. P. Jones, and N. V. C. Polunin. 2006. Multiple disturbances and the global degradation of coral reefs: are reef fishes at risk or resilient? Glob. Change Biol. 12:2220–2234.

Womble, J. N., and M. F. Sigler. 2006. Seasonal availability of abundant, energy-rich prey influences the abundance and diet of a marine predator, the Steller sea lion Eumetopias jubatus. Mar. Ecol. Prog. Ser. 325:281–293.

Young, K. A. 2004. Asymmetric competition, habitat selection, and niche overlap in juvenile salmonids. Ecology 85:134–149.

Zar, J. H. 1999. Biostatistical analysis, 4th ed. Prentice Hall, NJ.
### Appendix

Table A1. Proportional consumption of eight dietary categories by five butterflyfish across five locations. Data are pooled across *n* replicate feeding observations for each species of butterflyfish. Location abbreviations follow Figure 1. ACRO, *Acropora*; POR, *Porites*; MON, *Montipora*; POC, *Pocillopora*; FAV, *Faviidae*; OHC, other hard corals; NCS, noncoral substrates; OTH, other.

| Location | Site          | n  | ACRO | POC  | MON | POR | FAV  | OHC  | NCS | OTH |
|----------|---------------|----|------|------|-----|-----|------|------|-----|-----|
| C. auriga |               |    |      |      |     |     |      |      |     |     |
| HI       | Blue Pools    | 20 | 0.0  | 0.0  | 0.0 | 0.0 | 0.0  | 0.0  | 0.0 | 99.3| 0.7 |
|          | North Wistari | 20 | 1.2  | 0.0  | 0.0 | 0.0 | 0.0  | 0.0  | 0.0 | 94.0| 4.8 |
|          | 2nd Point     | 21 | 0.0  | 0.0  | 0.0 | 0.0 | 0.0  | 0.0  | 0.0 | 95.8| 4.2 |
| LI       | Bird Islet    | 20 | 0.0  | 1.6  | 0.0 | 0.0 | 0.0  | 0.0  | 0.0 | 98.4| 0.0 |
|          | Lizard Head   | 20 | 0.0  | 2.1  | 0.0 | 0.0 | 0.0  | 0.0  | 0.4 | 96.3| 1.2 |
|          | South Island  | 21 | 8.7  | 2.5  | 0.4 | 0.0 | 3.7  | 0.4  |     | 83.9| 0.4 |
| NC       | Ilot Nge      | 20 | 6.5  | 2.4  | 0.0 | 0.0 | 0.0  | 0.0  | 0.0 | 86.2| 4.9 |
|          | Recif Senez   | 20 | 0.7  | 0.0  | 0.2 | 0.0 | 0.0  | 0.0  | 0.0 | 99.1| 0.0 |
|          | Seche Croissant | 20 | 0.4  | 0.0  | 1.6 | 1.2 | 0.0  | 12.5 |    | 69.1| 15.2|
| FP       | Motu Ahi      | 20 | 0.0  | 0.0  | 0.0 | 1.0 | 0.0  | 0.0  | 0.0 | 99.0| 0.0 |
|          | Temea         | 20 | 0.0  | 0.0  | 3.2 | 10.4| 0.0  | 0.0  | 0.0 | 86.4| 0.0 |
|          | Tiahura       | 20 | 0.0  | 0.0  | 1.0 | 2.1 | 0.0  | 0.0  | 0.0 | 96.9| 0.0 |
| C. vagabundus |           |    |      |      |     |     |      |      |     |     |
| HI       | Blue Pools    | 22 | 0.0  | 0.4  | 0.0 | 0.0 | 0.0  | 0.0  | 0.0 | 86.2| 13.5|
|          | North Wistari | 22 | 0.6  | 0.3  | 0.0 | 0.0 | 0.3  | 0.0  | 0.3 | 95.7| 2.8 |
|          | 2nd Point     | 22 | 0.0  | 0.4  | 0.0 | 0.0 | 0.4  | 0.4  | 0.4 | 98.3| 0.4 |
| LI       | Bird Islet    | 20 | 7.7  | 0.0  | 0.0 | 0.0 | 0.3  | 0.0  | 0.0 | 92.1| 0.0 |
|          | Lizard Head   | 31 | 0.7  | 0.5  | 0.3 | 0.2 | 0.5  | 0.0  | 0.0 | 95.9| 2.0 |
|          | South Island  | 22 | 1.5  | 0.0  | 0.0 | 0.0 | 2.0  | 1.0  |    | 94.0| 1.5 |
| PNG      | Christine     | 20 | 1.7  | 2.8  | 7.6 | 10.4| 3.8  | 0.0  |    | 62.5| 11.1|
|          | Lubaluba      | 23 | 2.0  | 10.5 | 3.9 | 7.9 | 0.3  | 2.6  |    | 70.7| 2.0 |
|          | Susans        | 21 | 4.0  | 5.3  | 21.9| 6.4 | 4.3  | 0.5  |    | 50.1| 7.5 |
| NC       | Ilot Nge      | 20 | 9.9  | 9.3  | 2.5 | 0.3 | 0.0  | 0.0  | 2.0 | 71.3| 4.8 |
|          | Recif Senez   | 20 | 0.0  | 0.0  | 1.3 | 0.0 | 0.0  | 0.0  | 0.0 | 97.1| 1.6 |
|          | Seche Croissant | 20 | 0.6  | 8.2  | 0.0 | 1.5 | 0.9  | 3.2  |    | 48.8| 36.8|
| FP       | Motu Ahi      | 21 | 0.0  | 0.5  | 3.9 | 0.5 | 0.0  | 0.0  | 0.0 | 94.7| 0.5 |
|          | Temea         | 20 | 0.0  | 2.5  | 2.5 | 5.3 | 0.0  | 0.0  | 0.0 | 89.8| 0.0 |
|          | Tiahura       | 20 | 0.0  | 0.0  | 5.0 | 1.7 | 0.0  | 0.0  | 0.0 | 93.3| 0.0 |
| C. citrinellus |           |    |      |      |     |     |      |      |     |     |
| HI       | Blue Pools    | 24 | 51.1 | 3.0  | 3.9 | 0.7 | 0.7  | 5.3  | 0.0 | 35.3| 0.2 |
|          | North Wistari | 22 | 39.8 | 8.4  | 1.1 | 0.4 | 0.0  | 6.9  |    | 35.8| 7.6 |
|          | 2nd Point     | 20 | 8.2  | 2.8  | 4.1 | 0.8 | 3.4  | 13.1 |    | 65.7| 1.8 |
| LI       | Bird Islet    | 20 | 27.3 | 31.4 | 7.5 | 1.0 | 4.1  | 4.1  |    | 21.1| 3.6 |
|          | Lizard Head   | 20 | 27.0 | 23.5 | 1.8 | 0.5 | 1.4  | 1.8  |    | 34.2| 9.8 |
|          | South Island  | 20 | 24.8 | 30.9 | 5.0 | 0.6 | 6.5  | 7.1  |    | 21.3| 3.8 |
| PNG      | Christine     | 21 | 17.3 | 22.5 | 8.6 | 19.1| 9.5  | 8.4  |    | 14.5| 4.5 |
|          | Lubaluba      | 21 | 18.3 | 5.7  | 6.6 | 5.2 | 5.4  | 0.3  |    | 57.6| 0.9 |
|          | Susans        | 20 | 24.9 | 32.4 | 21.0| 6.4 | 1.5  | 2.2  |    | 11.5| 0.0 |
| NC       | Ilot Nge      | 20 | 40.4 | 17.3 | 1.1 | 1.1 | 3.1  | 0.2  |    | 36.8| 0.0 |
|          | Recif Senez   | 20 | 27.3 | 4.6  | 0.2 | 0.0  | 0.0  | 0.0  |    | 67.2| 0.7 |
|          | Seche Croissant | 20 | 13.8 | 2.6  | 19.8| 0.2 | 0.0  | 0.2  |    | 55.9| 7.6 |
| FP       | Motu Ahi      | 20 | 0.5  | 25.1 | 45.7| 4.8 | 0.0  | 0.2  |    | 23.7| 0.0 |
|          | Temea         | 21 | 0.8  | 7.7  | 7.1 | 7.9 | 0.5  | 0.0  |    | 75.8| 0.2 |
|          | Tiahura       | 20 | 0.7  | 1.6  | 8.4 | 7.1 | 0.2  | 0.4  |    | 81.6| 0.0 |

(continued)
### Table A1. Continued

| Location | Site          | n  | ACRO | POC | MON | POR | FAV | OHC | NCS | OTH |
|----------|---------------|----|------|-----|-----|-----|-----|-----|-----|-----|
| 
| C. lunulatus |               |    |      |     |     |     |     |     |     |     |

| ACRO | POC | MON | POR | FAV | OHC | NCS | OTH |
|------|-----|-----|-----|-----|-----|-----|-----|
| Blue Pools | 21  | 35.7 | 13.5 | 9.8 | 8.1 | 3.5 | 27.8 | 1.7 | 0.0 |
| North Wistari | 24  | 73.3 | 13.3 | 5.9 | 1.3 | 1.2 | 4.7 | 0.3 | 0.0 |
| 2nd Point | 20  | 32.5 | 4.5  | 35.8 | 3.1 | 2.6 | 18.4 | 2.6 | 0.5 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |
| Christines | 21  | 23.7 | 9.2  | 16.4 | 29.1 | 3.8 | 17.0 | 0.0 | 0.8 |
| North Wistari | 24  | 73.3 | 13.3 | 5.9 | 1.3 | 1.2 | 4.7 | 0.3 | 0.0 |
| 2nd Point | 20  | 32.5 | 4.5  | 35.8 | 3.1 | 2.6 | 18.4 | 2.6 | 0.5 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |

| PNG | C. trifascialis | 
|-----|----------------|
| ACRO | POC | MON | POR | FAV | OHC | NCS | OTH |
| Bird Islet | 21  | 23.7 | 9.2  | 16.4 | 29.1 | 3.8 | 17.0 | 0.0 | 0.8 |
| North Wistari | 24  | 73.3 | 13.3 | 5.9 | 1.3 | 1.2 | 4.7 | 0.3 | 0.0 |
| 2nd Point | 20  | 32.5 | 4.5  | 35.8 | 3.1 | 2.6 | 18.4 | 2.6 | 0.5 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |
| Bird Islet | 21  | 44.2 | 22.2 | 7.9 | 7.2 | 11.4 | 5.6 | 0.5 | 0.9 |
| Lizard Head | 23  | 53.7 | 27.8 | 7.5 | 2.1 | 4.5 | 3.8 | 0.7 | 0.0 |
| South Island | 23  | 56.3 | 25.0 | 3.2 | 5.2 | 6.3 | 3.4 | 0.7 | 0.0 |

* C. auriga not present at sampling sites in PNG.