Relationships among distribution, abundance and microhabitat specialisation in a guild of coral reef triggerfish (family Balistidae)

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ABSTRACT: The potential relationship between ecological versatility and local distribution and abundance for 5 species of triggerfish was examined at Kimbe Bay, Papua New Guinea. The abundances of juvenile and adult Balistapus undulatus, Melichthys vidua, Rhinecanthus verrucosus, Sufflamen bursa and S. chrysopterus were quantified in a range of habitats along a typical coral reef profile. Four of the 5 species displayed distinct and relatively narrow distributions across the reef profile, with the fifth species, B. undulatus, being broadly distributed across all zones and depths, and the most abundant species. For each species, the spatial distribution of juveniles closely matched that of adults and juvenile densities were greater in species with more abundant adults. A detailed description of depth distributions of individuals indicated that shallow species had narrower depth ranges. In terms of micro-habitat use, B. undulatus was the most generalised species, occupying all the microhabitats that were identified. The other 4 species were specialised to varying degrees on different microhabitats. Results from this study provide the first detailed description of patterns of distribution and abundance, habitat use and ecological versatility in triggerfish. Distribution and abundance could partially be explained by differences in the degree to which each species is specialised, both in terms of depth and micro-habitat selectivity. Hence, versatility in depth and microhabitat use may play an important role in determining the local distribution and relative abundance of coral reef fishes.

KEY WORDS: Abundance · Distribution · Ecological specialisation · Habitat versatility · Resource availability · Coral reef fish · Balistidae

INTRODUCTION

Ecological communities are made up of species that vary in the degree to which they are specialised on the available resources (Fox & Morrow 1981, Futuyma & Moreno 1988, McNally 1995). The factors affecting the degree of specialisation and the consequences of differing levels of ecological versatility have received considerable theoretical attention (e.g. McNaughton & Wolf 1970, McNally 1995, Morris 1996, Robinson & Wilson 1998). Generalists potentially have access to greater amounts of resources and consequently may achieve a greater local abundance, a wider distribution among local habitats (i.e. over the reef profile) and a greater geographic range (McNaughton & Wolf 1970, Brown 1984, Hengeveld 1990, McNally 1995, Hughes 2000). Specialists on the other hand, may be able to use certain resources more efficiently and may out-compete generalists in the acquisition of these resources (Emlen & Oring 1977, Futuyma & Moreno 1988). The ecological factors that may affect the relative costs and benefits of these 2 extremes are numerous with the theoretical relationship between resource specialisation, and the local distribution and abundance of species only recently tested (Hughes 2000).

Ecological versatility has been defined as ‘the degree to which organisms can fully exploit the available resources in their local environment’ (McNally 1995).
In terms of habitats occupied, versatile species are expected to be ubiquitous, occupying a broad range of habitats, while specialised species are confined to a limited number of habitats. Information on the patterns and consequences of ecological versatility in coral reef fish communities is particularly scarce (Jones et al. 2002). These highly diverse communities appear to be comprised of the full spectrum of species, from specialists closely associated with a few coral or other invertebrate species (Arvedlund & Nielsen 1996, Munday et al. 1997, Munday & Jones 1998) to generalists capable of occupying a wide range of biotic and abiotic habitat types (Green 1996, Symms & Jones 2000).

Relationships between reef fishes and habitat characteristics are best known for speciose families such as the gobies (e.g. Munday et al. 1997), damselfishes (Meekan et al. 1995), butterflyfishes (e.g. Hourigan 1989) and wrasses (e.g. Green 1996). Many species exhibit distinct patterns of habitat selection at settlement (Ormond et al. 1996). They may also display ontogenetic changes in resource utilisation, which may lead to major differences in the distribution and abundance of different life history stages (Lirman 1994, Green 1996). Hence, ontogenetic trends in the degree of habitat versatility may be critical in evaluating the processes affecting the distribution and abundance of adults.

Little is known of the factors affecting the habitat use and the distribution and abundance of triggerfishes (family Balistidae) on coral reefs. For example, it is not known how juvenile and adult triggerfish are dispersed among reef zones and habitats in relation to one another, or which habitat types are used. While they are not as diverse or numerous as many other reef fish families, they may be extremely important as consumers in coral reef habitats (McClanahan & Shafir 1990, McClanahan 1994). Balistids characteristically have highly omnivorous diets, with their strong teeth and jaws providing access to a wide range of plant and invertebrate food sources (Reinthal et al. 1984, Kuwamura 1991). However, they may acquire food resources by foraging in specialised habitats, with the family including both planktivorous and benthic feeding species (Randall et al. 1997). Hence, the balistid family may offer insights into the potential effects of ecological versatility on patterns of distribution and abundance.

In this study, we examined the potential relationships between microhabitat specialisation and local distribution and abundance patterns of juveniles and adults of 5 species of triggerfish at Kimbe Bay, Papua New Guinea. These were Balistapus undulatus, Melichthys vidua, Rhinecanthus verrucosus, Sufflamen bursa and S. chrysopterus. We made 2 predictions: (1) that species occupying a wide range of macrohabitats would reach greater local abundance (i.e. across the reef profile) than species restricted to a narrow range of habitats; and (2) distribution among macrohabitats and abundance would be greatest in species that are least specialised in terms of microhabitat and/or specific depth preferences. To evaluate these predictions, we first investigated the distribution and abundance of these species over a broad-scale, from the exposed front reef slope to sheltered back reef slope habitats. We then examined niche breadth in relation to a fine-scale description of depth strata and microhabitats used by individual fish, and determined how this related to broad-scale distribution and abundance.

**MATERIALS AND METHODS**

**Study sites.** This study was conducted in Kimbe Bay, West New Britain Province, Papua New Guinea, during February and March 1999 (5° 25' S, 105° 05' E) (Fig. 1a). Visual censuses of juveniles and adults of the 5 species of triggerfish over the reef profile were carried out on 4 discrete reefs: Garbuna, Madaro, Lady Di and Luba Luba (Fig. 1b). Each reef was divided into
3 broad habitat zones according to its orientation and exposure: front (exposed fore reef slope), middle (reef flat) and back (protected back reef slope). To determine depth distribution patterns at a finer scale and over a greater depth range, 5 additional reefs were sampled (Matane Walindi, Limuka, Crater, Gava Gava and Hanging Gardens) (Fig. 1b).

**Visual censuses of triggerfish.** Visual surveys were used to quantify the abundance of triggerfish at 4 reefs (Fig. 1b). Four replicate belt transects 50 × 4 m at depths of 0 to 1, 2, 6 and 10 m were censused at front and back reef zones for each of the 4 reefs. The middle reef zones were censused using similar methods, but were only sampled at a depth of 0 to 1 m. Tape measures were laid starting from a point parallel to the reef crest. Five metres were left between each transect at each depth. All triggerfish located within 2 m of the centre of the transect were identified to species and their size estimated to the nearest 1 cm standard length (SL). Size estimates were used to assign individuals into juvenile and adult categories. Diet and histological investigation of gonads indicated that the majority of individuals less than 6 cm SL consumed different prey items and were not reproductively mature compared to individuals greater than 6 cm SL (K. Bean unpubl. data). Juveniles of *Melichthys vidua* were, however, classified as individuals less than 12 cm SL as these individuals were still not reproductively mature and were consuming different prey items below this size class (K. Bean unpubl. data).

**Depth estimates.** Mean and variance in depth distributions were assessed in more detail by focal animal sampling over a greater depth range than that used for transects. Sampling involved descending to a depth of 20 m and slowly swimming in a zigzag pattern approximately 50 m wide from 20 m to the surface. These surveys were conducted in front and back reef zones. Each individual triggerfish observed was identified to species and its size estimated to the nearest 1 cm SL. The depth at which each fish was observed was recorded to the nearest metre by the same diver using the same depth gauge for each dive to remove variation associated with different observers and equipment. A total of 8 to 15 dives were conducted to determine the depths of 100 individuals of each species.

**Habitat use and availability.** For juveniles and adults of each of the 5 species of triggerfish, information on microhabitats occupied and availability was collected while carrying out transects and focal animal sampling. Microhabitat use was quantified by recording the substrata over which each individual was first observed during the visual census over the reef profile and while recording the depths of individual fish. In addition, on completion of the visual census of triggerfish on each transect, a second swim back along the transect was used to quantify habitat availability at each depth in each reef zone. This was quantified using 100 random points marked along the length of the transect tape.

Substratum type under each fish and random point was assigned to 1 of 15 substratum categories; sand, rubble, rock, macroalgae, turf algae, coralline algae, massive coral, foliaceous coral, columnar coral, encrusting coral, free living coral, branching coral, soft coral, sponges and solitary organisms (e.g. clams, sea anemones, shells).

**Data analysis.** Distribution and abundance of triggerfishes: The abundance of juvenile and adult *Balistapus undulatus*, *Melichthys vidua* and *Sufflamen bursa* (adults only) were each analysed separately using a 2-way analysis of variance (ANOVA) with reef zone (only the front and back reef zones were compared due to an unbalanced sample design) and depth as fixed factors. *S. bursa* and *S. chrysopterus* juveniles were not analysed in this way due to small sample size (n = 7 and 10, respectively). A 1-way ANOVA was used to compare the abundance of adult *B. undulatus* and adult and juvenile *Rhinecanthus verrucosus* within the 0 m depth categories (between the reef crest, middle and back zones). A 1-way ANOVA was also used to compare the abundances of adult *S. chrysopterus* among depths within the back reef zone due to the restriction of this species to the back reef. Q-Q plots were examined to assess normality and Levene’s test of equality of error variances was used to test for homogeneity of variance (Sokal & Rohlf 1995). Where heterogeneity was detected, data were log_{10} (x + 1) transformed. In all cases transformation stabilised variances. Unplanned multiple comparison tests using Tukey’s HSD were used to test for sources of differences in within-species abundances of adult and juvenile triggerfish. The mean and variance in the depth distributions of each species over a wider depth range and at a finer scale were compared graphically.

**Niche breadth:** Czekanowski’s Index (CI) was used to estimate the degree of similarity between the frequency distribution of habitat type used by individuals of each species and the frequency distribution of habitat available to those individuals. A niche breadth value (CI) for juveniles and adults of each of the 5 species of triggerfish was calculated using the formula:

\[
CI = 1 - 0.5 \sum_i |p_i - q_i|
\]

where \( p_i \) is the proportion of resource items in state \( i \) out of all items used by the population, and \( q_i \) is the proportion of \( i \) states in the resources base available to the population (Feinsinger et al. 1981).

**Microhabitat selection.** Use of resources by all 5 species in relation to availability was explored using
resource selection ratios. Resource selection ratios were chosen as they provide an indication of use in relation to availability and can therefore be used to determine preference for certain habitat types (Manly et al. 1993). Manly et al.’s (1993) Model 1 with Protocol A was used because it requires that animals are not identified individually and the used, unused or available resource units are sampled randomly for the entire study area. A resource selection ratio was calculated separately for juveniles and adults of each species of triggerfish for every substratum category with which they were associated, using the formula:

\[ \hat{w}_i = \frac{o_i}{\pi_i} \]

where \( o_i \) equals the proportion of resources used and \( \pi_i \) equals the proportion of resources available (Manly et al. 1993). Due to multiple comparisons between resources, Bonferroni Z-corrections were used to calculate 95% confidence intervals using the formula:

\[ \hat{w}_i \pm Z_{\alpha/2} \sqrt{\frac{o_i(1-o_i)}{U_i \pi_i^2}} \]

where \( Z_{\alpha/2} \) is the critical value of the standard normal distribution corresponding to an upper tail area of \( \alpha/2 \) and \( U_i \) is the number of used resources in all categories (Manly et al. 1993). The average percent cover of habitat in the reef zones and depths at which individuals were located were compared with the mean number of times individuals were found to associate with each habitat type.

### RESULTS

#### Distribution and abundance of triggerfish

*Balistapus undulatus* was the only species distributed across all habitats and depths. It also reached the greatest densities within any 1 macrohabitat (Fig. 2). There was no significant difference in the abundance of juvenile *B. undulatus* over reef zones or depths (Table 1). There was, however, a significant interaction between reef zone and depth for the abundance of adult *B. undulatus* (Table 1, Fig. 2). The densities of the other 4 species were generally lower and were restricted to a subset of reef zones and depths (Fig. 2).

**Table 1. Two-way ANOVA's assessing abundance patterns of adult and juvenile *Balistapus undulatus*, *Melichthys vidua* and *Sufflamen bursa* (adults only) among the fore and back reef zones and depths. *p < 0.05, **p < 0.01, ***p < 0.001. NA: not applicable**

<table>
<thead>
<tr>
<th>Species</th>
<th>Source of variation</th>
<th>Type III SS</th>
<th>df</th>
<th>MS</th>
<th>F</th>
<th>Tukey’s HSD</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>B. undulatus</em> (adults)</td>
<td>Zone</td>
<td>0.109</td>
<td>1</td>
<td>0.109</td>
<td>3.23</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>Depth</td>
<td>0.09</td>
<td>3</td>
<td>0.03</td>
<td>0.98</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Depth × Zone</td>
<td>0.556</td>
<td>3</td>
<td>0.185</td>
<td>5.48**</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Error</td>
<td>0.811</td>
<td>24</td>
<td>0.034</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>B. undulatus</em> (juveniles)</td>
<td>Zone</td>
<td>0.014</td>
<td>1</td>
<td>0.014</td>
<td>0.87</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Depth</td>
<td>0.553</td>
<td>3</td>
<td>0.184</td>
<td>1.16</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Depth × Zone</td>
<td>0.237</td>
<td>3</td>
<td>0.079</td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Error</td>
<td>3.828</td>
<td>24</td>
<td>0.159</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>M. vidua</em> (adults)</td>
<td>Zone</td>
<td>0.325</td>
<td>1</td>
<td>0.325</td>
<td>4.83*</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Depth</td>
<td>0.323</td>
<td>3</td>
<td>0.108</td>
<td>1.60</td>
<td>Front &gt; Back</td>
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<tr>
<td></td>
<td>Depth × Zone</td>
<td>0.021</td>
<td>3</td>
<td>0.007</td>
<td>0.11</td>
<td>NA</td>
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<td></td>
<td>Error</td>
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<td>24</td>
<td>0.067</td>
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</tr>
<tr>
<td><em>M. vidua</em> (juveniles)</td>
<td>Zone</td>
<td>0.049</td>
<td>1</td>
<td>0.049</td>
<td>1.16</td>
<td></td>
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<tr>
<td></td>
<td>Depth</td>
<td>0.083</td>
<td>3</td>
<td>0.028</td>
<td>0.65</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Depth × Zone</td>
<td>0.096</td>
<td>3</td>
<td>0.032</td>
<td>0.76</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Error</td>
<td>1.017</td>
<td>24</td>
<td>0.043</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>S. bursa</em> (adults)</td>
<td>Zone</td>
<td>1.423</td>
<td>1</td>
<td>1.423</td>
<td>29.6***</td>
<td>Back &gt; Front</td>
</tr>
<tr>
<td></td>
<td>Depth</td>
<td>1.999</td>
<td>3</td>
<td>0.666</td>
<td>13.9***</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Depth × Zone</td>
<td>0.527</td>
<td>3</td>
<td>0.176</td>
<td>3.6*</td>
<td>0 &lt; 2 = 10 &lt; 6 m</td>
</tr>
<tr>
<td></td>
<td>Error</td>
<td>1.153</td>
<td>24</td>
<td>0.048</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Adult and juvenile *Rhinecanthus verrucosus* were restricted to the reef top (Fig. 2). Although adult *R. verrucosus* reached greater abundance at the reef middle versus the reef front, this was not significantly different (Table 2, Fig. 2). Juvenile *R. verrucosus*, however, were more abundant on the back reef compared to the front reef zone (Table 2).

Adult *Sufflamen bursa* occurred on both the front and back reef below 2 m (Fig. 2), yet were significantly more abundant at the back reef zone and at depths of 6 to 10 m (Table 1, Fig. 2). The total number of juvenile *S. bursa* sampled was low, but their distribution closely matched that of adults (Fig. 2). In contrast, adult *S. chrysopterus* occurred in slightly shallower depth strata than *S. bursa* on the back reef (Fig. 2). Adults were significantly more abundant on the back reef at a depth of 2 m (Table 2) and were absent from both the middle and front reef zones (Fig. 2). Juvenile *S. chrysopterus* occurred over all zones but were restricted to a depth of 0 to 2 m (Fig. 2).

**Depth distribution and microhabitat use**

Fine-scale depth estimates for individual fish provided a more direct estimate of the degree to which different species are specialised on different depth strata. A visual comparison of depth distribution patterns among species indicated that *Balistapus undulatus* and *Sufflamen bursa* are generalists with respect to depth, being found at all depths sampled (Fig. 3). All other species were restricted to a narrower depth range (Fig. 3). *Melichthys vidua* occurred over a depth range of 0 to 9 m, occurring at a similar mean depth to

Table 2. One-way ANOVA’s assessing abundance patterns of adult *Balistapus undulatus* and juvenile and adult *Rhinecanthus verrucosus* among 0 m depth categories among reef zones, and adult *Sufflamen chrysopterus* among depths at the back reef zone. *p < 0.05, **p < 0.01. NA: not applicable
that of *S. chrysopterus* (Fig. 3). The most depth-restricted species was *Rhinecanthus verrucosus*, which occurred no deeper than 1 m (Fig. 3).

Niche breadth estimates revealed that the most abundant and broadly distributed species, *Balistapus undulatus*, was also the most generalised species in terms of its microhabitat use (Fig. 4). Those species which were less abundant and which were restricted to particular reef zones and depths had lower niche breadth values indicative of more specialised habitat use within the habitats they occupy (Fig. 4). Juveniles of all species except *Rhinecanthus verrucosus*, were slightly more restricted in their microhabitat use than conspecific adults (Fig. 4).

Use of available habitat indicated that juveniles and adults of the most abundant species, *Balistapus undulatus*, used the greatest number of substratum types (Table 3). Even though *B. undulatus* occupied a wide variety of substratum types, adults used rock, foliaceous coral, and branching coral more frequently than expected on the basis of availability (Table 3). Juvenile *B. undulatus*, however, used turf algae, sponge, sand, macroalgae, encrusting and soft coral more frequently than expected (Table 3, Figs. 5 & 6).

In contrast to *Balistapus undulatus*, juveniles and adults of the other 4 species used a smaller number of resources (Table 3). Adult *Melichthys vidua* were most often associated with massive and branching corals although they were found on other substrata (Table 3). Juveniles of this species were more specialised, using only 5 substratum types and using rock more frequently than expected by chance (Table 3, Fig. 6). Adult *Rhinecanthus verrucosus* were associated with macroalgae, rubble and sand more frequently than expected on the basis of availability (Table 3, Fig. 5). Juvenile *R. verrucosus* used a smaller number of resources (*n* = 5), also using rubble and macroalgae more frequently than expected (Table 3). Adult *Sufflamen bursa* appeared to avoid massive coral, branching coral, coralline algae and turf algae, while using rubble and sand more frequently than expected if non-selective (Table 3). Juveniles were restricted to sand and rubble substrata, using them more frequently than expected on the basis of availability (Table 3). Like adult *S. bursa*, adult *S. chrysopterus* occupied rubble and sand more often than availability would suggest, and used turf algae less frequently than expected (Table 3, Fig. 5). In contrast, juvenile *S. chrysopterus* used 4 substratum categories, using rubble more frequently than expected (Table 3, Fig. 6).
DISCUSSION

The distribution and abundance of *Balistapus undulatus* differs markedly from the other 4 triggerfishes examined in this study. *B. undulatus* exhibited the broadest distribution among reef zones, the greatest local densities and relatively uniform patterns of abundance, suggesting that on a macrohabitat scale *B. undulatus* is a generalist species. *B. undulatus* is also the most versatile species in terms of its microhabitat use among reef zones. It was observed on all substrata that we distinguished in our sampling design, and to a large extent, appeared to use them indiscriminately.

The 4 less abundant species exhibited narrow and distinct patterns of zonation across reef zones, with maximum abundance concentrated in different macrohabitats. *Rhinecanthus verrucosus* is a reef top specialist, *Melichthys vidua* a fore reef slope specialist and the 2 *Sufflamen* species are back reef slope specialists. For the 5 species examined here, the pattern observed was consistent with Brown’s (1984) observation that when closely related individuals of the same guild are compared, those species that have the highest local abundance tend also to have wider distribution patterns.

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Table 3. Outcomes of resource selection ratio calculations for habitat selection within reef zones and depths occupied by juveniles (J) and adults (A) of each of the 5 species of triggerfish. (+: habitat is being used more frequently than available; −: habitat is being used less frequently than available; ns indicates that a habitat is used in accordance with availability; and 0 indicates that a habitat is not used). FL = free living

<table>
<thead>
<tr>
<th></th>
<th><em>Balistapus undulatus</em></th>
<th><em>Melichthys vidua</em></th>
<th><em>Rhinecanthus verrucosus</em></th>
<th><em>Sufflamen bursa</em></th>
<th><em>Sufflamen chrysopterus</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Habitats: J A</td>
<td>132 479</td>
<td>17 131</td>
<td>85 259</td>
<td>15 166</td>
<td>37 92</td>
</tr>
<tr>
<td>Branching</td>
<td>− + ns +</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
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<tr>
<td>Columnar</td>
<td>− − 0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
</tr>
<tr>
<td>Coralline</td>
<td>− ns 0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
</tr>
<tr>
<td>Encrusting</td>
<td>+ − 0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
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<tr>
<td>Folioseous</td>
<td>− + 0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
</tr>
<tr>
<td>Laminar</td>
<td>− ns 0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
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<tr>
<td>Macroalgae</td>
<td>+ − 0 ns</td>
<td>+ +</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
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<tr>
<td>Massive</td>
<td>ns ns − +</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
<td>0 −</td>
</tr>
<tr>
<td>Rock</td>
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<td>ns ns − ns</td>
<td>+ +</td>
<td>+ +</td>
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</tr>
<tr>
<td>Sand</td>
<td>+ ns 0 −</td>
<td>− −</td>
<td>+ +</td>
<td>+ +</td>
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<td>FL coral</td>
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<td>0 0</td>
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<tr>
<td>Soft coral</td>
<td>+ ns 0 −</td>
<td>0 0</td>
<td>0 0</td>
<td>0 0</td>
<td>0 0</td>
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<tr>
<td>Turf algae</td>
<td>+ ns 0 − ns ns</td>
<td>0 ns</td>
<td>ns ns</td>
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</tr>
<tr>
<td>Dead coral</td>
<td>− − ns 0 0</td>
<td>0 0</td>
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</tr>
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</table>

Fig. 5. Proportion of habitat used by adult *Balistapus undulatus*, *Melichthys vidua*, *Rhinecanthus verrucosus*, *Sufflamen bursa* and *S. chrysopterus* compared with the availability of those substrata within the reef zones and depths each species occurred.
Depth distribution patterns also suggest that *Balistapus undulatus* is one of the most versatile species in this guild. While *B. undulatus* appears to prefer depths of 2 to 8 m, it was regularly found to at least 20 m with *Sufflamen bursa* also found over this depth range. Comparing all species, there appears to be a pattern of increasing variance in the depth range with increasing average depth. That is, there were no examples of a deep species with a narrow depth range. It is possible that distributions among macrohabitats are explained by differences among species in their ability to use deeper water, regardless of habitat structure. However, different abilities between species to use deeper water would not explain the differences in the distribution of some species between front and back reef habitats. Clearly, depth and habitat structure are closely linked and further work is required to isolate their effects over the whole reef profile.

The preference of adult *Melichthys vidua* for massive coral, and in particular front reef locations, may be related to *M. vidua* being a planktivore, compared to the other species which are omnivores (K. Bean unpubl. data). By occupying a shallow position on reef fronts, exposure to currents carrying plankton would likely be increased (Hobson 1991). Preference for massive coral may therefore reflect high abundance in this area and its subsequent use by *M. vidua* as a shelter site, rather than for food acquisition. Juvenile *M. vidua* displayed a similar distribution pattern to adults although they were more often associated with branching coral. Again, preference for substrata of higher topographic complexity may be greater at the juvenile stage when they may be more susceptible to predators (Hixon & Beets 1993, Caley & St. John 1996).

In contrast, *Rhinecanthus verrucosus* was restricted to reef tops and therefore had the narrowest depth distribution. Adult and juvenile *R. verrucosus* were associated with macroalgae and rubble more frequently than expected. The peak in abundance of adult and juvenile *R. verrucosus*, however, occurred at different locations on the reef top. Other factors such as competitive interactions or differential habitat requirements between juveniles and adults may influence the observed pattern, resulting in disjunct distributions. Competitive interactions between adults and juveniles where resource requirements overlap has been found in previous studies to result in juveniles being excluded from adult habitats (Jones 1987, Levin 1993). Hence, an understanding of behavioural interactions between adults and juveniles may be required to understand distribution patterns of *R. verrucosus* within the reef top zone. Alternatively, differences in diet and the distribution of food items may influence differences in the distribution of adult and juvenile *R. verrucosus*. Prey availability has previously been shown to be important in determining the distribution of the grey triggerfish *Balistes capriscus*, for which there was a positive relationship between sand dollar abundance and the abundance of adult triggerfish (Kurz 1995). Diet may therefore influence patterns of distribution due to differences in the distribution of preferred prey items.

*Sufflamen bursa* and *S. chrysopterus* exhibited disjunct distributions on the back reef slope, with very little overlap in their depth distributions. Both species,
however, displayed high overlap in habitat use, preferring sand and rubble as both juveniles and adults, and occurring primarily in the back reef zone. High overlap in habitat use suggests that depth distribution patterns on the back reef zone may be affected by competitive interactions between the 2 Sufflamen species (Ebeling & Laur 1986). Behavioural interactions may therefore be an important mechanism shaping the distribution patterns of S. bursa and S. chrysopterus across the reef profile, where the competitively dominant species may occupy the preferred depth, pushing the other species into deeper or shallower water. In this case, the more specialised of the 2 species would be expected to out-compete the more generalised species (Futuyma & Moreno 1988).

The guild of balistids studied here provides some insight into how distribution and abundance patterns may be influenced by microhabitat specialisation and the availability of resources. Although this study provides one of the first tests of ecological versatility for coral reef fish, other factors may be influencing the observed degree of specialisation, including interactions among and within fish species. A greater range of species from this and other taxa, however, must be examined to confirm this pattern. Therefore, experimental studies are needed to test whether such behavioural interactions influence microhabitat use. For example, can the 4 specialist species out-compete Balistapus undulatus within their preferred habitat zones?

Clearly, local patterns of distribution and abundance are likely to result from complex interactions among species, competitors and resources. Disentangling these potential causes of observed distributions is worthy of greater attention in the future.

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