MEALS FOR TWO: FORAGING ACTIVITY OF THE BUTTERFLYFISH *Chaetodon striatus* (PERCIFORMES) IN SOUTHEAST BRAZIL

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(With 2 figures)

**ABSTRACT**

The banded butterflyfish (*Chaetodon striatus*) from the tropical and subtropical western Atlantic is a territorial, diurnal forager on benthic invertebrates. It is usually seen moving singly or in pairs, a few meters above the sea floor. We studied the foraging activity of *C. striatus* on rocky reefs in southeastern Brazil. This fish spent about 11 h and 30 min per day on feeding activities, and preferred colonies of non-scleratinian anthozoans over sandy and rocky substrata while foraging. The lowest feeding rates were recorded in the early morning and late afternoon, but we found no further differences between feeding rates throughout the day. We also found no differences between the feeding rates of paired and single individuals.

*Key words*: Chaetodontidae, foraging, substrate preferences, southwestern Atlantic.

**INTRODUCTION**

The butterflyfishes (Chaetodontidae) are conspicuous components of the reef community on tropical and subtropical coral reefs (Pitts, 1991). These fishes swim by daylight, often in pairs, and reside for up to five years in the same reef area and with the same partner (Reese, 1973; Fricke, 1986; Driscoll & Driscoll, 1988). Consequently, mates, food, and shelter are potentially defensible (Driscoll & Driscoll, 1988; Neudecker & Lobel, 1982). Swimming in pairs and mutual partner guarding are regarded as a joint territorial advertisement that minimizes agonistic interactions between territory holders (Fricke, 1986).
Chaetodontidae are mostly bottom-foragers, although some species forage for plankton in the water column (Hobson, 1974, 1991; Motta, 1988). The diet of the bottom-foraging species is composed mostly by anthozoans, polychaetes, small crustaceans, and mollusk eggs (Hiatt & Strasburg, 1960; Randall, 1967; Sano, 1989). Most of the knowledge about feeding activity of butterflyfishes originates from studies in the Caribbean (e.g., Birkeland & Neudecker, 1981; Pitts, 1991) and the Pacific (e.g., Hobson, 1974; Irons, 1989; Tricas, 1989).

The banded butterflyfish, Chaetodon striatus, is found throughout the Western Atlantic, from New Jersey in the U.S.A. to Santa Catarina in Brazil (Carvalho-Filho, 1999). It is often found in pairs on shallow coral and rocky reefs (Menezes & Figueiredo, 1985). This fish is mostly a bottom-forager (Pitts, 1991), although it may forage for plankton in the water column (Sazima & Sazima, 2001). Its diet is composed primarily by non-scleratinian anthozoans and polychaetes (Randall, 1967; Motta, 1989; Pitts, 1991).

We studied the foraging of C. striatus in the SW Atlantic, a region where information about the ecology of this butterflyfish is scarce (Menezes & Figueiredo, 1985; Carvalho-Filho, 1999), especially that related to feeding (Sazima & Sazima, 2001). Our study addressed the three following questions about the foraging activity of C. striatus: is there any substrate selection for foraging? Do the feeding rates differ throughout the day? Do the feeding rates of paired and single individuals differ?

**MATERIAL AND METHODS**

**Study site**

The field study was conducted on the Ilha de São Sebastião (23°46'S, 45°21'W), off the coast of São Paulo in southeastern Brazil, from February to June 2002. The study site was located on rocky reefs covered mostly by algae; by zoanthid colonies, mostly *Palythoa caribaeorum*, and *P. variabilis*; and by the hard coral *Mussismilia hispida*. It also covered adjacent sandy areas. During our study, horizontal visibility was 2-7 m, and water temperature was 26-27°C. Depth at the study site ranged from 1 to 3 m.

**Procedure**

The foraging activity of C. striatus was observed in eight nonconsecutive days, while snorkelling. During observational sessions of 60-150 min, “focal animal” and “all occurrences” samplings (Lehner, 1979) were used in 371 min of direct observation. The foraging activity of C. striatus was assessed by following individuals for 1-5 min and counting the number of bites on the three available substrate types (anthozoan colonies, and rocky as well as sandy bottoms). To avoid the risk of biased samples, individuals were not followed over successive periods (Birkeland & Neudecker, 1981).

The observations were conducted at daytime, from sunrise (5:30 h) to sunset (18:45 h). No nocturnal observations were conducted, as C. striatus is inactive at night (Starck & Davis, 1966, pers. obs.). The distance between the pair members changed during their foraging, making impracticable the observation of both individuals at the same time. Thus, only one of the partners was followed during the observational sessions. We considered as single an individual whose partner was not visible throughout a 1-5 min session. In a total of 100 foraging bouts, 58 occurred in the morning (5:00 h-13:00 h) and 42 in the afternoon (13:01 h-19:00 h). Additionally, of these 100 bouts, 62 focused on paired individuals and 38, on single ones.

We assessed the availability of the three foraging substrate types (anthozoan colonies, and rocky and sandy bottoms) with six 10 x 2 m transects (modified from Birkeland & Neudecker, 1981). The transects started on the rocky shore and ended in the sandy area where C. striatus was still observed feeding.

Kruskal-Wallis one-way analysis of variance was used to compare the number of bites per min for successive periods of 120 min of foraging time throughout the day. As the Kruskal-Wallis was significant, a posteriori tests were made to compare pairs of 120 min periods, for identifying which period(s) caused the differences. The goodness-of-fit test was used to determine the substrate selection by C. striatus. Two-way ANOVA was used to test for differences in feeding rates between paired and single individuals, and between morning and afternoon. The time spent during feeding bouts was
compared between paired and single individuals by means of the Student’s t test (Zar, 1999).

RESULTS

The feeding activity of *C. striatus* started about 30 min after sunrise (5:45 h) and ended shortly before nightfall (18:26 h), totaling about 11 h and 30 min of feeding activity per day. The feeding rate of *C. striatus* was $1.60 + 1.76$ bites min$^{-1}$ (average + standard error, $n = 100$). The fish foraged predominantly on anthozoans over sandy and rocky bottoms ($X^2 = 26.44, p < 0.001$; Fig. 1), and selected only the non-scleratinians (zoanthids) *Palythoa variabilis* and *P. caribaeorum*.

Feeding frequency varied throughout the day ($H_6 = 19.92, p = 0.003$; Fig. 2). The lowest feeding rates were recorded in the early morning and late afternoon (in some observational sessions *C. striatus* did not forage at all). We found no significant differences between the morning and afternoon feeding rates nor between paired and single individuals ($F_1 = 0.24, p = 0.63$ (Table 1). On the other hand, the foraging bouts of paired individuals ($x = 247 + SD = 72 s$) were longer than those of the single ones ($x = 195 s + SD = 92 s$) ($t_{96} = 3.14, p < 0.005$).

DISCUSSION

We found that the time *C. striatus* spends foraging is high when compared to the values obtained for *C. trifascialis* from the Johnston Atoll (16°45'N, 169°31'W) in the Pacific, the only other study in which data were collected from sunrise to sunset (Irons, 1989). For *C. trifascialis*, a total of 10 h per day were spent feeding (Irons, 1989), a value slightly below that obtained in our study (11 h and 30 min). The slightly higher foraging activity here recorded for *C. striatus* may be related to the higher latitude and greater number of daylight hours (our study took place in austral summer). Since *C. trifascialis* and *C. striatus* are diurnally active (Starck & Davis, 1996; Irons, 1989), it was to be expected that *C. striatus* in São Sebastião island would forage for a larger period than did *C. trifascialis* in Hawaii.

Our estimates of the feeding rates of *C. striatus* are low when compared to the values found for *C. capistratus* (5.88 + 1.96 bites min$^{-1}$) in the Caribbean (Birkeland & Neudecker, 1981) and *C. trifascialis* (8.45 + 0.72 bites min$^{-1}$) in the Pacific, both of which feed almost exclusively on hard corals (Randall, 1967; Birkeland & Neudecker, 1981; Motta, 1989). However, the feeding rates of *C. striatus* are similar to those recorded for the butterflyfish *Prognathodes aculeatus* in the Caribbean (2.44 + 0.92 bites min$^{-1}$), which feeds on invertebrates such as polychaetes and crustaceans (as *C. striatus* also does, see Randall, 1967; Birkeland & Neudecker, 1981; Motta, 1989). Butterflyfish species that feed on hard coral exhibit significantly more bites than species feeding on polychaetes and crustaceans (Birkeland & Neudecker, 1981). Thus, the differences recorded for the feeding rates of *C. capistratus* and *C. trifascialis* versus *P. aculeatus* and *C. striatus* are due to diet, as invertebrates have a higher caloric value than do hard corals (Birkeland & Neudecker, 1981; Motta, 1989).

In our study *C. striatus* selected non-scleratinian anthozoans, whose colonies shelter polychaetes and crustaceans (Gleibs et al., 1995). Therefore, it is possible, although yet to be verified, that *C. striatus* from southeast Brazil also includes these associated invertebrates in its diet, as already has been recorded for the Caribbean (Randall, 1967; Pitts, 1991).

**TABLE 1**

Two-way ANOVA for the number of bites per min recorded for *Chaetodon striatus* in two daytime periods: morning (5:00 h-13:00 h) and afternoon (13:01 h-19:00 h), and two social categories (paired and single individuals).

<table>
<thead>
<tr>
<th>Sources</th>
<th>df</th>
<th>Mean square</th>
<th>F</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Periods (morning/afternoon)</td>
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<td>1.768</td>
<td>0.564</td>
<td>0.45</td>
</tr>
<tr>
<td>Categories (paired/single)</td>
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<td>0.751</td>
<td>0.240</td>
<td>0.63</td>
</tr>
<tr>
<td>Interaction</td>
<td>1</td>
<td>3.743</td>
<td>1.194</td>
<td>0.277</td>
</tr>
</tbody>
</table>
Fig. 1 — Number of expected and observed bites by *Chaetodon striatus* on three feeding substrate types in São Sebastião island, São Paulo, southeast Brazil.

The feeding frequency variation throughout the day here recorded for *C. striatus* is mostly due to the beginning and the end of its feeding activity, at which times it feeds little or not at all. Also, the nonsignificant differences between morning and afternoon feeding rates are similar to what was recorded in the Caribbean for *P. aculeatus* (Birkeland & Neudecker, 1981), whose diet is similar to that of *C. striatus* (Randall, 1967). On the other hand, *C. trifascialis*, which feeds on almost exclusively on *Acropora* hard corals that have maximal lipid production during this period (Irons, 1989), does so at a significantly higher rate in the early afternoon (Irons, 1989). Furthermore, we found no significant differences between the feeding rates of paired and single individuals of *C. striatus*, which contrasts with the findings of Fricke (1986) for *Chaetodon chrysurus* in the Red Sea.

Butterflyfishes defend territories in pairs and move with mutual partner guarding, a behavior regarded as a joint territorial advertisement (Fricke, 1986; Roberts & Ormond, 1992). The removal of one member in a *C. chrysurus* pair resulted in territories being reduced to one quarter of the same area held when defended by pairs (Fricke, 1986). In addition, the distance covered per time unit by the remaining fish greatly increased, primarily because they now had to defend the area against an increased number of intruders. Thus, we probably found no differences between the feeding rates of paired and single individuals because in our study the partners stayed away from each other for short

Fig. 2 — Comparison limits of bites by *C. striatus* per min. on three substrate types during the day. Bars showing the same pattern are not significantly different.
periods only, which were presumably insufficient to cause an increase of intruders within the butterflyfish territories. We also surmise that the removal of one of the *C. striatus* partners will result in reduction of the remaining partner’s feeding rate, in a way similar to that recorded for *C. chrysurus* (Fricke, 1986).

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**REFERENCES**


