Changes in foraging as a response to predation risk in two gobiid fish species, *Pomatoschistus minutus* and *Gobius niger*

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ABSTRACT: The influence of predation risk and hunger level on the foraging rate and swimming activity in the sand goby *Pomatoschistus minutus* and the black goby *Gobius niger* was tested in aquarium experiments. The fish were given a choice of 2 habitats, viz. vegetation without food, and open sediment with natural prey density. Both species had a lower food consumption in the presence of predators than without predators, but starved fish had a higher consumption rate than fed fish both in the presence and in the absence of predators. Starved fish thus took higher risks of being eaten than did fed fish. *P. minutus* decreased their swimming activity in the presence of predators, which may have caused the lower food intake. However, *G. niger* did not change their activity with predator presence. Habitat choice must also be of importance for the foraging rate in gobies. *G. niger* were mostly found in the vegetation, which is their natural habitat. *P. minutus* to a greater extent inhabit open areas in the field and therefore might gain more from decreasing their activity than seeking cover at high predation risk.

INTRODUCTION

Many studies have shown that the risk of being eaten by a predator may influence the foraging of an animal (Milinski & Heller 1978, Krebs 1980, Sih 1980, Dill 1983, Lima et al. 1985, Metcalfe et al. 1987, Magnhagen 1988). There are often trade-offs to be made between different behaviours (Sih 1980, Dill & Fraser 1984, Lima et al. 1985, Milinski 1986), and it is important to examine the interactions between different demands such as the need for predator avoidance, feeding and mating, in order to understand the foraging patterns of an animal (Sih 1987). One goal for a forager should be to maximize its probability of survival (Mangel & Clark 1986, McNamara & Houston 1986), and different behaviours could be profitable in different situations depending for example on predation risk and hunger level. There are several ways to avoid predators. To decrease the risk of detection an animal can either use refuges (Mittelbach 1981, Cerri & Fraser 1983, Werner et al. 1983, Power 1984, 1987, Schlosser 1987) or reduce its activity (Dill 1983, Dill & Fraser 1984, Godin 1986, Sih 1986, Giles 1987, Prejs 1987). Both these strategies, however, can be costly in terms of lost opportunities to feed. A hungry animal may have to take higher risk of being eaten, in order to get the food it requires, than a less hungry one (Mangel & Clark 1986, McNamara & Houston 1986). According to the foraging models developed by Mangel & Clark (1986) and McNamara & Houston (1986) the 'metabolic state' of an animal (e.g. energy level or gut contents) is the factor that decides the choice of actions made by the animal, since the state determines the probability of survival given a certain behaviour.

The sand goby *Pomatoschistus minutus* (Pallas) and the black goby *Gobius niger* L. (Gobiidae) are marine benthic fish species occurring in great numbers in shallow soft-bottom areas along the Swedish coast. *G. niger* has a strong preference for vegetated habitats, while *P. minutus* is found both in vegetation and in open areas (Magnhagen & Wiederholm 1982, Wiederholm 1987). Both species feed on invertebrate prey, mostly crustaceans. Pihl (1982) reported that cod *Gadus morhua* L. preys upon gobies, especially in the autumn, and other potential predators in these areas include sculpins *Myxocephalus scorpius* (L.) and eels *Anguilla anguilla* (L.).

In this study I examine the foraging behaviour in these 2 gobiid fishes, more specifically the effect of predation risk and hunger level on foraging.
experiments were performed to look at food intake, habitat choice and activity, in the presence and absence of a predator. I wanted to test whether the foraging rates were affected by predation risk, and if so, what mechanisms could explain the changes. Furthermore, would hunger level affect the response to predators as predicted by Mangel & Clark (1986) and McNamara & Houston (1986), i.e. would a starved fish take higher risks to be eaten in order to get food than a fish with a higher energy reserve?

METHODS

The study was carried out in July and August 1987 at Klubbans Biological Station, Fiskebackskil, on the west coast of Sweden (58°15'N, 11°28'E). Pomatoschistus minutus were caught with a hand-trawl at depths ≤ 1 m in a shallow bay in the vicinity of the laboratory, and Gobius niger were caught in an eelgrass (Zostera marina L.) meadow at depths of 1 to 2 m using a beam trawl. The fish were taken to the laboratory and kept in storage tanks. They were either starved for 1 wk or fed twice a day with commercial salmon pellets, mussels and live invertebrates, in order to obtain fish with 2 hunger levels.

Feeding experiment. To study food intake and habitat choice in the presence and absence of predators, 4 fish tanks offering the choice between 2 habitats, one with food and the other with protection from predators, were used. The volume of the fish tanks was ca 1800 l, with a bottom area of 2.3 m². The bottom substrate in half of each tank consisted of sediment taken from the same area as Pomatoschistus minutus, scooped up in such a way as to minimize disturbance of the infauna (mostly Corophium volutator (Pallas) and Nereis diversicolor O. F. Müller) and to provide natural prey density. In the other half the bottom was covered with sand from the beach which contained no prey organisms. On the sand were placed plastic aquatic plants, imitating eelgrass, to provide cover. The water was continuously renewed, with temperatures around 15 °C. Twelve gobies were put in each tank at 23:00 h. Observations on habitat choice started at 08:00 h the following morning. Once every hour, the number of fish seen in the 2 habitats was registered. At 19:00 h the fish were removed, anaesthetized in MS222, and frozen. Their stomachs were later analyzed, number of prey counted and their stomach contents dried for 48 h at 60 °C and weighed.

In 2 of the tanks the gobies used were starved for 1 wk before the experiment started, while in the other 2 they had previously been fed twice a day. In 2 of the tanks there was 1 cod (body length ca 20 cm) to provide predation risk (the cod actually ate a few gobies during each run). The experimental setup thus created 4 different treatments: starved fish with and without predator, and fed fish with and without predator. The experiments were repeated 5 times each for Pomatoschistus minutus and Gobius niger. Mean body length was 55.1 ± 9.4 mm (N=182) for P. minutus and 56.8 ± 10.5 mm (N=200) for G. niger.

Activity experiment. The swimming activities of Pomatoschistus minutus and Gobius niger, in the presence and absence of predators, was measured using photo-cells registering number of passages across an infra-red beam. The aquaria used were octogonal (diam. = 60 cm) with double walls, 10 cm apart, creating 2 separate sections, an outer and an inner (Fig. 1).

![Fig. 1. Design of activity aquarium, shown from above. A: predator chamber; B: goby chamber; - - - - location of infrared beams; - - - - masonite screens](image)
'predation risk' situation and the control. In one of the 'predation risk' aquaria, the gobies had been starved for ca 1 wk, while in the other fish that previously had been continuously fed were used. In the control aquarium the fish were previously fed. The experiments were repeated 5 times with Pomatoschistus minutus and 6 times with Gobius niger. Each individual fish was used only once in the experiments.

RESULTS

Feeding experiments

The observations on habitat choice were pooled for each treatment within each replicate. The proportions of fish found in the open habitat were arcsin $\sqrt{x}$ transformed and the influence of predator presence, hunger level and species was tested in a 3-way ANOVA. Predator presence and species were found to have a significant influence on habitat choice ($F_{1,24} = 23.5, p < 0.001; F_{1,24} = 45.6, p < 0.001$, respectively). In the presence of predators more fish were found in the open habitat than in the absence of predators (Fig. 2). More Pomatoschistus minutus than Gobius niger were found in the open. Usually only a low proportion of the fish were in the open habitat; only P. minutus in the presence of predators occupied the open area to more than 50%.

Food intake was calculated as the mean dry weight of the stomach contents from the fish in each replicate treatment. A 3-way ANOVA on the log$(x+1)$ transformed data tested the influence of predator presence, hunger level and species. All 3 factors had a significant influence on the food intake ($F_{1,31} = 9.0, p < 0.01$ (predator presence); $F_{1,31} = 12.6, p < 0.01$ (hunger level); $F_{1,31} = 6.7, p < 0.05$ (species). Starved fish had a higher consumption than fed fish, and Pomatoschistus minutus ate more than Gobius niger (Fig. 3). Fish at the same hunger level had a higher food intake in the absence of predators.

![Fig. 2](image)

Fig. 2. (a) Pomatoschistus minutus and (b) Gobius niger. Proportion of fish found in the open habitat in the foraging rate experiments. Data points show the means ($\pm$1SD) from 4 experiments for both fish species.

![Fig. 3](image)

Fig. 3. (a) Pomatoschistus minutus and (b) Gobius niger. Mean dry weight (mg) $\pm$ SE of stomach contents from the fish in the foraging rate experiments.

Activity experiments

The swimming activity is expressed as the number of times per hour the fish passed either of the 2 photocells in each aquarium (Fig. 4). The means for the light periods and the dark periods were used when testing the effect of predator presence, hunger and light on the activity (3-way ANOVA, Table 1).

In the Pomatoschistus minutus experiments a higher activity was found during the first 24 h in the aquarium without predator compared to the others (Table 1; Fig. 4). On the second day, when the predators were removed, however, the fish in the 'predation risk' aquaria increased their activity and there were no differences between the 2 aquaria during Day 2 (Fig. 4). During the dark period the activity was lower during both Day 1 and Day 2 (Table 1).

In the Gobius niger experiments light influenced the activity only during Day 1 (Table 1). This was mainly due to the high activity during the first hours the fish spent in the aquarium (when disregarding the data from the first 4 h, there was no significant difference between day and night). During Day 2 the starved fish had a significantly lower activity than the fed fish. There were no differences in activity aquaria with or without predator.
Fig. 4. (a) Pomatoschistus minutus and (b) Gobius niger. Activity measured as number of passages per hour by 3 fish through an infrared beam. Data points show the means from 5 (P. minutus) or 6 (G. niger) replicate experiments. (○-○) Starved fish with predator present; (●-●) fed fish with predator present; (⋯⋯) fed fish without predator; bars: dark periods; arrows: removal of predators.

Table 1. Pomatoschistus minutus and Gobius niger. Three-way ANOVA on the mean activity per hour measured as number of passages by 3 fish through an infra-red beam. P: predator presence; H: hunger level; L: light condition. On Day 2 the predators were removed, and P is then comparing the aquaria that previously had a predator present with the control aquarium.

<table>
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**DISCUSSION**

Both Pomatoschistus minutus and Gobius niger changed their foraging as a response to predation risk. Furthermore, the response differed depending on hunger level. However, the higher food intake in the absence of predators was not due to increased utilization of the habitat with the higher prey density as would be expected. In the presence of a predator both species, but especially P. minutus, were found more commonly in the open area than otherwise, even if most of G. niger still remained in the vegetation. This can depend on the location of the predator. The cod spent ca 70% of their time in the vegetation, and thus the choice for the gobies was between seeking cover or maximizing their distance from the predator. For P. minutus, which is not as strongly associated with vegetation as G. niger, the structure of the sediment may be more important than vegetation. At least to the human eye, P. minutus were actually more cryptic on the open sediment, which was more heterogeneous, with holes from burrowing animals etc., than on the smooth sand in the other area. Furthermore, P. minutus has the ability to burrow which also suggests that type of substrate may be more important than presence of cover.

The lower average food intake in Gobius niger compared to Pomatoschistus minutus was probably due to the fact that the former occupied the vegetation to a higher extent. Thus, in G. niger which usually lives in vegetation, cover seems more important than type of substrate, and the differences between the 2 species...
found in the experiments probably reflect the differences in their natural habitats. Obviously, in my experiments the division of the tanks into 2 habitats was of minor importance for the results, but the difference between treatments in food intake depended instead on differences in foraging activity.

An animal is more visible when moving than when stationary, hence decreasing activity would lower the risk of being detected by a visually searching predator as well as decreasing prey encounter rate (Sih 1987). Banded killifish Fundulus diaphanus (LeSueur) reduced the rate of feeding attempts in the presence of a predator (Godin 1986). Prejs (1987) found a decrease in feeding activity in small tropical stream fishes in the presence of gape-limited predators compared with predator-free areas. Similarly, small crayfish Orconectes propinquus (Girard) reduced their activity and chose substrates offering protection as a response to predation risk (Stein & Magnusson 1976). It is likely that a lower swimming activity in the presence of predators, as shown in the activity experiments for Pomatoschistus minutus, will lead to lower food consumption. This should also apply to Gobius niger since they too consumed less food in the presence of a predator. However, in this species there was no decrease in activity in the experiments. Earlier studies on activity in G. niger showed that their activity was lower in the presence of cover (Hesthagen 1976). In my experiments no vegetation or any other kind of cover were provided, and therefore the activity might have been higher than it would otherwise have been, and maybe higher than in the foraging rate experiments, where vegetation was present.

Starved fish had a higher food intake than fed ones, both in the presence and in the absence of predators. Obviously, they were taking higher risks in order to get food than did the fed fish. This probably reflects a trade-off between energy requirements and the risk of being eaten by a predator. Situations like this are likely to arise in habitats and at seasons with low food availability. Hungry animals have previously been found to be less responsive to predators, e.g. fish (Milinski & Heller 1978, Dill & Fraser 1984, Maghnagen 1986), insects (Bailey 1986, Wellborn & Robinson 1987), and birds (Krebs 1980). The tendency to take higher risks at higher hunger levels is a behaviour that is predicted by the foraging models developed by Mangels & Clark (1986) and McNamara & Houston (1986). The animal is expected to always choose the behaviour that maximizes the probability of survival. By looking at the 'metabolic state' of the animal it is possible to predict what risks it should take, its habitat choice, activity level, etc. The changes in foraging behaviour made by Pomatoschistus minutus and Gobius niger as a response to predation risk and hunger level thus agree with predictions made by Mangel & Clark (1986) and McNamara & Houston (1986). However, choice of activity made by an animal may also be influenced by demands other than energy intake and predator avoidance, such as reproduction and territorial defence.

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