

Complexity affects habitat preference and predation mortality in postlarval *Penaeus plebejus*: implications for stock enhancement

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ABSTRACT: Global attempts to offset declines in fishery populations through stock enhancement have had varied levels of success due to the absence of preliminary studies to determine which habitats best support release species and the mechanisms controlling their distribution. Habitat preference was examined as a possible mechanism driving distribution of postlarval *Penaeus plebejus*, a current candidate prawn for stock enhancement in Australia. Occupancy of complex (artificial macrophyte) and simple (bare sand and mud) habitats by postlarvae was compared in the presence and absence of a choice between the habitats. Predation mortality was also compared amongst these habitats. *P. plebejus* settled into the different habitats randomly during the night, but actively selected macrophyte over the simple habitats during the day. Mortality caused by the predatory fishes *Centropogon australis* and *Acanthopagrus australis* was higher in simple habitats than in complex habitats, but was similar across habitats when large penaeid prawns, *Metapenaeus macleayi* (which are tactile rather than visual feeders), were used as predators. Postlarvae may select macrophyte habitats during the day to lower predation risk, but because nighttime foraging efficiency is reduced in their predators, which are primarily visual hunters, this may preclude the need of postlarvae to obtain shelter in macrophyte habitats at night. Predation mortality of stocked *P. plebejus* may be minimized by releasing postlarvae directly into macrophyte habitats. Studies such as these must precede all stock enhancement attempts because they identify optimal release strategies and allow ecological and financial costs of enhancement to be weighed against projected benefits, and thereby assess the practicality of enhancement as a management option.

KEY WORDS: Stock enhancement · Habitat · Predation · Macrophytes · Fisheries management · Prawns · Penaeid · Eastern Australia

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INTRODUCTION

Global declines in fishery stocks due to overexploitation and habitat fragmentation (Pikitch et al. 2004, Anderson et al. 2008) have regenerated interest in stock

enhancement as a means of replenishing depleted populations through the release of hatchery reared individuals into the wild (Rothlisberg et al. 1999). However, this management approach is rarely successful and is often characterized by low survival rates

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amongst released individuals (Blankenship & Leber 1995, Grimes 1998). Historically, the failure of enhancement events has been caused by impetuous hatchery production and indiscriminate releases that have hampered development of a scientific basis for how best to optimize stock enhancement or whether this approach should even be used (Leber 2002). Previous assessment of the ecological factors that affect the survival and growth of stocked species can dramatically reduce post-release mortality and can help evaluate the practicality of enhancing a species within a given system (Crowl et al. 1992, Eades & Steinkoenig 1995, Einum & Fleming 2001). One such ecological factor is the capacity of different habitats to support a stocked species (Rothlisberg 1998), which is partly determined by the species' habitat preference and habitat related predation mortality. Our study evaluates these determinants of carrying capacity for the eastern king prawn *Penaeus plebejus* Hess, 1865, which has been recently identified as a candidate for stock enhancement in New South Wales, Australia.

Penaeus plebejus is one of the most economically important penaeid prawns found in Australia, and has an annual commercial production valued at ~AUS \$40.5 million (Courtney 2002) and a significant recreational fishery (Ives & Scandol 2007). High exploitation rates, habitat loss and recruitment variability have led to significant declines in the abundance of many penaeid species (Rothlisberg et al. 1999) and trial stock enhancement has been assessed in different regions to address these factors with limited success (Su & Liao 1999, Loneragan et al. 2007). Understanding which types of habitats best support these species and the mechanisms controlling their habitat use is critical to improving the success of future enhancement efforts for penaeids.

Penaeid prawns show a high degree of association with distinct habitats (e.g. Dall et al. 1990a). For some species, structurally complex habitats such as macrophyte beds generally support more individuals (Heck & Thoman 1984, Hill & Wassenberg 1993), especially during their juvenile stages (Haywood et al. 1995). *Penaeus plebejus* juveniles are usually more abundant in seagrass beds than adjacent bare habitats at small spatial scales (Young 1978), and their densities are often positively correlated with seagrass cover (Skilleter et al. 2005). Similar relationships in other penaeids have been attributed to greater protection from predators, increased feeding resources or a combination of these factors (Bell & Westoby 1986, Liu & Loneragan 1997). Despite abundant data supporting these theories in other species (e.g. Loneragan et al. 1997, Primavera 1997, Macia 2004), it is still unclear whether the immediate cause of higher densities of *P. plebejus* within macrophyte beds is simply a result of

lower post-settlement mortality (via predation or starvation) within this habitat or an innate preference for macrophyte beds during settlement. If *P. plebejus* postlarvae show a preference for structured habitats, then behavioral mechanisms could be influencing their distribution before post-settlement mortality through predation or starvation. Although other factors that are distinct from behavioral or interactive processes between species may also influence the distribution of *P. plebejus* amongst habitats (e.g. small scale habitat-related variation in hydrodynamics may allow more postlarvae to settle more frequently in particular habitats), the research presented here focuses on the influence of preference and predation mortality.

We investigate whether the distribution of *Penaeus plebejus* amongst habitats is a result of active selection during its postlarval phase, and whether selection differs between day and night by comparing selection of macrophyte, bare sand and bare mud habitats during these diel periods. The habitats examined represent the most prevalent habitat types in local estuaries where enhancement of *P. plebejus* is being considered. They also differ in characteristics (other than complexity) that may influence a postlarva's ability to use them. For example, the anoxic nature of mud and its smaller pore spaces relative to sand may limit respiration in postlarvae (Williams 1958, Mnaya et al. 2006). In contrast, the high organic content of mud compared with sand may increase food supply within muddy habitats (Williams 1958, Van Luijn et al. 1999, Mnaya et al. 2006). Accordingly, some penaeid species occur more frequently on sandy substratum than muddy substratum (e.g. *P. duorarum*), whilst other penaeids exhibit the opposite pattern (e.g. *P. aztecus* and *P. setiferus*) (Williams 1958). This research also tests whether postlarval predation mortality differs between macrophyte, sand and mud habitats. We conclude by considering the implications of our results in relation to stock enhancement, providing a scientific basis for releasing *P. plebejus* into estuarine systems.

MATERIALS AND METHODS

Rearing and transport of *Penaeus plebejus*. Hatchery-reared *Penaeus plebejus* postlarvae were produced by a commercial supplier, using wild brood stock collected off the central east coast of Australia (between 30° 16' 49" S, 153° 12' 6" E and 24° 45' 6" S, 153° 1' 22" E). At 17 d old (carapace length [CL] ~ 4 mm), the postlarvae were transferred to a flow-through, coarse-filtered aquarium facility in Cronulla, New South Wales (34° 4' 21" S, 151° 08' 56" E) by air and road in foam boxes containing 10 l of water within sealed plastic bags. Postlarvae were acclimated to

water conditions in the aquaria (11.5 mg dissolved oxygen [DO] l⁻¹, 12.3°C, 36.6 salinity) by suspending the bags in an aquarium holding tank (100 l) for 40 min. Postlarvae were fed hatchery feed pellets and kept in the holding tanks for a period of 3 d before the commencement of the habitat preference trials and for 9 d before the commencement of the aquaria based predation trials. During this time, tanks were siphoned every second day to clean out unconsumed food and waste. The predation trials were repeated in the field in Wallagoot Lake, New South Wales (36°47'18" S, 149°56'31" E). Postlarvae, 18 d old and CL ~ 4 mm, were transported to the field site from the hatchery as described previously. Field experiments commenced immediately after postlarvae had acclimated to the conditions within experimental cages suspended in the lake (7.8 mg DO l⁻¹, 24.3°C, 35.7 salinity).

Habitat preference. The settlement patterns of *Penaeus plebejus* postlarvae into bare mud, bare sand and artificial macrophyte habitats were examined using a design similar to that described by Olabarria et al. (2002). Sand and mud were collected from a coastal bay near the aquaria facilities and placed in shallow bins to a depth of 4 cm. Seawater was then added to the bins to submerge the substratum entirely. The substratum was 'aged' in a sunlit area with continuous air supply over a 7 d period to allow a film of biota to develop on the surface. The artificial macrophyte habitat consisted of AquaMat®, a positively buoyant synthetic matting with a high microscopic surface area that is supplied divided into blades (shoots) (each blade was ~15 cm high and ~1.5 cm wide and there were ~67 blades per meter of AquaMat). AquaMat effectively simulates the blades and leaves of macrophytes in the wild (Arnold et al. 2005). At their base, the AquaMat blades were attached to weighted 30 cm lengths of the same synthetic matting. The AquaMat was placed within a natural, sunlit seagrass bed adjacent to the aquaria facilities for 21 d to facilitate colonization of epiphytes and a microbial film. Any macroinvertebrates that settled on the AquaMat during this time were located and removed before use in the experiment since these organisms may have included newly settled *P. plebejus* postlarvae from the wild and invertebrates capable of consuming the experimental postlarvae.

At the beginning of each experiment, each substrate type was established within 6 separate aluminum habitat trays (15 × 32 × 4 cm). Both mud and sand habitat trays were entirely covered with a 4 cm layer of their respective substrate types. When retrieved from the 'ageing' bins, these substrates were transferred to the habitat trays with minimal disruption to the surface layer to preserve the surface film of biota. Each macrophyte habitat tray had two 30 cm lengths of weighted

AquaMat base implanted into a 4 cm layer of sand, with the attached AquaMat blades (with equivalent shoot density of 837.5 shoots m⁻² and a surface area of 1.88 m² m⁻² of substrate) emerging above the sand. Each tray had two 50 cm twine handles to allow the trays to be removed from the tanks upon completion of experiments.

In each of 3 independent experimental tanks (50 l, 55 × 34 × 25 cm), one of each of the sand, mud and macrophyte habitat trays were placed along the tank (Fig. 1). These tanks were used to determine whether postlarvae occupied a particular habitat at higher densities when the alternative habitats were also available. In each experimental tank, 20 postlarvae were initially released over 1 tray only, either the sand, mud or macrophyte habitat tray. Three separate 50 l tanks were used as controls, each containing 3 habitat trays of only 1 habitat type (Fig. 1). These tanks allowed us to attribute greater occupancy of a particular habitat to the active behavior of preference to habitat type, as they eliminated the possibility of confounding preference with other artifacts (e.g. varied accessibility among the habitats) that may cause a greater number of postlarvae to remain within a habitat. Twenty postlarvae were initially released over only one of the habitat trays present in each control tank. During release and for the first hour after release, each tray within each tank was segregated from adjacent trays with 1 mm mesh netting. After the first hour, the mesh netting was removed so that the postlarvae could access all trays within a given tank.

Six replicate trials (n = 6) of the experiment were conducted (with renewed postlarvae in each trial). The placement of habitat trays along each tank was alternated among replicate trials so that all possible arrangements (total of 6) of the 3 habitat trays in a single tank were used. Treatments were randomly assigned to each tank during each trial and indepen-

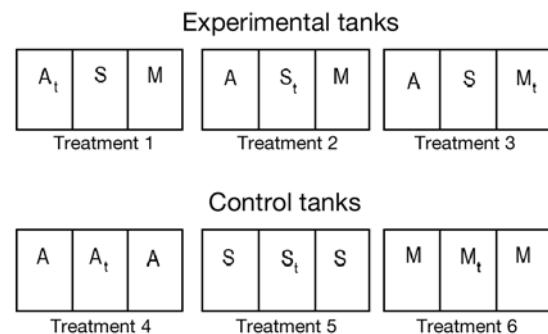


Fig. 1. *Penaeus plebejus*. Tanks used to test for preference of artificial macrophyte (A), bare sand (S) or bare mud (M) habitats by postlarval prawns. The subscript t associated with A, S and M indicates the habitat over which postlarvae were released at the beginning of the experiment for each tank

dent habitat trays (with renewed substrate and seawater) were used for each trial. Postlarvae were initially released into each tank at 06:00 h and the experiment ran for 12 h in an artificially illuminated environment to simulate daytime conditions. After 12 h, the habitat trays were segregated with the mesh netting, trays were removed from tanks and the number of postlarvae occupying each tray was counted. The entire series of trials was repeated in dark conditions (from 18:00 to 06:00 h) using new postlarvae and habitat trays.

Predation mortality. Laboratory experiment: Predation mortality of *Penaeus plebejus* postlarvae in sand, mud and macrophyte habitats was examined using adult eastern fortesque *Centropogon australis* (White, 1790) (9 to 10 cm total length [TL]), juvenile yellowfin bream *Acanthopagrus australis* (Owen, 1853) (12 to 15 cm TL) and late juvenile to sub-adult school prawns *Metapenaeus macleayi* (Haswell) (6 to 10 cm TL, 3.5 to 5 cm CL) as predators. Predators were sampled from the George's River, New South Wales (34° 00' 11.62" S, 151° 06' 27.61" E) with an otter trawl and transported to the aquaria facilities in separate aerated 80 l tanks where they were acclimated to aquarium conditions and released into a 2500 l (fish species) or a 100 l (*M. macleayi*) holding tank. Whilst in the holding tanks, the fish were fed commercially purchased prawn flesh every second day and the *M. macleayi* were fed a mixture of commercially purchased adult prawn flesh and hatchery pellets every second day. Tanks were siphoned every second day to remove unconsumed food and waste

Two 100 l tanks (25 cm radius, 50 cm high) were prepared with sand habitat, 2 with mud habitat and 2 with macrophyte habitat. Each habitat was collected and 'aged' as described previously. The sand and mud habitat were established to a depth of 4 cm on the base of each of their respective tanks. Sand was added to the base of the macrophyte habitat tanks to a depth of 4 cm and three 40 cm lengths of fouled AquaMat were then implanted into the sand with attached 25 cm high AquaMat blades emerging above (equivalent to a shoot density of 409.5 shoots m⁻² and a surface area of 1.53 m² m⁻² of substrate). One of the 2 tanks containing each habitat type was haphazardly assigned as the experimental tank and had 1 predator added to it at the beginning of each experimental trial. The remaining tank was the control tank and contained no predator. The control tanks were used to identify any variables other than predation that may cause mortality of postlarvae. There were 3 independent replicate tanks of each treatment and 2 replicate trials of the experiment (n = 6) were undertaken with each predatory species. Substrates, seawater and predators were renewed for each experimental trial and tanks were

haphazardly re-assigned to a treatment before each trial.

At the beginning of each experimental trial, 20 postlarvae were released into each tank and left to settle for 2 h, after which a single predator was added using a large fine-mesh dip net. The process of adding a predator to a tank was simulated in the control tanks by dipping an empty dip net that had previously contained a predator into the water for 10 s. The predators were removed from the experimental tanks after 24 h with a dip net and each tank was drained of water to determine the number of postlarvae remaining.

Field experiment: Nine juvenile *Centropogon australis* (6 to 9 cm TL) obtained from Merimbula Lake, New South Wales (36° 53' 50.89" S, 149° 53' 04.73" E) with a beach seine were used as predators. The predators were housed in 2 aerated tanks (~50 l) for a period of 78 h. During this time, the *C. australis* were fed commercially purchased adult prawn flesh every day. The water in each tank was renewed on a daily basis about 2 h after the predators had ceased feeding.

The field experiment was conducted in 6 cages (2 mm stretch mesh), each 1 m³ in size with an enclosed base, which were half-submerged in Wallagoot Lake. Sand, mud or macrophyte habitat were each added to 2 of the cages. Sand and mud were collected from Wallagoot Lake and deposited directly into the cages 5 d before the experiment began. The macrophyte habitat was prepared and established within the cages in the same manner and at similar quantities as described for the aquaria experiments, but the AquaMat was 'aged' in Wallagoot Lake for 38 d before the experiment. One of the cages containing each habitat type was haphazardly assigned as the experimental cage and the other was assigned as the control cage. There were 3 independent replicate cages for each treatment (n = 3).

Forty minutes after 50 postlarvae were released into each field cage and acclimated to the conditions therein, a single *Centropogon australis* was added to each experimental cage by means of a dip net. The process was simulated within the control cages. After 5 d, each cage was moved intact to the shoreline where its contents were emptied into a plastic container, preserved in formalin and transported to laboratory facilities. The *C. australis* from each experimental cage was captured and preserved separately. The contents of the containers were examined 3 wk later within the laboratory to quantify the proportion of postlarvae remaining at the end of the experiment.

Statistical analyses. All statistical analyses were carried out in Stat Graphics Plus 5. Cochran's test was used to examine homogeneity of variances among treatments for each experiment. Where variances dif-

ferred significantly ($p < 0.05$), the proportional data were arcsine transformed.

For the habitat preference experiments, *a priori* planned contrasts incorporated into a 1-way ANOVA (with 12 levels; each of macrophyte, sand or mud with the other habitats either present or absent in each of night and day conditions) were used to test the effects of habitat treatment on the proportion of postlarvae remaining within each release tray during the day and at night. Only the data from a single tray from each tank (the release tray) was used in the ANOVA, thus avoiding the problem of non-independence amongst treatments.

We predicted that the macrophyte habitat would be the habitat preferred by postlarvae and would be seen in either diurnal period. Three alternative hypotheses, which were tested using the *a priori* planned contrasts, had to be supported for each period (Table 1). The first hypothesis (H_1) stated that if released into a tray with one of macrophyte, sand or mud habitats and given a choice to move into 1 of the 2 alternative habitats (Treatments 1, 2 and 3, respectively, in Fig. 1), fewer postlarvae would move out of the macrophyte tray compared with the sand and mud trays. H_1 also predicted that when released into a macrophyte tray and given no choice of other habitats in which to move (i.e. when all thirds of a tank consisted of macrophyte trays) (Treatment 4 in Fig. 1), fewer postlarvae would actively move out of the macrophyte tray-of-release compared with the number moving out of the release trays in Treatments 2 and 3 in Fig. 1. The last 2 hypotheses (H_2 and H_3) predicted that more postlarvae would move out of sand and mud release trays, respectively, when the macrophyte habitat was also available (Treatments 2 and 3 in Fig. 1) compared with when each of the sand and mud habitats were available alone (Treatments 4 and 6 in Fig. 1). The residual mean

squares (MS) used in the planned contrasts were taken from the omnibus 1-way ANOVA.

For the laboratory predation experiments, three 2-way ANOVAs were used to analyze the survival data for each predatory species separately. Both factors in the ANOVAs, habitat (3 levels: sand, mud or macrophyte) and predator treatment (2 levels: present or absent), were fixed. For the field based experiments, one 2-way ANOVA with the same fixed factors was used to analyze the survival data. Each ANOVA tested 2 alternative hypotheses that (1) proportion of surviving postlarvae within each habitat would be dependent on the presence or absence of predators, and (2) in the presence of predators this proportion would be higher in the macrophyte habitat compared with the sand and mud habitats. Where an interaction between predator treatment and habitat was detected, Tukey's HSD test was used to determine which treatments differed from each other.

RESULTS

Habitat preference

For the night and day experiments 100 and 99.4 %, respectively, of the postlarvae released into all tanks were recovered. Variances were homogeneous amongst all release trays used in the experiment (Cochran's test: $C = 0.23$, $n = 6$, $k = 12$, $p > 0.05$). Similar proportions of postlarvae remained in each habitat's release tray at night (ranging from 0.3 to 0.35), irrespective of whether postlarvae were given a choice of habitats to move into (Fig. 2). During the day, how-

Table 1. *Penaeus plebejus*. Hypotheses used to test for habitat preference and examine effects of diel period on preference patterns. n_{A_i} , n_{S_i} or n_{M_i} represent the number of postlarvae left at the end of the experiment in the habitat tray where they were initially released (artificial macrophyte [A_i], sand [S_i] or mud [M_i], respectively); N_i is the total number of postlarvae recovered from a treatment tank at the end of the experiment and $i = 1$ to 6 indicates the treatment tank (see Fig. 1)

Alternative hypothesis	Test
H_1	$\frac{n_{A_1}}{N_1}, \frac{n_{A_4}}{N_4} > \frac{n_{S_2}}{N_2}, \frac{n_{M_3}}{N_3}$
H_2	$\frac{n_{S_2}}{N_2} < \frac{n_{S_5}}{N_5}$
H_3	$\frac{n_{M_3}}{N_3} < \frac{n_{M_6}}{N_6}$

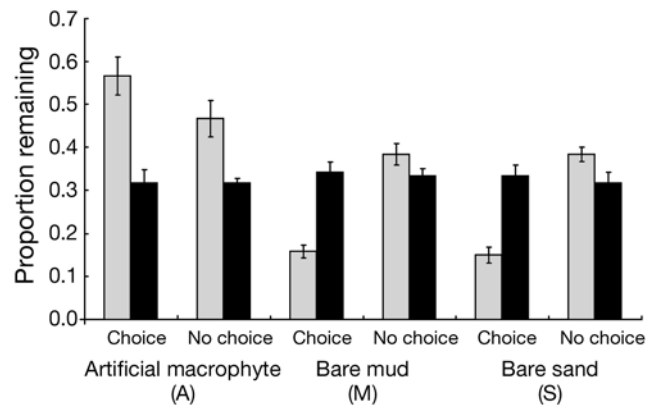


Fig. 2. *Penaeus plebejus*. Average (\pm SE) proportion of postlarvae remaining in artificial macrophyte (A), bare mud (M) or bare sand (S) after release into each of these habitats either when each was present with equal amounts of the 2 alternate habitats also available (Choice) or when each habitat was the only habitat present (No choice). The plot contrasts the behavior of the postlarvae in simulated daytime (grey bars) and nighttime (black bars) conditions

ever, release trays within the different treatment tanks had different proportions of postlarvae remaining in them (Fig. 2). The proportion was highest in the macrophyte release trays, irrespective of whether postlarvae were given alternative habitats to move into. Intermediate proportions remained in the sand and mud release trays when the alternative habitats were absent. The lowest proportions were observed in sand and mud release trays when the alternative habitats were also present. Consistent with this overall pattern, the planned contrasts revealed that H_1 , H_2 and H_3 were supported during the day, but not at night when there was no significant difference between the habitat treatments involved in each of the planned comparisons (Table 2, Fig. 2). With the exception of each of the sand and mud release trays that were available with no choice of alternative habitats, all daytime trays were dissimilar from their nighttime equivalents (Fig. 2).

Predation mortality

Whilst the treatments in the laboratory experiment in which *Acanthopagrus australis* were used as predators had homogenous variances (Cochran's test: $C = 0.31$, $n = 6$, $k = 6$, $p > 0.05$), variances were significantly different for the laboratory experiment in which *Centropogon australis* were used as predators (Cochran's test: $C = 0.69$, $n = 6$, $k = 6$, $p < 0.05$). These data were consequently arcsine transformed to produce a data set with homogeneous variances (Cochran's test: $C = 0.29$, $n = 6$, $k = 6$, $p > 0.05$). There was a significant interaction between habitat and predator presence/absence in the *A. australis* and *C. australis* laboratory trials (Table 3). Survival of postlarvae was close to 100% in all control tanks and varied among habitats in the experimental tanks, being significantly higher (55 to 60% survival) in the macrophyte habitat tanks com-

pared with the sand and mud tanks (15 to 25% survival; Table 3, Fig. 3a,b). When *Metapenaeus macleayi* were used as predators in the laboratory, variances were not significantly different among treatments (Cochran's test: $C = 0.39$, $n = 6$, $k = 6$, $p > 0.05$). There was also no significant interaction between habitat and predator presence/absence (Table 3, Fig. 3c). Although survival was significantly higher in the con-

Table 2. *Penaeus plebejus*. Omnibus ANOVA and planned contrasts ($\alpha = 0.05$) comparing average proportions of postlarvae remaining in 12 treatments. The different treatments were: Day A_i 1, Day S_i 2, Day M_i 3, Day A_i 4, Day S_i 5, Day M_i 6, Night A_i 1, Night S_i 2, Night M_i 3, Night A_i 4, Night S_i 5 and Night M_i 6, where A_i , M_i and S_i were trays with artificial macrophyte, bare mud or bare sand, respectively, over which postlarvae were initially released when they had either a choice of habitats to move between (1 to 3) or no choice (4 to 6) in either daytime (Day) or nighttime (Night) conditions. Contrasts 1 and 4 tested the hypotheses that Day A_i 1 and Day A_i 4 > Day S_i 2 and Day M_i 3, and Night A_i 1 and Night A_i 4 > Night S_i 2 and Night M_i 3, respectively;. Contrasts 2 and 5 tested the hypotheses that Day S_i 2 < Day S_i 5 and Night S_i 2 < Night S_i 5, respectively. Contrasts 3 and 6 tested the hypotheses that Day M_i 3 < Day M_i 6 and Night M_i 3 < Night M_i 6, respectively. p-values in **bold** denote statistically significant differences; $n = 6$ for each treatment

	df	MS	F	p
Omnibus ANOVA				
Groups	11	0.077	18.65	< 0.05
Residual	60	0.004		
Total	71			
Planned contrasts				
Contrast 1	1	0.79	189.85	< 0.05
Contrast 2	1	0.16	39.33	< 0.05
Contrast 3	1	0.15	36.57	< 0.05
Contrast 4	1	0.003	0.63	> 0.05
Contrast 5	1	8.33×10^{-4}	0.20	> 0.05
Contrast 6	1	2.08×10^{-4}	0.05	> 0.05

Table 3. *Penaeus plebejus*. Two-way ANOVAs ($\alpha = 0.05$) comparing the average number of postlarvae surviving in different habitats in the presence and absence of predators. *Metapenaeus macleayi*, *Acanthopagrus australis* or *Centropogon australis* were used as predators in the laboratory and *C. australis* were used in the field. The habitats compared were simulated macrophyte, bare mud and bare sand. p-values in **bold** denote statistically significant differences; $n = 6$ for each laboratory treatment and $n = 3$ for each field treatment

	<i>A. australis</i> (laboratory)				<i>C. australis</i> (laboratory) ^a				<i>M. macleayi</i> (laboratory)				<i>C. australis</i> (field)			
	df	MS	F	p	df	MS	F	p	df	MS	F	p	df	MS	F	p
Predator (P)	1	3.87	744.6	< 0.05	1	35.012	351.92	< 0.05	1	0.54	165.47	< 0.05	1	684.50	40.13	< 0.05
Habitat (H)	2	0.11	20.98	< 0.05	2	891.85	8.96	< 0.05	2	0.00	0.06	> 0.05	2	134.00	7.86	< 0.05
H × P	2	0.11	21.75	< 0.05	2	521.47	5.24	< 0.05	2	0.00	0.28	> 0.05	2	92.67	5.43	< 0.05
Residual	30	0.005			30	99.49			30	0.00			12	17.06		
Total	35				35				35				17			

^aData for *C. australis* (laboratory) were arcsine transformed for analysis

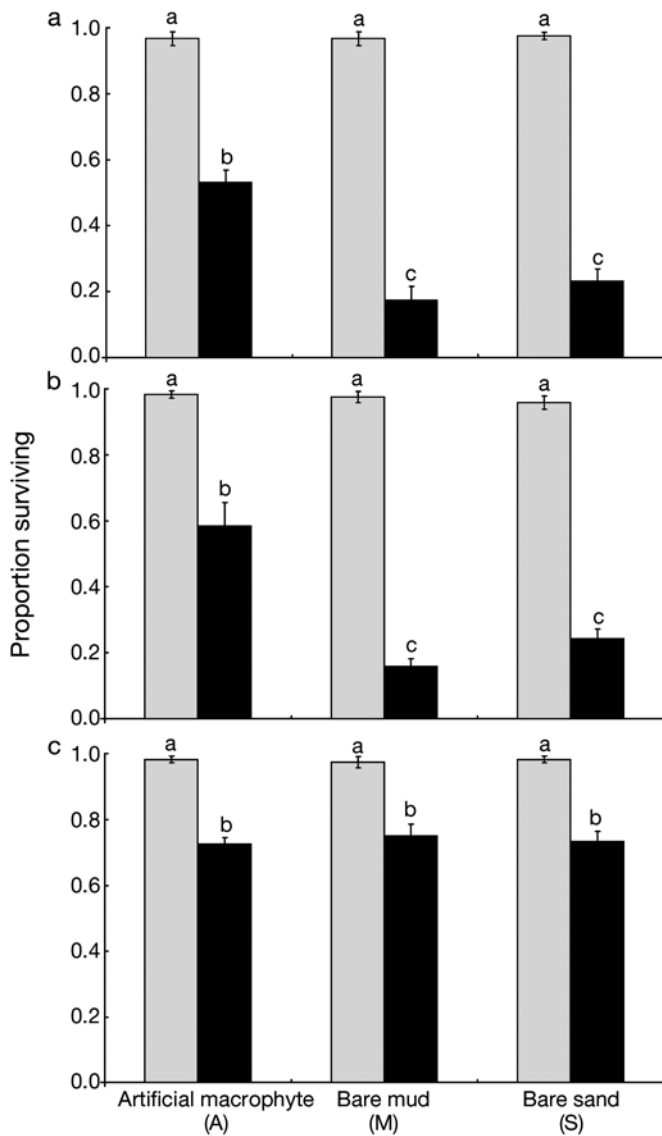


Fig. 3. *Penaeus plebejus*. Average (\pm SE) proportion of postlarvae surviving in laboratory tanks with artificial macrophyte, bare mud or bare sand in absence (grey bars) and presence (black bars) of (a) *Acanthopagrus australis*, (b) *Centropogon australis* or (c) *Metapenaeus macleayi* as predators. Letters a, b and c ($a > b > c$) each represent groups of treatments that were statistically homogeneous according to Tukey's HSD tests ($\alpha = 0.05$, $n = 6$)

trol tanks (~95%) compared with the experimental tanks (~75%), similar proportions of postlarvae survived across all habitats in the presence of predators (Table 3, Fig. 3c).

For the field-based experiments, variances were homogeneous amongst treatments (Cochran's test: $C = 0.26$, $n = 6$, $k = 6$, $p > 0.05$) and there was a significant interaction between habitat and predator presence/absence (Table 3). Tukey's HSD test showed that in the absence of *Centropogon australis*, all habitat types had

similar proportions of postlarvae surviving within them (60 to 75% survival); however, in the presence of *C. australis*, significantly more postlarvae (~60%) survived in the macrophyte cages than in the sand and mud cages, which had similar survival rates (~30%; Fig. 4).

DISCUSSION

Habitat complexity can affect ecological interactions directly by facilitating coexistence among competitors and influencing the success of predators and prey, or indirectly by influencing behavior and, thus, altering the outcome of interactions (Crowder & Cooper 1982, Brown 1988). This study showed that complexity affects predation mortality and settling behavior in postlarval *Penaeus plebejus*.

Habitat preference

Penaeus plebejus did not show a preference for complex habitat during the night, with bare sand, bare mud and artificial macrophyte retaining similar numbers of postlarvae, regardless of whether alternative habitats were available. In contrast, when postlarvae were released into each habitat during the day and able to move between different habitats, higher proportions moved out of the non-complex habitats, sand and mud, than into the macrophyte habitat, suggesting a preference for macrophytes. A larger number of postlarvae left the bare habitats when the macrophyte habitat was also available compared with the number

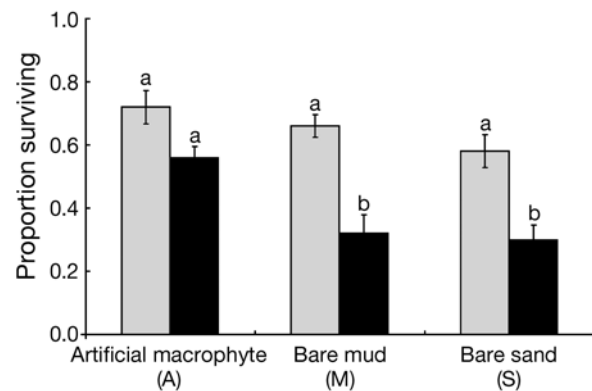


Fig. 4. *Penaeus plebejus*. Average (\pm SE) proportion of postlarvae surviving in field cages with artificial macrophyte (A), bare mud (M) or bare sand (S) in the absence (grey bars) or presence (black bars) of *Centropogon australis* as predators. Letters a and b ($a > b$) each represent groups of treatments that were statistically homogeneous according to Tukey's HSD tests ($\alpha = 0.05$, $n = 3$)

that left the macrophyte habitat when it was the only one available, providing further evidence of a greater aversion to the bare habitats relative to the macrophyte habitat.

Several studies have attempted to examine habitat preference in other penaeids (e.g. Hughes 1966, Macia 2004, Gribble et al. 2007), but their methods tested for association with habitat rather than preference per se (see Underwood et al. 2004 for distinction). Key exceptions include studies on juvenile *Penaeus aztecus*, *P. esculentus* and *P. semisulcatus*, which display similar affinities for complex habitats as reported here, studies on juvenile *P. setiferus*, which exhibit no preference for either vegetated or un-vegetated habitat (Minello & Zimmerman 1985, Hill & Wassenberg 1993, Kenyon et al. 1997), and studies on juvenile *Melicertus latisulcatus*, which selected bare sand over vegetated habitat regardless of diel period (Tanner & Deakin 2001). These varied results indicate that there is no common pattern among penaeids in general.

Habitat preference can play an important role in the survival of species as it can optimize the use of resources or limit negative interactions (Huey 1991, Real 1991, Rosenzweig 1991). For *Penaeus plebejus*, preference may reduce exposure to predators. Since many of the fish that feed on penaeids are visual daytime hunters (Brewer et al. 1995) and complex habitats can reduce their foraging efficiency (Heck & Thoman 1981), postlarval *P. plebejus* may be selecting macrophyte beds when they are most at risk of predation (i.e. daytime). As penaeids increase in size, however, burrowing into sand may become a more effective means of predator avoidance than hiding among macrophytes (Hill & Wassenberg 1993, Liu & Loneragan 1997), resulting in an ontogenetic change in preference in favor of bare sand. Although this change may explain the distinct lack of preference for complex habitat in juvenile *P. setiferus* and *Melicertus latisulcatus* (Minello & Zimmerman 1985, Tanner & Deakin 2001), a number of other penaeids still select structured habitats over bare habitats as juveniles (Kenyon et al. 1997). Therefore, ontogenetic shifts in preference vary among penaeids and must be independently assessed for *P. plebejus*.

Although the diurnal habitat use patterns of postlarval *Penaeus plebejus* combined with our predation results suggest habitat preference is driven by predator avoidance, we cannot conclude that this is the only mechanism at play. Penaeid feeding activity is mainly nocturnal (Primavera & Leбата 1995), with macrophytes and their epiphytes often contributing to penaeid diets (Kitting et al. 1984, Loneragan et al. 1997). Optimal foraging theory and a cost/benefit function incorporating predation risk and habitat profitability consequently predict that a postlarva will select macrophyte beds during the day to maximize energy

intake per unit of time and simultaneously lower predation risk. During the night, however, reduced predator efficiency may preclude the need to shelter within macrophytes, thereby allowing the postlarva to forage in various habitats for alternative sources of food (MacArthur & Pianka 1966, Sih 1980, Werner et al. 1983). Experiments examining selection of complex habitats by *P. plebejus* in the presence and absence of macrophyte-based food sources must be conducted to assess this theory. Another plausible explanation for the diurnal change in habitat use by postlarvae is an inability to perceive the visual cues required to select protective habitat in darkness leading to random settlement at night. However, penaeids are generally tactile and have well developed chemoreceptors (Zimmer-Faust 1989, Dall et al. 1990a), and these attributes may make them adept at perceiving surrounding habitat in the absence of visual reception.

Predation mortality

Predation on *Peneaus plebejus* postlarvae by *Centropogon australis* and *Acanthopagrus australis* was lower in artificial macrophytes compared with habitats without physical structure. These findings were consistent with studies on other members of Penaeidae (Minello & Zimmerman 1985, Kenyon et al. 1995, Primavera 1997) and Caridea (Coen et al. 1981, Heck & Thoman 1981). For penaeids and many other prey species, complex habitats limit predation by providing refuge or by reducing predatory foraging efficiency (Laprise & Blaber 1992, Kenyon et al. 1995, Almany 2004). Postlarval *P. plebejus* were observed remaining motionless amongst vegetative structures in the presence of predators. This behavior may provide the best protection against predators during this life stage because burrowing into substrate to avoid detection by predators is not as prevalent in penaeid postlarvae (Hill & Wassenberg 1993). Only 0.4 and 2.5% of the postlarvae used in our habitat preference and predation experiments, respectively, were observed burrowing.

When juvenile *Metapenaeus macleayi* were used as predators, the survival of *Peneaus plebejus* did not differ between complex and non-complex habitats. Where similar results have been recorded for other penaeids, aspects of the chosen predator's physiology and behavior may make it equally or more adept at detecting and capturing prey within complex habitats. Predation mortality in *P. aztecus* was unaffected by the presence of vegetative structure for predators that used non-visual chemosensory mechanisms to detect their prey (Minello & Zimmerman 1983, Primavera 1997). Crustaceans such as *M. macleayi* possess sensi-

tive chemoreceptors on their antennae, pereopods and mouthparts with an olfactory function (Zimmer-Faust 1989, Dall et al. 1990b). These receptors trigger foraging responses such as walking faster and probing into the substratum (Carr & Derby 1986, Dall et al. 1990b) whenever prey are detected in the absence of visual cues. Compared with *Acanthopagrus australis* and *Centropogon australis*, whose chemosensory foraging abilities have not been previously documented, *M. macleayi* may have been less reliant on visual detection of prey due to their chemosensory abilities and, therefore, were able to forage effectively in macrophyte habitat. Predation by passive ambush predators, such as *C. australis* (Harmelin-Vivien & Bouchon 1976) and *A. australis* (Shand et al. 2000), can also be significantly reduced by the presence of structure. In contrast, predators that chase their prey may be unaffected by complexity because they squeeze between structures and occupy the narrow crevices where prey attempt to hide (Primavera 1997). This active pursuit behavior has been documented in penaeid species (Racek 1959, Dall et al. 1990b) and may have added to the efficiency of *M. macleayi* as predators in complex habitats.

Implications for stock enhancement

Stock enhancement is one of the more expensive and least successful fishery management options available (Grimes 1998, Kellison et al. 2003). To reduce the ecological uncertainties that often limit its success, a candidate release species' optimal habitat must be known together with this habitat's carrying capacity and how to minimize losses through predation (Rothlisberg 1998). The present study provides important information on these elements for *Peneaus plebejus*.

Macrophyte beds provide optimal habitat for postlarval *Peneaus plebejus* and habitat structure can significantly reduce post-release predation mortality. Therefore, enhancement of *P. plebejus* must be undertaken in lakes and estuaries with extensive macrophyte beds. Experiments similar to those done here in a range of macrophyte habitats may help determine exactly which types of macrophytes best support the species. If *P. plebejus* is to be grown to larger sizes before release, habitat selection at these sizes must be investigated and the release habitats selected accordingly. Potential predators within a system should also be identified before enhancement. If predators such as *Acanthopagrus australis* and *Centropogon australis* are present at high densities relative to macrophyte cover, enhancement must be reassessed based on rigorous analyses that take into account potential losses through predation. If wild penaeids such as *Metapenaeus macleayi*

exist at high densities, predation on released postlarvae may be significant across all habitats and restocking may only be viable if *P. plebejus* postlarvae are released into lakes and estuaries during the *M. macleayi* offshore migratory period or if *P. plebejus* are cultured to a size at which they will no longer be vulnerable to predation by large penaeids. If other potential predators are identified within target systems, experiments should be conducted to determine whether these predators will limit the survival of *P. plebejus* in the available habitat.

Stock enhancement efforts are slowly undergoing a paradigm shift from poorly investigated and hurried mass releases, to releases based on scientific theory and precautionary accountability (Leber 2002). This shift must continue to incorporate comprehensive pilot assessments of the ecological and life history patterns of candidate release species, previous investigations into the habitat characteristics of the targeted release sites and comparisons of the relative fitness of wild and stocked animals (Blankenship & Leber 1995, Araki et al. 2007). Such studies will allow the financial costs and predicted ecological consequences of an enhancement program to be weighed against its projected economic, biological and social benefits and, therefore, assist in determining whether stock enhancement is a better response to fisheries problems than reduction of fishing effort or restoration of habitats.

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