Size-specific locomotion rate and movement pattern of four common Indo-Pacific sea stars (Echinodermata; Asteroidea)

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ABSTRACT: The ecology of sea stars appears to be related to their locomotive abilities. This relationship was studied for the sea stars Acanthaster planci, Archaster typicus, Linckia laevigata, and Protoreaster nodosus in the coastal waters of Samal Island, the Philippines between May and July 2008. In order to avoid the sensory interruptions that sea stars exhibit when moving across natural substrate, a tarpaulin (2 × 2 m) was placed on the seafloor to create a uniform habitat. Mean (±SD) locomotion rate of Archaster typicus was 45.8 ± 17.0 cm min⁻¹ but increased with mean radius (R). Locomotion rate increased from 17.8 to 72.2 cm min⁻¹ for specimens with R of 1 and 5 cm respectively. Mean locomotion rate of L. laevigata, P. nodosus, and Acanthaster planci was 8.1 ± 1.9, 18.8 ± 3.9, and 35.3 ± 10.0 cm min⁻¹ respectively, and was not related to R. Maximum speed of Archaster typicus was 76.0 ± 31.9 cm min⁻¹ and also increased with R. Maximum speed of L. laevigata, P. nodosus, and Acanthaster planci was 10.6 ± 2.7, 27.7 ± 7.5, and 50.9 ± 14.0 cm min⁻¹ respectively. An inter-specific comparison of locomotion rates using the number of body moves per minute suggests that the effect of body size partially overrules morphological and behavioral adaptations. The movement patterns of Acanthaster planci, L. laevigata, and P. nodosus were highly directional (ratio of displacement to total distance traveled [D:Wₜₜ] ≥ 0.86), whereas Archaster typicus displayed a more random movement pattern (D:Wₜₜ = 0.51). Water-current direction did not affect movement directions of the 4 species.

KEY WORDS: Acanthaster planci · Archaster typicus · Davao Gulf · Linckia laevigata · Protoreaster nodosus · Starfish · Speed

INTRODUCTION

The ability to migrate and the speed of migration has multiple implications for the biology and ecology of an animal (e.g. Hedenström 2003, Dingle & Drake 2007). Benthic reef organisms such as sea stars, that move relatively slowly, may have to spend their entire life at the reef where they once settled. This implies that such organisms will reproduce in this spatial area and that a pelagic larval phase provides the only opportunity for long-distance dispersal (Williams & Benzie 1993). Despite being relatively slow-moving animals, most Indo-Pacific sea stars have comparatively large distribution areas. The larval periods of the Indo-Pacific sea stars Acanthaster planci (Linnaeus 1758) (Moran 1988), Linckia laevigata (Linnaeus 1758) (Williams & Benzie 1993), and Protoreaster nodosus (Linnaeus 1758) (Bos et al. 2008a) are approximately 3 to 4 wk and provide the potential for long-distance dispersal.

Sea stars move intermittently to search for food or to avoid physical stress (Feder & Christensen 1966). For example, Archaster typicus (Müller et Troschel 1840) and Acanthaster planci may display locomotion rates of 15.1 to 24.9 cm min⁻¹ (Run et al. 1988) and 33.3 cm min⁻¹ (Moran 1990) respectively. Maximum speed in sea stars was assumed to be based on the positive rela-
tionship between body size, and number (Gaymer & Himmelman 2008) and length of tube feet (e.g. Westheide & Rieger 1996). Hence, larger specimens move faster than smaller ones, both inter-specifically and intra-specifically. Maximum speed may also be a measure of the ability to avoid predation, e.g. for Archaster typicus. This sea star inhabits similar habitats as the larger omnivorous species Protoreaster nodosus, but does not share habitats with the latter. While Archaster typicus inhabits intertidal habitats such as sandy shores and seagrass meadows, juvenile P. nodosus are exclusively found in the adjacent shallow subtidal sand and seagrass areas (Bos et al. 2008a,b). This could be an indication that Archaster typicus avoids predation by P. nodosus (Bos et al. 2011). In contrast, the 2 species occurring in shallow soft-bottom habitats, Linckia laevigata and the multi-rayed corallivore sea star Acanthaster planci, are commonly encountered from the intertidal up to a depth of 60 m (Moran 1990, Schoppe 2000).

Locomotion rates and movement patterns of sea stars have been determined by marking tracks of individuals on natural substrates (e.g. Ferlin 1973, Scheibing 1981, Himmelman et al. 2005). However, limited observation time results in poor resolutions in time and space. Recently developed techniques such as time-lapse video analysis (e.g. Dale 1997, Lauzon-Guay et al. 2006, Dumont et al. 2007, McClintock et al. 2008a, b) provide researchers with highly accurate data but require costly equipment. The use of an artificial substrate attached to the sea floor combines the advantage of observations under natural conditions and the avoidance of movement interruptions when sea stars encounter prey, which is likely to happen on natural substrates.

The present study aims to determine the size-specific locomotion rate and maximum speed and describe the movement patterns of 4 common Indo-Pacific sea stars: Acanthaster planci, Archaster typicus, Linckia laevigata, and Protoreaster nodosus.

**MATERIALS AND METHODS**

**Sea star collection and measurements.** Sea stars were collected in the coastal waters of Samal Island in the Davao Gulf, The Philippines from May to July 2008 (Acanthaster planci: 4 June to 2 July, Archaster typicus: 5 to 20 May, Linckia laevigata: 11 to 26 June, and Protoreaster nodosus: 5 May to 2 July). Collection was usually done by means of snorkeling, whereas the collection of larger specimens of L. laevigata and Acanthaster planci was done using SCUBA equipment. Since sea stars cease movement after handling (Dale 1997), specimens were acclimated for 2 to 12 h in a submerged net-covered container (45 × 20 × 30 cm) to minimize handling stress.

Radius (distance from arm tip to center of oral disk) of each arm was measured with calipers (accuracy: 1 mm) at the oral side to calculate mean radius (R). In the case of Acanthaster planci, the diameter (= 2R) was measured with a ruler (accuracy: 5 mm) at the aboral side before collection, because this species curls itself when handled.

Specimens of Archaster typicus (R = 1 to 5 cm) were categorized in nine 0.5 cm size classes. Specimens of the other 3 species were categorized in 1 cm size classes, due to their larger sizes (Acanthaster planci: R = 4 to 19 cm, Linckia laevigata: R = 4 to 15 cm, Protoreaster nodosus: R = 2 to 14 cm).

**Experimental design and observations.** A tarpaulin (2 × 2 m) with a printed grid (10 × 10 cm squares) was laid on the surface of the substratum. Squares were vertically lettered and horizontally numbered (Fig. 1). The tarpaulin, weighted with a metal chain along its entire circumference, was firmly attached to the sea floor with pegs in an intertidal sandy area (grid always facing north). Observations were performed in water with depth ranging from 0.3 to 2.0 m while we snorkeled above the tarpaulin.

Each sea star was carefully positioned in the center of the tarpaulin always with its madreporite to the south. This orientation was not possible for Acanthaster planci, which is a multi-madreporite species. As soon as a sea star started moving, we started a stopwatch (accuracy: 1 s). Time and square label were recorded each time an individual reached another square of the grid. An observation was terminated when a sea star reached the edge of the tarpaulin or after performing a traveling path of >20 squares. Slow movement of Linckia laevigata made it necessary for us to terminate observations after measuring 10 consecutive squares. Due to difficulties in recording the exact traveling path on a slate, an observation was also terminated when a sea star visited a particular square a third time.

The following environmental parameters were recorded before each observation: (1) water depth (estimated in 5 cm intervals), (2) bottom water temperature (0.5°C accuracy), (3) wave activity (0 = no waves, 1 = small waves, 2 = medium waves, 3 = choppy), (4) water current speed (measured with float and stopwatch in cm s⁻¹), and (5) water current direction (Table 1).

**Data analysis.** Displacement of sea stars was calculated trigonometrically for each square of the tarpaulin. Sea star velocity was calculated by dividing the displacement (D) by the time needed to move to a neighboring square. An observation started when a sea star began its traveling path in the center of the tarpaulin (starting point, S) and lasted until it left the tar-
paulin (ending point, E) (Fig. 1). Therefore, one observation consisted of a number of consecutive sections. Locomotion rate was determined by calculating the mean speed of all sections within one observation, whereas maximum speed was the highest velocity.

Body moves per minute were calculated by dividing locomotion rate by the diameter ($2R$) for each specimen to study inter-specific locomotive differences independent of size. The pattern of movement was characterized as the ratio of $D$ to total distance traveled ($W$) (Ferlin 1973 modified by Scheibling 1981). $D$ is the shortest distance between S and E, whereas $W_{\text{all}}$ is the total distance traveled by a specimen during one observation (Fig. 1). An individual moving in a straight line displays a directional movement pattern and has a $D:W_{\text{all}}$ of 1. The more the actual movement path deviates from a straight line, the smaller $D:W_{\text{all}}$ and the less directed the movement pattern becomes. $D:W_{\text{all}}$ values >0.7 are considered to be highly directional, values >0.5 partially directional, whereas values <0.5 are considered to be undirected (Ferlin 1973, Scheibling 1981). $D:W_5$ was calculated by only considering the first 5 squares of the observations to test for size-dependent effects within the movement patterns.

Differences in mean locomotion rate, maximum speed, and $D:W_{\text{all}}$ and $D:W_5$ ratios were tested with 1-way ANOVA if homogeneity of variance was shown (Levene’s test). In case of heterogeneity, a Welch ANOVA was used. Post hoc analysis was done with Tukey’s HSD test.

The influence of water currents on the movement direction of sea stars was tested by calculating the deviation of the movement direction from the direction of the current. The Rayleigh test (Zar 1999) was used to determine whether the angles between movement direction and direction of the current were uniformly distributed around the 360° circle (which is expected when there is no influence of currents on the movement direction).

**RESULTS**

*Linckia laevigata*’s locomotion rate was the lowest, with a mean of 8.1 cm min$^{-1}$, while *Protoreaster nodosus* and *Acanthaster planci* moved at approximately double and 4 times that rate (Table 2). *Archaster typicus* was the fastest-moving sea star, with a mean locomotion rate of 45.8 cm min$^{-1}$ (Table 2).

*Archaster typicus*’ locomotion rate significantly increased with $R$ from 17.8 to 72.2 cm min$^{-1}$ (Fig. 2). Similarly, a significant linear relationship was found between maximum speed and $R$ for *A. typicus*. There was no significant relationship between both mean and maximum locomotion rates and $R$ for the other 3 species (Fig. 2).

The number of body moves per minute was interspecifically related to $R$ (Fig. 3). Small specimens ($R < 2$ cm) performed 6 to 12 body moves min$^{-1}$, whereas specimens with $R > 5$ cm could only perform <4 body moves min$^{-1}$.

<table>
<thead>
<tr>
<th>Species</th>
<th>Depth (cm)</th>
<th>Temperature (°C)</th>
<th>Current velocity (cm s$^{-1}$)</th>
<th>Wave action</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Range</td>
<td>Mean ± SD</td>
<td>Range</td>
<td>Mean ± SD</td>
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<tr>
<td><em>Acanthaster planci</em></td>
<td>30–185</td>
<td>29.5–31.0</td>
<td>0–6.3</td>
<td>5.4 ± 1.3</td>
</tr>
<tr>
<td><em>Archaster typicus</em></td>
<td>40–190</td>
<td>nd</td>
<td>0–11.7</td>
<td>5.0 ± 2.9</td>
</tr>
<tr>
<td><em>Linckia laevigata</em></td>
<td>65–170</td>
<td>29.5–30.5</td>
<td>0–14.5</td>
<td>7.8 ± 4.5</td>
</tr>
<tr>
<td><em>Protoreaster nodosus</em></td>
<td>30–185</td>
<td>29.5–32.0</td>
<td>0–8.9</td>
<td>5.0 ± 2.2</td>
</tr>
</tbody>
</table>
The mean \( D:W \) ratio of \textit{Acanthaster planci}, \textit{Linckia laevigata}, and \textit{Protoreaster nodosus} ranged from 0.86 to 0.89 (Table 2), whereas \textit{Archaster typicus}' mean \( D:W \) ratio of 0.51 was significantly lower (Tukey's test, \( p < 0.05 \)). No significant difference was found among the other 3 species (ANOVA, \( p > 0.05 \)).

The mean \( D:W \) ratio of \textit{Acanthaster planci}, \textit{Linckia laevigata}, and \textit{Protoreaster nodosus} ranged from 0.88 to 0.93 and for \textit{Archaster typicus} it was 0.68. \textit{Acanthaster planci}'s and \textit{Archaster typicus}' mean \( D:W \) ratio was significantly higher than their \( D:W \) ratio (Tukey's test, \( p < 0.05 \)). No significant difference was found for the other 2 species (ANOVA, \( p > 0.05 \)).

The distribution of the deviation angles between the movement direction and the direction of water currents showed that there was no preferential angle of movement (Fig. 4). The angle distribution was not significantly different from uniformity (Rayleigh test, \( p > 0.05 \)).
DISCUSSION

Substrates may affect locomotion rates and movement pattern of sea stars. Ferlin (1973) reported that *Astropecten aranciacus* moved at higher velocities on fine sand than on coarse sand. In contrast, Scheibling (1981) found that the locomotion rate of *Oreaster reticulatus* did not vary between fine and coarse sand. Velocities measured on tarpaulin could therefore differ from values measured on natural substratum. However, the mean locomotion rate of *Acanthaster planci* of 35.3 cm min\(^{-1}\) (Table 2) closely resembles the rate of 33 cm min\(^{-1}\) on sand reported by Moran (1990). Therefore the use of a tarpaulin to compare inter- and intra-specific movement of sea stars seems to be a suitable in-field method, even if results must be extrapolated to the natural environment with some caution.

Movement patterns of animals are an integrated component of their feeding strategy. Pyke et al. (1977) stated in the optimal foraging theory that natural selec-
tion may optimize movement patterns in relation to distribution and abundance of prey over evolutionary times. The goal of optimal foraging is to maximize the net rate of food uptake (Pyke et al. 1977, Krebs 1978). A highly directional movement pattern, as earlier described for sea stars (Scheibling 1981), optimizes foraging efficiency on substrata containing a relatively uniform distribution of food resources because it prevents re-grazing of substrata and extends the foraging range (Scheibling 1981). This seems to be the case for Acanthaster planci, Linckia laevigata, and Protoreaster nodosus, which all had a D:W all of approximately 0.9. Their movement was highly directional, whereas the movement path of Archaster typicus showed a partially directional pattern, with a D:W all of about 0.5.

Inter-specific interactions have the potential to determine sea-star foraging behavior under natural conditions (Himmelman & Dutil 1991, Morissette & Himmelman 2000, Gaymer et al. 2002, Gaymer & Himmelman 2008). In the present experiment, however, these were excluded because specimens were placed on the tarpaulin one at a time.

Beddingfield & McClintock (1993) found that Astropecten articularis exhibited a directional pattern of movement in the absence of food. When encountering prey, however, both the frequency and magnitude of angular deviations in the foraging path increased, resulting in intensified foraging in areas with high prey density. This seems to be a highly efficient foraging strategy that may be used by many organisms. Generally, undirected movement patterns may be an adaptation to living in high density in an environment with locally high prey density (McClintock & Lawrence 1985). This may be the case for Archaster typicus, which has densities of up to 131 ind. m$^{-2}$ (Bos et al. 2011) in the prey-abundant intertidal zone.

The relative small body size of Archaster typicus (up to 15 cm in diameter; Janssen et al. 1984) compared to the size of the other 3 sea stars in the present study may also explain the more undirected pattern of movement of this species. Sea stars may generally show directed movement for a certain distance that is related to their body size before changing their movement direction. Consequently, small sea stars would move a shorter distance before changing direction than larger ones. In fact, when just considering the first 5 squares of our observation (D.W.), Archaster typicus’ movement path was more directed. This was also the case for Acanthaster planci, which is much larger than Archaster typicus. In contrast, Linckia laevigata’s and Protoreaster nodosus’ D.W ratios do not differ significantly from their D.W all ratios, although they are about the same size as Acanthaster planci. Furthermore, when comparing the D.W ratio throughout all size classes of Archaster typicus or within the other 3 species, there is no evidence for a size-dependent relationship between pattern of movement and body size.

Water currents are used by numerous mobile marine organisms for orientation and to detect food. Sea stars have been variously reported to move upstream (e.g. Castilla & Crisp 1973) or cross-stream, which increases the chance of detecting waterborne chemical stimuli (Rochette et al. 1994). In our experiments, all 4 sea star species did not show a preferential angle of movement in relation to the water current direction. This suggests that the use of a tarpaulin as substrate worked well in avoiding sensory interruptions.

Large sea stars are thought to move faster than smaller specimens (e.g. Westheide & Rieger 1996), based on the assumption that velocity depends on the number (Gaymer & Himmelman 2008) and length of tube feet and thus on body size (Westheide & Rieger 1996). Indeed, Moran (1990) reported that small Acanthaster planci were slower than large specimens. However, Ferlin (1973) found body size and velocity of the sea star Astropecten aranciarius not to be related. In the present study, we found a significant linear relationship between R and both locomotion rate and maximum speed for Archaster typicus, which confirms that larger sea stars move faster than smaller ones. However, although Archaster typicus displayed the highest locomotion rate, it was the smallest species studied. A relationship between body size and velocity was not found for the other 3 species though. Hence, the relationship between body size and velocity of sea stars may just be species-specific. Nevertheless, morphological differences may explain inter-specific variation in locomotion rate. Blake (1989) hypothesized that sea-star growth either stresses strength and sturdiness or flexibility. The relatively small, thickened disk and the moderately enlarged marginal plates of Linckia laeve-gata and Protoreaster nodosus provide sturdiness and protection, whereas the relatively larger and flattened disk as well as the less-developed marginal plates of Acanthaster planci and Archaster typicus feature flexibility, which promotes locomotive activity. This supports our findings that Acanthaster planci and Archaster typicus perform higher locomotion rates than L. laevigata and P. nodosus. Moreover, Acanthaster planci is a multi-armed sea star with a large disk and relatively high number of tube feet and therefore presumably has increased predatory abilities such as locomotion (Blake 1989). Furthermore, tube-foot morphology is likely to affect locomotion rates. Santos et al. (2005) hypothesized that tube-foot morphology is influenced by the adaption to habitats and varying substrate types (Blake 1990) but within limits imposed by the evolutionary lineage (Vickery & McClintock 2000). In accordance to this theory, all 4 species in the present study feature the typical valvaid simple disk-
Acanthaster planci and Linckia laevigata are usually encountered in highly structured, 3-dimensional hard-substrate and soft-bottom habitats; and so the main functions of their tube feet are locomotion and fixation (Santos et al. 2005). In contrast, Archaster typicus and P. nodosus both occur in less-structured soft-bottom habitats and the main function of their tube feet is primarily locomotion. Therefore, it would be likely that the latter 2 species have higher locomotion rates than the 2 species that are adapted to highly structured environments, especially under our experimental conditions on a 2-dimensional tarpaulin. And in fact, L. laevigata showed the lowest locomotion rate of all 4 species tested. However, Acanthaster planci had the second highest locomotion rate. A possible explanation might be that its relatively fast locomotion rate may allow Acanthaster planci to move over sandy areas in search of coral prey and may provide it with the potential to migrate from one reef to another (Moran 1988).

This could also explain the successful spreading over large coral reef areas during outbreaks of Acanthaster planci (e.g. Moran 1986, Babcock et al. 1994, Bos 2010).

Other explanations for variation in inter-specific locomotion rates may be given by differences in prey abundance, prey quality, and feeding behavior. Linckia laevigata is known to be a microphagous feeder grazing on detritus and small invertebrates living in algal mats (Schoppe 2000). For this species, there seems to be no need to move fast, because its prey does not move (or only slowly) and is usually abundant. Protoreaster nodosus on the other hand is an opportunistic microphagous feeder (Schoppe 2000), also feeding on mobile invertebrates such as heart urchins (Bos et al. 2008a). While foraging, P. nodosus often covers longer distances (B. Mueller et al. pers. obs.), which makes it beneficial to move at a considerable speed. Acanthaster planci, a coralivore, may need to migrate to other feeding grounds when food becomes scarce, which may occur during population outbreaks. Moreover, large numbers of A. planci were observed to move from deeper water up the slopes of reefs during early stages of outbreaks (Moran 1988), Moran (1988) suggested that larvae had settled at the base of the reefs about 2 yr earlier and spent their juvenile life stages there feeding on coralline algae. When maturing, feeding behavior changes and sub-adults migrate to shallower reefs to prey on coral polyps.

One further reason to explain inter-specific differences in locomotion rate may be the existence of migration as part of a species’ life cycle. Bos et al. (2008a) found juvenile Protoreaster nodosus (R < 8.0 cm) exclusively in shallow sandy habitats with abundant seagrass in water ≤ 2 m deep, whereas adults (R = 10 cm) were found at a water depth ranging from 0 to 37 m independent of seagrass presence. They suggested that juvenile P. nodosus displays a gradual migration to adult habitats, which would underline the biological need for a certain level of locomotion.

Our findings on the size-dependent locomotion rate of the 4 tested valvate sea-star species suggest that there is a relationship between body size and locomotion rate, irrespective of species affiliation. Specimens of all 4 species moved at the same speed in relation to their body size. Hence, if this is the case, the body size of a specimen may partly overrule species-specific characteristics such as body structure and tube-foot morphology and therefore morphological and behavioral adaptations.

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LITERATURE CITED


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