Mucus-net feeding on organic particles by the vermetid gastropod *Dendropoma maximum* in and below the surf zone

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ABSTRACT: *Dendropoma maximum* is a sessile marine gastropod that inhabits reef flats along a distinct, shallow, sub-tidal zone and uses mucus nets for passive suspension feeding. We tested the hypothesis that surface waves improve food capture by animals inhabiting the surf zone. *D. maximum* feeds mainly on suspended detrital particles that are highly abundant in its habitat. Its feeding strategy depends on ambient currents, not only for the transport of suspended particles to the filtering apparatus but also for the spreading of the mucus net and for determining its shape and size. Using an in situ experiment, we observed a 1.7-fold decrease in prey capture rates among animals transplanted from the surf zone to 5 m depth, where most of the flow was unidirectional. Part of the reduction in feeding rate could be attributed to a lower concentration of detrital particles at the deeper habitat. However, laboratory experiments in a recirculating flume showed that the mucus net produced by the animals under oscillatory flow was larger than under unidirectional flow. *D. maximum* is well-adapted for trapping detrital particles under the unique hydrodynamic conditions in the surf zone.

KEY WORDS: Suspension feeding · Oscillatory flow · Sea waves · Detritus

INTRODUCTION


The vermetid gastropod *Dendropoma maximum* Sowerby, 1825 is a common inhabitant of Indo-Pacific coral reefs. It is most abundant on shallow reef flats, at sites exposed to waves (Hughes & Lewis 1974, Smalley 1984). This sessile animal has an irregularly uncoiled shell, cemented to the substrate (Keen 1961). It feeds by excreting a mucus net that originates from a large pedal gland and spreads into the water via special, grooved tentacles. The net remains in the water for 20 to 40 min, close to the substrate, after which it is pulled...
to the mouth and ingested together with all the trapped particles (Morton 1950, Hughes & Lewis 1974). The animal's diet consists mostly of plankton, meio-benthos and detritus (Kappner et al. 2000). The use of mucus nets for suspension feeding is unique because unlike most passive suspension feeders that have a fixed filtering organ, the mucus net can change size and shape in response to different current conditions and its area can be larger than the animal itself. Hughes & Lewis (1974) observed that individuals held in containers with still water usually fail to produce mucus nets and that the direction in which the net is extended corresponds to the flow direction. Hence, currents appear to affect net spreading. Hughes & Lewis (1974) and Smalley (1984) suggested that the distribution of D. maximum on exposed reef flats is determined by feeding. However, very little is known about the mechanisms of mucus net formation, spreading, and food capturing.

Waves generate flow patterns with cyclic periods of acceleration and deceleration and rapid current reversals. Peak velocities can be much higher than the average current speed and the flow is generally turbulent. At depths greater than half the wave length, the currents are relatively unidirectional with a well-developed benthic boundary layer with slow flow. The turbulent benthic boundary layer of oscillatory flow varies in thickness and fluid velocity during the wave cycle, and as a result, large shear forces and accelerations are augmented on the seabed (Denny 1988, Miller et al. 1992). Apart from the different flow environment, waves may affect both the rate and type of particles encountered by the animal’s trapping mechanism (Muschenheim 1987, Miller & Sternberg 1988). The higher turbulence in the surf zone can markedly increase particle transport from the water column to the bottom but can also resuspend bedload particles (Denny & Shibata 1989). Miller et al. (1992) reported a change in feeding behavior of several suspension feeders in response to increased oscillatory flow. However, our understanding of suspension feeding under wave conditions is rather poor.

This study examines the effects of unidirectional and oscillatory flows on in situ and in vitro suspension feeding by Dendropoma maximum.

**MATERIALS AND METHODS**

**Study site.** Field and laboratory work were conducted at the H. Steinitz Marine Biology Laboratory (MBL), on the west shore of the Gulf of Aqaba (29° 30' 40'' N, 34° 55' 58'' E). Waves propagate towards these shores at a sharp angle, low amplitudes and with a frequency of approximately 0.4 Hz, due to the year-round north wind, low fetch and seabed topography. Oceanographic and meteorological conditions in the Gulf of Aqaba have previously been described by Reiss & Hottinger (1984), Genin et al. (1995b) and Genin & Paldor (1999). Occasional southern winds generate stronger waves due to higher fetch, but such instances are rare. The amplitude of the sea-level tide ranges from 30 to 100 cm, exposing the reef flat during extreme low tides, which occur a few times a year, usually during late summer (Loya 1976). The experimental work described herein focused on the shallow subtidal zone, where the actual depth of the seabed changes with tide. For simplicity, all the depths reported here are relative to the reef flat (i.e. the yearly minimum water level) and not to actual water depth.

**Density distribution.** The distribution of Dendropoma maximum was sampled across the reef flat by randomly placing 20 quadrates (1 × 1 m) along each of five 100 m long transects (A to E in Fig. 1). All transects were parallel to the seaward edge of the reef flat. On the flat, the distance between transects B to D was 7 m. Transects A and E were along the upper 1 m of the vertical wall of the flat’s sea- and lee-ward ends, respectively. Due to the scarcity of D. maximum in other parts of the reef, its incidence in the lagoon and at depths greater than approximately 2 m below the reef flat (G and F, respectively) was surveyed qualitatively.

**Transplantation procedure.** Individuals used in the in situ experiment were collected from horizontal surfaces at a depth of 0.5 to 1 m. The animals were detached from the bottom using a chisel and hammer, and food capturing.

**Fig. 1.** Schematic drawing (not to scale) of a cross-shore transect along the reef flat. Arrows A to E indicate the location of the long-shore transects in relation to the reef flat. A: upper 1 m of the vertical wall facing seawards; B: the edge of the reef flat facing the coming waves; C: on the reef flat 7 m further towards the shore (approximately the center of the reef flat); D: the horizontal edge of the reef flat facing away from the coming waves; E: the vertical wall of the reef flat facing away from the waves. Two additional locations were only scanned for individuals of Dendropoma maximum. F: the deeper part (>2 m) below the reef flat on the vertical wall facing the waves; G: the bottom of the shallow (<1.5 m) lagoon.
with the substrate they grew on (mainly dead corals) to avoid any damage to the shells. The animals were transported to the lab in a large seawater container and returned to sea at a depth of 1 m within an hour of collection. There, the animals were cemented onto large (30 x 70 cm) limestone plates using underwater epoxy (A-778 Splash Zone Compound, Kop-Coat) and left to acclimatize for a period of 1 wk. Daily observations were made to verify that all the transplanted animals were producing mucus nets. Afterwards, the animals were randomly divided into 2 groups: 13 were transplanted to a shallow site, 1.5 m below the reef flat, and 10 to a deep site, 5 m below the flat, (hereafter, referred to as the ‘shallow-transplanted’ and the ‘deep-transplanted’ groups, respectively). The horizontal distance between the sites was 40 m. In addition to the transplanted individuals, 17 intact Dendropoma maximum, original inhabitants of the shallow site, were individually tagged (hereafter, referred to as the ‘non-transplanted’ group). Seven individuals from the non-transplanted group neighbored the shallow-transplanted animals (at 1.5 m depth), while the other 10 were 4 m away, on the reef flat (0 m depth), where D. maximum was highly abundant. The survival and feeding behavior of the non-transplanted group (17 animals in total) was monitored as a control for the effects of the transplanting procedure.

Environmental conditions at the sites. The current velocity was recorded every 0.5 s (2 Hz) for 40 min at each of the shallow and deep sites using an electromagnetic current meter (Model S4, InterOcean) mounted 50 cm above the seabed. Since only 1 current meter was available, the currents were first recorded at the deep site and then at the shallow site. At the same time, 2 submerged pumps (one at each site), located 60 cm above the seabed, delivered water to the beach (~200 l min–1) through 5 cm diameter PVC pipes. Plankton samples were obtained by filtering the pumped water through a 100 µm plankton net for 45 min. The exact pumping rate was measured before each sample collection, to allow calculations of plankton concentrations. The collected plankton was fixed in 4% formalin in seawater and sub-samples, taken with a 2.5 ml Stemple Pipette (Omori & Ikeda 1984), were counted under a dissecting microscope. The number of sub-samples counted was such that at least 400 organisms were counted from each sample to assure reliable representation of all abundant planktonic species. Copepod carcasses (Genin et al. 1995a) were counted separately from live copepods. Detritus particles, including fragments of macroalgae, animal parts and unidentified organic particles were pooled into a single category, hereafter referred to as ‘fragments’. Chlorophyll a (chl a) concentrations were measured at each site by taking a water sample (280 ml) 40 cm above the seabed, filtering the water on a GF/F filter (Whatman) and extracting the chl a for 24 h in 90% acetone at 4°C as in Strickland & Parsons (1972). The extracted chl a concentration was measured using a TD 700 fluorometer (Turner device).

The entire sampling procedure was repeated each sampling day (n = 41 d). A sampling day started with a dive to the deep site where a 280 ml water sample was collected for chl a analysis, and the current meter deployed. Two divers measured the area of mucus nets, and proceeded to collect them as described below. This work lasted approximately 40 min, after which the divers repeated the procedure at the shallow site. The plankton pumps were set to work simultaneously with the dive to each site. The average current speed, zooplankton density and chl a concentration for each site in each sampling day were used for inter-site comparison of currents and food.

Analysis of mucus nets. We sampled newly-secreted nets precisely 20 min after the initiation of their excretion. First, the existing (old) mucus net was removed. This stimulated Dendropoma maximum to start producing a new net within less than a minute. Next, a 0.5 mm metal wire was laid on the outside perimeter of the animal’s shell aperture, forming a ring (3 cm diameter) through which the mucus net was excreted unobstructed. Twenty min later, the mucus was collected by raising the ring and turning it 3 to 5 times to roll and bundle the mucus net around it. Prior to net collection, the area of the mucus net was estimated by placing a 20 x 20 cm metal grid a few centimeters above the net and counting the number of 1 cm² squares that overlapped with the net. When sampled at night, mucus net area was not measured, and the collection of mucus nets was performed using a dimmed flashlight covered with a red filter. Collecting the mucus after 20 min ensured that the mucus net would be sampled prior to the start of its retrieval by the animal. Preliminary observations (G. Ribak et al. unpubl. data, n = 102) showed that, during the day, D. maximum retrieves its net >20 min after initiating its spreading in >90% of the cases, (see also Kappner et al. 2000).

The collected nets were transferred to the laboratory and kept frozen (~70°C) for later analysis. Food particles were dislodged from the net by disintegrating the mucus in 0.35% sodiumhypochlorite solution (NaOCl) in water (A. Hudson pers. comm.) and filtering the solution through a 100 µm plankton net. We tested the adequacy of the technique, and the ideal concentration of the NaOCl solution by running preliminary trials using a known number (5 to 6) of copepods at a gradient of concentrations of the solution (prepared from a stock solution of 10% active chlorine, NaOCl, Frutarom). All copepods were retrieved up to a concentration of 0.5%. At concentrations >0.5%, the external...
integument of the copepods showed signs of digestion. At concentrations <0.35%, the separation of the particles from the mucus was incomplete. No visible change in the size of fragments was detected as the result of treatment with the 0.35% NaOCl; however, some pigmented fragments showed a tendency to bleach. Food particles were thereby counted separately for organisms (plankton and meiofauna) and fragments. Zooplankton was counted under a dissecting microscope as described above. To estimate the biomass of fragments, we recorded their size using a CCD video camera, mounted on a dissecting microscope and connected to a SONY EVO-9800P VCR. The area of each fragment was measured using a Scion LG-3 frame-grabber and image-analysis software (ScionImage, Scion).

**Flow tank experiments.** The effect of flow regime (unidirectional and waves) and phytoplankton concentration on the feeding rate of *Dendropoma maximum* was examined in the recirculating flume described by Trager & Genin (1993). Flow was generated by a propeller and a small electric motor connected to a desk computer, allowing full control of flow speed and direction. Waves were generated by changing the direction of the flow at a frequency of 0.4 Hz (similar to the average observed in situ). The waves generated were asymmetrical, with a weak residual forward flow (Fig. 2). The flow in the flume, at the height of the animal’s aperture (3 cm above bottom) was measured, without the animals being present, using video tracking of suspended particles. The flow was adjusted so that the RMS (Root Mean Square) speed under the oscillatory flow would equal that of the unidirectional flow. Two RMS speeds were used: 3.5 and 5.5 cm s\(^{-1}\), with the average particle displacement in the corresponding oscillatory motions equal to 1.5 and 1.3 cm s\(^{-1}\), respectively. Each experimental trial used a single *D. maximum*, cemented to a thin smooth plate (20 × 30 cm), positioned at the center of the flume’s working section (20 × 30 × 80 cm) with the animal’s aperture oriented upcurrent. The experimental animals were kept at sea between trials and were brought to the flume after their mucus net was removed. Trials began as soon as the animal started to secrete the new mucus net in the flume (typically <1 min after transfer), and lasted precisely 20 min. At the end of the trial, the mucus net was collected using a pre-weighted wire ring, as described above. Each animal (n = 5) was tested 5 times at each combination of current regime (unidirectional, oscillatory) and speed (3.5, 5.5 cm s\(^{-1}\)). The choice of flow speeds used in the flume experiment was not random. The lower flow speed (3.5 cm s\(^{-1}\)) we used was typical for the deep site (an average current speed of 3.2 cm s\(^{-1}\) was measured at our study site 8 cm above the sea bed, Genin et al. 2002). The faster flow speed (5.5 cm s\(^{-1}\)) was taken as an upper value that deep suspension feeders may rarely encounter at strong currents.

Prior to each trial, the water in the flow tank was filtered through a 100 µm plankton mesh to remove larger particles and fragments (but retaining phytoplankton). A water sample (280 ml) was then taken to determine the concentration of chl \(a\) in the flume. The water sample was filtered on a GF/F (Whatman) filter and the filter incubated in 10 ml of 90% acetone for chl \(a\) extraction as mentioned above. After each trial, the chl \(a\) content of the mucus net was similarly determined by incubating the ring with the attached mucus in 20 ml of 90% acetone for 24 h to extract the chl \(a\). The mucus itself has no auto-fluorescence that could bias chl \(a\) measurement (Kappner et al. 2000). After extraction, the ring with the remaining attached mucus was dried at 60°C for 24 h, and weighed. Weight loss due to acetone extraction and ring handling was evaluated by weighing dried nets, extracting the chl \(a\) in acetone and then drying and re-weighing. The average loss was 31.3% (SD = 11.4, n = 17). To compare chl \(a\) capture at the different flow regimes, the volume of water cleared of chl \(a\) per unit of time (clearance) was calculated as: clearance = (g chl \(a\) accumulated on the net in 20 min)/(g chl \(a\) 1\(^{-1}\) in the flume).

Clearance per weight of net was calculated by dividing the clearance by the weight of the dried net after chlorophyll extraction.
**Estimating total carbon per net.** To estimate the relative contribution of phytoplankton, zooplankton and fragments, we converted all values to organic carbon based on a chl a:C ratio of 1:60 reported from our study site by Yahel et al. (1998). The contribution of phytoplankton could not be calculated *in situ*, as both phytoplankton and some fragments included chl a. The carbon gain through phytoplankton feeding was, therefore, estimated by combining the observed clearance rate at 5.5 cm s⁻¹ in the flume experiment with the ambient *in situ* concentration of phytoplankton in fragment-free samples at the sites.

Biomass from the zooplankton counts was achieved by sorting animals into 3 size categories according to body length (<300 µm; 300 < length < 700 µm; >700 µm). The middle value of each size class (200, 500 and 800 µm, respectively) was then converted to organic carbon weight according to Rodriguez & Mullin (1986) and multiplied by the number of organisms within the category. This conversion is likely to overestimate the organic carbon weight as the equation is based on zooplankton samples that included mainly crustaceans, while our zooplankton included mainly foramenifera and ostracods that have lower carbon content per body size than copepods.

The carbon contribution via fragments was estimated based on our direct size measurements of the fragment trapped on the mucus nets *in situ*. The total area of fragments (mm²) from each net was converted to ash free dry weight (AFDW, mg) based on the following calibration curve:

\[
\log(\text{AFDW}) = 0.6129 \log(\text{area}) - 0.9682
\]

This relationship \( r^2 = 0.9722, n = 7 \) was obtained by measuring the areas and AFDW of the fragments in 7 different aliquots drawn from a single pump sample. AFDW was obtained after filtering the measured fragments samples through pre-weighed, pre-combusted GF/A filters (Whatman) and combusting the filters in 450°C for 4 h, as described by Omori & Ikeda (1984). Carbon was assumed to contribute 40% of the AFDW of organic matter as reported (for plankton) by these authors.

**Sample size.** Currents and plankton concentrations were measured at the shallow and deep sites in October 1998 (17 days and 4 nights) and in April to May 1999 (16 days and 4 nights). Mucus nets were measured for area and collected, at each site, simultaneously with current measurements and plankton collection, as described above. Since the net area was not measured during the night, we obtained a total of 41 samples of environmental conditions and mucus nets, only 33 of which had measurements of the net area. Due to the tedious quantification of fragment size, the food content of the mucus nets was only analyzed from the nets of 12 randomly-selected sampling dates (8 days, 4 nights) in October 1998 and 13 sampling dates (9 days, 4 nights) in April to May 1999. Within each sampling day, we only analyzed the nets of 6 animals from the deep-transplanted group and 6 from the shallow-transplanted group. In total, we analyzed the content of 300 mucus nets from 25 sampling days. The nets from all other *Dendropoma maximum* in the shallow-transplanted, deep-transplanted and non-transplanted groups were measured for area and collected each sampling day without further processing in order to treat all the transplanted and non-transplanted animals equally.

**Statistical analysis.** All statistical tests were performed using STATISTICA (StatSoft). Due to logistical limitations, our samples were replicated in time, rather than sites. Our statistical analysis is, therefore, based on repetitive (daily) monitoring of the flow and plankton conditions at the same deep and shallow sites. Similarly, mucus nets were sampled on different days from the same individuals at the 2 sites. We, therefore, used repetitive measures design (StatSoft 1995). Inter-site differences in specific environmental conditions (currents, chl a) were tested using a paired t-test. The inter-site difference in plankton concentration was tested using Repeated Measures ANOVA (RMANOVA). The average mucus net area of the 4 treatment groups described above was compared by a 1-way ANOVA for repeated measures. The average biomass of the fragments (total AFDW net⁻¹) on the deep and shallow mucus nets was compared using paired t-test, where the different sampling days served as repetitions in all cases. ANOVA assumptions (homo- geneity of variance and the normal distribution of residuals) were tested and data transformation was performed when necessary (Underwood 1981)

**RESULTS**

**Density distribution.**

The density of *Dendropoma maximum* along the reef-flat increased towards the seaward edge (Fig. 3). High density was found on the upper 1 m of the seaward vertical wall. *D. maximum* was absent deeper than 3 m below the reef flat and was rare in the shallow lagoon. The few individuals found in the lagoon were living on rocks likely dislodged from the near by reef flat.

**Environmental conditions at the sites.**

Daily variations in the concentrations of phytoplankton and most zooplankton groups at the deep
and shallow sites were highly correlated (Table 1). The currents, however, were very different at the 2 sites, with oscillatory currents dominating the shallow site (Fig. 4), producing significantly faster currents \((p < 0.01; \text{paired } t\text{-test})\) than the typical unidirectional flow at the deep site. No significant correlation was found in the current speed at the 2 sites \((r = 0.29, p > 0.067, n = 41, \text{Table 1})\). Zooplankton concentrations were significantly higher at the shallow site \((p < 0.001, \text{RMANOVA})\), mostly due to enhanced concentrations of carcasses, fragments, ostracods and eggs \((p < 0.03 \text{ for all 4 categories, Tukey post hoc test})\). All other planktonic groups were similar at both sites. The average numerical abundance of fragments and carcasses at the shallow site was about twice that at the deep site.

### Analysis of mucus nets

The size of mucus-net area in *Dendropoma maximum* in its natural habitat (non-transplanted group) on the reef flat ranged from 7 to 20 cm², with an average of 12.8 cm² \((SD = 2.5)\) This area was significantly larger than that of the deep-transplanted and shallow-transplanted groups, and the non-transplanted individuals below the flat \((p < 0.001, \text{1-way RMANOVA, Table 2})\). Net size did not differ between shallow-transplanted and deep-transplanted animals \((p > 0.28, \text{Tukey post hoc test})\), nor were the net sizes of the 2 groups correlated in different days \((r = 0.13, p > 0.4, n = 33)\). At the shallow site (below the reef flat), the net size of transplanted and non-transplanted groups did not differ significantly \((p > 0.16, \text{Tukey post hoc})\), exhibiting a significant correlation in time \((r = 0.39, p < 0.03, n = 33)\). The area of the net was significantly correlated with current speed at the deep site \((r = 0.5, p < 0.01, n = 33)\), but not at the shallow site \((r = -0.1, p > 0.42, n = 33)\).

The mucus net of *Dendropoma maximum* contained a few planktonic organisms (Fig. 5), with no apparent difference in the taxonomic composition at the 2 depths \((p > 0.292, \text{RMANOVA})\) and no significant difference between day and night \((p > 0.147, \text{RMANOVA})\). In terms of carbon, the average catch per net was 11.4 \((SD = 12.2)\) and 11.5 \((SD = 16.9)\) \(\mu g \text{ C net}^{-1}\) for the shallow and deep animals, respectively. Most of the diet, however, consisted of fragments. The average AFDW of the fragments trapped on the nets at the shallow site was 1.7 times higher than that at the deep site \(0.369 \text{ (SD = 0.296) versus 0.213 (SD = 0.125) mg AFDW net}^{-1}\), respectively \((p < 0.03, \text{paired } t\text{-test, Fig. 6})\).
A significant increase in clearance rate by a factor of 2.42 (SD = 0.33) (p < 0.006, RMANOVA) occurred when the unidirectional current of 3.5 cm s\(^{-1}\) was changed to oscillatory flow with similar RMS speed. The enhanced clearance rate was probably due to a 3-fold (SD = 1.4) increase in the weight of the mucus net (p < 0.013, RMANOVA), as clearance rates per net weight did not differ among the 2 flow regimes (Fig. 7). No such difference in clearance rate was found when the flow speed was 5.5 cm \(s^{-1}\) (Table 3). An increase in flow speed from 3.5 to 5.5 cm \(s^{-1}\) caused an increase in both net weight and clearance rate (p < 0.012, RMANOVA).

The rate of phytoplankton clearance was correlated with net weight (\(r = 0.783, p < 0.001, n = 100\)). Surprisingly, no significant correlation was found between the amount of chl \(a\) caught per g net and the chl \(a\) flux (speed \(\times\) concentration) (\(r = 0.097, p > 0.335, n = 100\)).

**Total carbon per net**

In terms of carbon, the feeding rate per animal per single net cycle was as follows: phytoplankton 17.3 and 15.8 \(\mu\)g C net\(^{-1}\) cycle\(^{-1}\) for the deep- and shallow-transplanted animals, respectively. The carbon content of the \textit{in situ} trapped fragments (40\% of AFDW) was 85 and 148 \(\mu\)g C net\(^{-1}\), while that of zooplankton was 11.5 and 11.4 \(\mu\)g C net\(^{-1}\), at the deep and shallow sites, respectively. Thus, on average at the shallow site, fragments constituted 84.5\% of the carbon ingested by \textit{Dendropoma maximum}, phytoplankton 9.0\% and zooplankton 6.5\%, while at the deep site, the proportions were 74.7, 15.2 and 10.1\%, respectively. The total carbon gained by a \textit{D. maximum} in a single round of mucus net was 0.114 mg C net\(^{-1}\) 20 min\(^{-1}\) in the deep site and 0.175 mg C net\(^{-1}\) 20 min\(^{-1}\) in the shallow site.

**DISCUSSION**

\textit{Dendropoma maximum} living at shallow depths (1.5 m below the reef flat) obtained more food per mucus net than individuals living at greater depths. Our \textit{in situ} and flume experiments indicated that this inter-depth difference was mostly due to the higher food flux at the shallower site and depth-related changes in the flow regime. This may suggest that the density distribution of \textit{D. maximum} (present study, Hughes & Lewis 1974) is related to feed-

**Table 2. \textit{Dendropoma maximum}. The average area (±SD) of the mucus nets in the different groups, over 33 sampling days. Superscripted letters (a, b) indicate statistical similarity between groups (p < 0.05, Tukey post hoc test). TD: Deep-transplanted group; TS: shallow-transplanted group**

<table>
<thead>
<tr>
<th>Group</th>
<th>No. of animals</th>
<th>Area (cm(^2)) ± (SD)</th>
<th>Range (cm(^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>TD(^a)</td>
<td>6</td>
<td>9.2 (3.26)</td>
<td>5.5–14.47</td>
</tr>
<tr>
<td>TS(^b)</td>
<td>8</td>
<td>9.51 (2.23)</td>
<td>5.88–16</td>
</tr>
<tr>
<td>Non-transplanted (1.5 m)(^a)</td>
<td>6</td>
<td>10.62 (2.71)</td>
<td>5.57–16.67</td>
</tr>
<tr>
<td>Non-transplanted (reef-flat)(^b)</td>
<td>10</td>
<td>12.77 (2.48)</td>
<td>6.67–19.93</td>
</tr>
</tbody>
</table>
The maximum density was observed on the reef table facing the waves and individuals were rarely found below 3 m or in the lagoon. Our measurements show that, at the shallow site, *D. maximum* encountered faster currents, higher fragment concentrations and had larger mucus nets. That individuals transplanted to a 5 m depth had a higher mortality rate than those transplanted to a shallow site (37 versus 7%, respectively; G. Ribek unpubl. data), suggests that the absence of *D. maximum* at deeper depths could be related to lower feeding rates. However, the precise cause for mortality and the lack of *D. maximum* at deeper depths awaits further investigation.

*Dendropoma maximum* is well-adapted to feeding in the physical environment of the surf-zone. Both the field measurements and the flume experiment indicated that net size was larger when flow was stronger or when a weak unidirectional current became oscillatory. The flume experiment, where changes in the weight of the excreted net were induced by changes in the flow regime, indicated that net size might be actively determined by the animal. An alternative, however not mutually exclusive, feeding strategy may be an enlargement of the net area due to the enhanced drag provided by waves and stronger currents, causing further spread of the net area. Regardless of the reason, the outcome was an increased net and greater particle entrapment by the animal.

Although located below the zone of maximum population density, our shallow site was within the natural depth range of *Dendropoma maximum* as indicated by the presence of non-transplanted animals. Our shallow site was a logistical compromise since we could not secure the large plates on which *D. maximum* was transplanted to the reef flat in a manner that would prevent their dislodgment during the occasional winter storms. We chose to cement the animals to plates, rather than directly to the natural bottom, to eliminate the variance of substrate structure. Above our shallow site, at the point of maximum population density (the reef flat), animals may experience even greater feeding rates due to exposure to stronger currents and waves. Indeed, the area of mucus nets measured from non-transplanted individuals, naturally found on the reef flat, was the highest of all groups.

A mere difference of 3.5 m in depth caused a great difference in environmental conditions. Waves dominated *Dendropoma maximum*’s natural habitat, while unidirectional flow (tidal currents) dominated the deeper zone. While the concentration of living plankton was similar at the shallow and deep sites, detrital particles (fragments and carcasses) were more abundant (~2-fold) at the shallow site. Previous studies of *D. maximum* diet also observed detritus on mucus nets (Kappner et al. 2000), but did not provide a quantitative estimate for the contribution of those particles to the animal’s diet. Our measurement of the carbon content in the trapped fragments indicates that this type of food contributes most of the carbon ingested by individuals.
The higher abundance of fragments at the shallow site was due either to local generation or accumulation. The action of the waves on the seabed may erode the turf on the shallow reef, releasing suspended fragments. Additionally, waves greatly enhance resuspension of bedload particles by augmenting acceleration, shear and turbulence (Denny & Shibata 1989). Since detrital fragments are negatively buoyant (G. Ribak pers. obs.), enhanced resuspension by waves should increase their availability for being trapped by the encrusting net of *D. maximum*. An elevated availability of such fragments 60 cm above the bottom was observed at the shallow site using our pumps.

Phytoplankton only contributes a little amount of carbon (9 to 15%) to the diet of *Dendropoma maximum*. The ca. 10% higher concentration of phytoplankton at the deeper site, therefore, had no significant effect on the animal’s *in situ* diet.

Our estimate of the total carbon gained by *Dendropoma maximum* per net cycle at the shallow site is an order of magnitude lower than that of Kappner et al. (2000), obtained at a nearby site. This discrepancy can be partially attributed to the differences in methodology. Kappner et al. (2000) measured total carbon content of mucus nets collected on the reef flat and then subtracted a fixed value of the carbon content of ‘empty’ nets (obtained from animals in a container filled with filtered seawater). Our results show that the weight of the mucus net is dependent on current speed and can change almost an order of magnitude in size when flow speed changes from 3.5 to 5.5 cm s⁻¹. Thus, the weight of empty nets obtained in a container with little or no water motion could have substantially underestimated the carbon content of the net excreted by the animal *in situ* under substantial water motion, thereby overestimating the carbon content of the food trapped by the net. Another difference between the method we use and that of Kappner et al. (2000), is that we always used fresh nets that were deployed for exactly 20 min of net exposure to the flow (see text).
(Table 2). However, the suggestion by Kappner et al. (2000) that the daily ingestion of *D. maximum* amounts to 41% of its weight, should be taken with caution. This ration is greater than that of flying endothermic animals (e.g. 24 to 10% for piscivorous birds, Nilsson & Nilsson 1976). The order-of-magnitude lower values of food ingested per net reported in the present study, suggest a much lower ration and appear to be more realistic for this type of ectothermic, sessile mollusc.

Our current measurements were made as close to the bottom as was possible (50 cm). The current speed at this depth is expected to be stronger than at <1 cm above the bottom. Genin et al. (2002) measured the flow at 8 cm above the bottom at a depth of 8.5 m and found that the current speed was <3 cm s⁻¹ 67% of the time. Hence, the absence of oscillatory motions greatly reduced the spreading of the mucus net, as indicated by our flume experiments at 3.5 cm s⁻¹.

The mucus net of *Dendropoma maximum* is well-adapted for trapping near-bed particles in the wave zone. The net is flexible and strongly adheres to the substrate, allowing a collecting area remarkably larger than the animal’s aperture. The flexibility of the mucus net may reduce the impact of the waves (Koehl et al. 1991) and allows its large size. That the animal can modulate the size of the net as a function of the flow, allows further optimization of its usage.

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