Significance of copepod emergence to benthic, pelagic, and phytal linkages in a subtidal seagrass bed

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ABSTRACT: The role of meiobenthic copepod emergence in linkages among benthic, pelagic, and phytal habitats was examined in a subtidal seagrass (Thalassia testudinum) bed at the mouth of Tampa Bay, Florida, USA. On 4 dates the ability of emerging copepods to affect the density and composition of benthic and phytal assemblages was tested during 2 h periods of minimum and maximum emergence. Copepod exchanges between benthic and phytal habitats and the duration of pelagic excursions also were examined throughout a night by measuring settlement into 2 types of sediment traps and onto natural and defaunated seagrass blades. The ability of emerging fauna to disperse through the water column was determined from horizontal collectors. Changes in copepod densities and composition during periods of active emergence did not indicate an unequivocal exchange between sediment and seagrass blade assemblages. Copepod resettlement onto the sediments was greater during periods of increased emergence and suggested that emerging copepods typically did not remain in the water column for a prolonged (> 2 h) time. The dispersal of copepods through the water column was haphazard and unaffected by prevailing currents. Species and habitat affiliation influenced both the dispersal and settlement behaviors of emerging copepods. Although appreciable numbers of sediment-associated copepods can enter and disperse through the water column, our results suggest that the effects of emergence on linkages between benthic, pelagic, and phytal habitats are minimal or limited in duration.

KEY WORDS: Benthic-pelagic-phytal coupling · Emergence · Meiobenthic copepods · Seagrass · Resettlement

INTRODUCTION

Adult and postlarval stages of sediment-dwelling invertebrates from diverse taxonomic groups periodically enter the water column in a variety of habitats (Walters 1988, Jacoby & Greenwood 1989, Lewis & Boers 1991), and this emergence may affect significantly both benthic and pelagic environments. Frequent and substantial emigrations of individuals from the sediments would influence density dependent processes that occur within benthic populations (see Service & Bell 1987, Walters 1991). While in the water column, emerging metazoans might interact with organic aggregates (Shanks & Edmondson 1990), phytoplankton (Decho 1986), zooplankton, and nekton (Robertson & Howard 1978). The extended distances that emigrating adults could disperse would affect the spatial scales over which benthic population phenomena occur (sensu Wiens 1989). Current models of benthic community development that focus exclusively on larval recruitment (i.e. Underwood & Fairweather 1991) also may need to be expanded to account for the effects of adult and postlarval recruitment.

Harpacticoid copepods numerically dominate emerging fauna in most habitats (Cahoon & Tronzo 1988, Walters 1988, Armonies 1989), and a significant proportion of sediment-dwelling copepod assemblages frequently enter the water column (Walters & Bell 1986, Walters 1991). Copepod emergence is influenced by ambient light levels (Armonies 1988a), tidal (Bell et al. 1988) or diurnal periods (Walters 1988, 1991), current velocities (Armonies 1988b), copepod densities (Service & Bell 1987, Walters 1991), and species behaviors (Walters 1991). Possible reasons for harpacticoid...
copepods emerging from the sediments include mating (Bell et al. 1988), foraging (Decho 1986), or dispersal (Kurdziel & Bell 1992).

The impact of emergence on benthic and pelagic environments will depend partly on the time spent in the water column, the distances dispersed, and the final destination for emerging copepods. Emergence effects would be limited if copepods entering the water column immediately resettled to the same position in the sediments. Historically, researchers have targeted the processes governing either movement into or settlement out of the water column, but not both (e.g. Armonies 1988a, Fegley 1988, Kern 1990, Walters 1991). Separate studies of copepod advection (Palmer & Gust 1985) and the colonization of defaunated sediments in hydrodynamically active habitats (Sherman & Coull 1980, Chandler & Fleeger 1983, Kern & Taghon 1986, Fegley 1988) suggest a direct connection between pelagic dispersal and benthic recruitment. In unvegetated habitats a few studies have documented the simultaneous emergence and resettlement of copepods (Alldredge & King 1980, Cahoon & Tronzo 1988, Jacoby & Greenwood 1988, 1989). Unfortunately, interpretations of results are hampered by 2 problems: (1) potentially biased settlement trap designs (e.g. Gardner 1980a, Butman 1986) that prevent an accurate assessment of the origin (emergence or advection) and rate of copepod settlement, and (2) trap deployments typically >10 h that limit conclusions about the length of time emerging copepods spend in the water column.

The resettlement of emerging copepods from a subtidal seagrass bed was examined in field mesocosms by Bell et al. (1989), but those results may not reflect normal behavior because the availability of natural substrates for settlement was limited (<30% of the total surface area) and mesocosm isolation might have affected copepod behavior. Because of the restricted focus and technical limitations of past studies, little is known about the fate of emerging copepods after leaving the sediments in the field.

In vegetated habitats, understanding the connections among emergence, dispersal, and settlement is complicated by the presence of phytal structures. Above-ground phytal structures can alter patterns of near-bottom flow and affect settlement (Eckman 1983). Phytal habitats represent a potential destination for sediment-emerging copepods. Macrophytes also are populated by unique copepod species (Rutledge & Fleeger 1994) and species that are capable of entering the water column (Hicks 1986, Palmer 1986, Kurdziel & Bell 1992). Phytal copepods in seagrass habitats can disperse through the water column and colonize blades >20 m from a source bed (Bell & Hicks 1991, Kurdziel & Bell 1992). Although copepod morphology can identify potential habitat preferences and behavior in seagrass beds, many species that reside on blades also emerge from the sediment (Bell et al. 1987). In mesocosms sediment-emerging copepods resettle onto seagrass blades (Bell et al. 1989), and Webb & Parsons (1992) suggest benthic copepods readily colonize blades. The presence of phytal structures as an additional source of and sink for emerging copepods increases the complexity of possible linkages between benthic and pelagic environments.

In the present study we examine how emergence contributes to linkages among benthic, pelagic, and phytal habitats. Copepod densities and species composition in the sediments and on seagrass blades were determined during periods of minimum and maximum emergence to investigate exchanges between benthic and phytal habitats. Colonization of sediments and blades was examined during frequent nighttime intervals to compare simultaneously patterns of emergence with benthic and phytal resettlement and assess copepod persistence in the water column. Connections between copepod emergence and pelagic dispersal also were investigated. Our goal was to determine the fate of emerging meio-benthic copepods in seagrass habitats and evaluate whether emerging copepods represent a significant link among benthic, pelagic, and phytal habitats.

**METHODS**

**Study site.** All samples were collected from a subtidal seagrass bed adjacent to the eastern end of Mullet Key at the mouth of Tampa Bay, Florida, USA (Fig. 1). The study site was located within a gently sloping embayment predominated by extensive *Thalassia testudinum* König beds beginning ca 20 m offshore. The site is not exposed to wave action or intense tidal currents. In Mullet Key Channel (Fig. 1), the mean diurnal tidal range is 0.7 m and maximum current speeds are <50 cm s⁻¹. Current velocities within the vegetated study site located behind Mullet Key at least 1.5 km away from a main tidal channel (Fig. 1) were not measured directly in this study, but maximum velocities in similar seagrass beds average <5 to 6 cm s⁻¹ (Halls pers. comm.). On average, sediments were fine sands with <7.0% silt/clay and <4.0% total organic content (Walters 1987).

**Field collections.** Copepod movements among benthic and phytal habitats were examined initially by comparing the densities and species compositions of emerging, benthic, and phytal assemblages in field collections. A reduction in densities for individual copepod species in the sediment coincident with increased numbers emerging and on seagrass blades would suggest movements between benthic and phy-
tal habitats. The distinct periodicity of emergence in subtidal seagrass beds (Walters 1988, 1991) enabled us to compare benthic and phytal densities and compositions during times when copepods were known to enter the water column.

Samples of emerging copepods and seagrass blades with associated fauna were collected from presunset and postsunset periods, times of minimum and maximum movement out of the sediments (Walters 1988, 1991), on 19 March, 20 May, 12 August, and 30 September 1983. Except for September, sediment samples also were taken and all sampling occurred within 2 h of high tides (0.5 to 0.6 m above MLW). Emergence samples were collected using traps described by Walters (1988) that consisted of an 82 cm² PVC base, a clear acrylic catch-tube with an attached inverted funnel, and a 63 μm mesh cap (Fig. 2). At the beginning of a sampling period, traps were positioned in the field between seagrass culms to exclude Thalassia testudinum blades and only sampled emerging, sediment-dwelling fauna. Organisms migrating 15 cm from the sediment surface and trapped in the catch-tube were collected during 2 h periods. Some emergence data have been presented earlier (Walters 1991), but new analyses are included to allow a direct comparison with seagrass blade and sediment data. Resident fauna on T. testudinum were sampled in a manner similar to that of Hall & Bell (1988) by placing a clear plastic tube filled with filtered seawater over haphazardly chosen individual blades that were then excised at the sediment surface. Sediment samples were taken from an undisturbed area adjacent to emergence traps using a 6.2 cm² corer to a depth of 3.5 cm. A total of 5 emergence, 8 to 10 blade, and 8 to 10 sediment samples were collected at the end of each sampling period. Data were standardized to numbers per 10 cm² sediment or seagrass blade surface. Emergence, blade, and sediment data were used to test whether copepod emergence coincided with increased numbers on seagrass blades and decreased numbers in the sediments.

Field experiments. Copepod emergence, horizontal transport, and settlement onto sediments or seagrass blades were examined on 20–21 April 1987. During the study tides ranged from 0.6 m MLW (ca 75 cm of water at the study site) at 17:11 h before the start of
sampling to 0.1 m MLW (ca 25 cm of water at the study site) at 02:26 h. Emerging copepods were collected in traps \( n = 5 \) only during 2 h presunset and postsunset periods as described previously.

Organisms transported in the water column were sampled by horizontal traps positioned above both sediments and seagrass blades (Fig. 2). Each trap consisted of an 82 cm\(^2\) PVC collar horizontally attached to a stake placed within the seagrass bed into which a 100 mm polypropylene funnel and attached collection jar were fitted. To enable water but not organisms to flow through the trap, the bottom of the collection jar was removed and fitted with 63 \( \mu \)m mesh. Two arrays of 4 traps facing north, east, south, and west were placed in the field for 2 h periods. Prevailing water movement in the seagrass bed on outgoing tides when samples were collected is to the east-southeast (Fig. 1). Collection of greater copepod numbers was expected in traps facing the prevailing currents (N and W) if flow affected the water column dispersal of emerging copepods. Traps were positioned above the extended tips of seagrass blades, ca 30 cm above the sediment surface, to prevent the accidental collection of organisms directly from blades that might have contacted trap surfaces. Horizontal traps were deployed only during 2 periods covering sunset (20:00 to 22:00 h) and just after sunset (22:00 to 24:00 h). After 24:00 h, water levels prevented the placement of traps above seagrass blades. To determine if sediment-associated copepods were dispersed in the water column, data on the presence/absence of individual species in emergence and horizontal traps were compared.

Two cylinder traps with different collectors, petri dish and funnel, were employed to measure copepod settlement to the sediment surface (Fig. 2). Petri traps were similar to many open tray resettlement samplers that allow unrestricted faunal ingress and egress (e.g. Alldredge & King 1980). Funnel traps were intended to restrict the reemergence of settled copepods but were not similar to settlement traps which have been shown to have a significant sampling bias (Gardner 1980a, Butman et al. 1986). Both traps consisted of a 10.7 cm diameter PVC base that was pushed into the sediments until ca 10 cm remained above the surface and into which petri or funnel collectors were placed. Positioning traps above the sediment surface prevented the inadvertent collection of nonemerging or seagrass blade copepods. In the petri traps, a 60.7 cm\(^2\) covered petri dish filled with heat-treated (60 °C for > 24 h) seagrass bed sediments and filtered seawater was placed on a wire mesh shelf 4.5 cm above the sediment surface inside the PVC base (Fig. 2). Heating the sediments killed resident copepods and rendered remaining exoskeletons easily identifiable. Funnel traps consisted of a 100 mm polypropylene funnel and collection jar similar to horizontal traps. The top of the funnel was placed 4.5 cm below the top of the PVC base and the collection jar filled with a small amount of heat-treated sediment (Fig. 2). Settlement samples \( n = 5 \) were collected and traps replaced every 2 h throughout the night beginning before sunset and ending after sunrise. Traps provided data on copepod species recruitment to the sediment surface that could be compared directly to data on benthic copepod emergence.
Settlement trap design and placement represented a compromise between considerations of the hydrodynamic factors known to affect trapping efficiency, the physical characteristics of the seagrass environment, and the hypotheses being addressed. A suite of physical parameters including trap Reynolds number, particle fall velocities, and trap aspect ratio all can affect collection efficiencies (Gardner 1980a, b, Butman et al. 1986, Baker et al. 1988). The smaller the aspect ratio and greater the current velocity the greater the chance that particles will be resuspended and bias trap results (Gardner 1980b, Butman 1986). In areas of slack current flow such as the conditions under which our study was conducted, trap dimensions appear to be less critical to collection of an unbiased resettlement sample. Differences in trap aspect ratios (Gardner 1980a) or trap diameter for a fixed aspect ratio (Baker et al. 1988) had a limited effect on trapping efficiency at low (<9.5 cm s⁻¹) current velocities. Although the aspect ratio of both traps was <1, maximum current velocities in seagrass beds typically are <5 to 6 cm s⁻¹ and resuspension was not considered a major source of sample bias. Horizontal and vertical positioning and resultant secondary circulation patterns also can bias settlement trap results (Gardner 1980b, 1985). Our placement of traps 10 cm above the sediment surface was necessary to limit the collection of non-emerging or seagrass blade copepods. In low flow conditions, secondary circulation patterns likely would bias our results less than contamination of settlement traps with copepods that move or are transported along the sediment surface but do not emerge. Although physical factors affect the settlement of living organisms (Butman 1989, Yund et al. 1992), swimming and other behaviors also can influence trapping efficiencies (Fegley 1988, Coale 1990). A comparison between the 2 different settlement traps, one that restricted egress and one that did not, enabled us to examine the possible effects repeated emergence might have on copepod settlement patterns.

The movement of copepods onto and off Thalassia testudinum was examined using defaunated and natural blades. To defaunate blades, natural blades were collected from the field, treated with an MgCl₂ solution (72 g l⁻¹) for 15 to 20 min to remove motile epifauna, and stored in 63 μm filtered seawater for later use. Post-treatment examination indicated that copepods were removed effectively from defaunated blades. At the beginning of each sampling period defaunated blades were attached to 10 cm² plastic mesh squares haphazardly placed within the seagrass bed and anchored into the sediments. Throughout the night 5 defaunated and natural blades were collected from the field after each 2 h sampling period using blade collection methods described above. Copepod numbers on seagrass blades were standardized per 10 cm² of blade surface area and, along with species data, provided information on the movement of phytal copepods and the colonization of blades by emerging benthic and phytal copepods.

All samples were preserved in the field using a 10% buffered formaldehyde and rose bengal solution. In the lab, seagrass blades were agitated vigorously and visually inspected to remove all attached fauna. Sediment samples were processed using a shake and decant procedure (Wieser 1960) which removed greater than 95% of the copepods (Walters unpubl.). All copepods were enumerated at 25x and identified to the lowest taxon possible.

**Statistical analyses.** Copepod densities were analyzed using ANOVA and Ryan's Q multiple comparison test (Day & Quinn 1989) on appropriately transformed data. Nonparametric Kruskal-Wallis and Dunn's multiple comparison test procedures were used to compare all species densities because homogeneity assumptions of ANOVA were not satisfied by a majority of the individual species data. All tests were run on an IBM PS/2 using SAS 6.04 and the GLM or NPAR1WAY statistical routines (Joyner 1985). The statistical power of individual tests was calculated using Cohen (1988).

**RESULTS**

Harpacticoid copepods were the first or second most abundant group of adult metazoans in all samples and represented from 16.5 to 65.6% of all emerging, 2.4 to 30.9% of all seagrass blade, and 1.2 to 13.0% of all sediment fauna. In samples not dominated by copepods, nematodes and/or crustacean nauplii were the most abundant metazoans. Numbers of copepods emerging, on blades, or in the sediments varied between presunset and postsunset periods in 1983 (Fig. 3). Significantly greater numbers of copepods emerged during postsunset periods in March (F = 10.93; df = 1.8; p < 0.05) and May (F = 48.28; df = 1.8; p < 0.0005), but not August (F = 2.63; df = 1.8; p > 0.1) or September (F = 4.32; df = 1.8; p > 0.05). Copepod densities on postsunset seagrass blades increased significantly in March (F = 9.64; df = 1.17; p < 0.01) and August (F = 34.02; df = 1.18; p < 0.0005) but did not increase on all dates (Fig. 3). Observed postsunset decreases in blade copepod densities were significant in September (F = 5.16; df = 1.18; p < 0.05) and not significant in May (F = 2.64; df = 1.16; p > 0.1). The average blade area sampled also varied among dates, from a low of 18.4 cm² in March to a high of 70.7 cm² in August. The presunset variation in copepod numbers on seagrass blades enabled detection of a 93% (March), 76% (May), 54% (August), and 50% (Sep-
The composition of predominant emerging and seagrass blade species or species groups in 1983 was similar within but varied among sampling dates (Table 1). Combined densities of *Mesocha pygmea*, *Ectinosomatidae* spp., *Harpacticus* sp., *Laophontidae* spp., *Metis holothuriae*, *Dactylopodia* spp., and *Tisbe lurcata* constituted between 36.0% to 74.9% of all emerging and 35.8% to 83.4% of all seagrass blade copepods. No consistent changes in species densities on blades coincided with the postsunset period of increased copepod emergence (Table 1). March densities of *Ectinosomatidae* spp. ($\chi^2 = 4.34$, df = 1, $p < 0.05$) and *Laophontidae* spp. ($\chi^2 = 6.00$, df = 1, $p < 0.05$) were significantly greater on postsunset blades, but neither species group emerged in significant numbers during either presunset or postsunset periods (Table 1). Densities of *ectinosomatid* spp. ($\chi^2 = 5.67$, df = 1, $p < 0.05$) in May and *Harpacticus* sp. ($\chi^2 = 6.00$, df = 1, $p < 0.05$) in September were significantly lower on postsunset seagrass blades while both species exhibited an increase in postsunset emergence from the sediments (Table 1). Only *M. holothuriae* densities in August suggested any connection between significantly greater densities on postsunset blades ($\chi^2 = 8.64$, df = 1, $p < 0.005$) and species emergence during presunset and postsunset periods (Table 1).
Table 1 Mean (± SE) presunset and postsunset densities (per 10 cm² sediment or blade surface area) for copepod species or species groups that constituted >10% of the individuals found emerging or on seagrass blades during 1983.

<table>
<thead>
<tr>
<th>Date</th>
<th>Period</th>
<th>Emerging Species/Group</th>
<th>Density</th>
<th>Seagrass blades Species/Group</th>
<th>Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>19 Mar</td>
<td>Presunset</td>
<td><em>Tisbe furcata</em></td>
<td>0.0 ± 0.0</td>
<td>Ectinosomatidae spp.</td>
<td>1.6 ± 0.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Mesochra pygmea</td>
<td>0.2 ± 0.1</td>
<td>Laophontidae spp.</td>
<td>0.9 ± 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Laophontidae spp.</td>
<td>0.9 ± 0.8</td>
<td>Harpacticus sp.</td>
<td>2.2 ± 0.6</td>
</tr>
<tr>
<td></td>
<td>Postsunset</td>
<td><em>Tisbe furcata</em></td>
<td>2.5 ± 0.6</td>
<td>Ectinosomatidae spp.</td>
<td>6.3 ± 2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Mesochra pygmea</td>
<td>1.5 ± 0.3</td>
<td>Laophontidae spp.</td>
<td>4.5 ± 1.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Laophontidae spp.</td>
<td>0.1 ± 0.1</td>
<td>Harpacticus sp.</td>
<td>2.0 ± 0.6</td>
</tr>
<tr>
<td>20 May</td>
<td>Presunset</td>
<td>Laophontidae spp.</td>
<td>4.6 ± 1.0</td>
<td>Ectinosomatidae spp.</td>
<td>1.2 ± 0.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ectinosomatidae spp.</td>
<td>0.9 ± 0.4</td>
<td>Laophontidae spp.</td>
<td>1.0 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>Postsunset</td>
<td>Laophontidae spp.</td>
<td>21.4 ± 4.5</td>
<td>Ectinosomatidae spp.</td>
<td>17.5 ± 3.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ectinosomatidae spp.</td>
<td>19.6 ± 6.1</td>
<td>Laophontidae spp.</td>
<td>17.4 ± 1.8</td>
</tr>
<tr>
<td>12 Aug</td>
<td>Presunset</td>
<td>Metis holothuriae</td>
<td>5.3 ± 1.5</td>
<td>Metis holothuriae</td>
<td>0.6 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>Postsunset</td>
<td>Metis holothuriae</td>
<td>2.5 ± 0.8</td>
<td>Metis holothuriae</td>
<td>4.5 ± 1.3</td>
</tr>
<tr>
<td>30 Sep</td>
<td>Presunset</td>
<td>Harpacticus sp.</td>
<td>4.4 ± 0.6</td>
<td>Harpacticus sp.</td>
<td>13.5 ± 2.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dactylopodia spp.</td>
<td>3.8 ± 0.6</td>
<td>Dactylopodia sp.</td>
<td>7.1 ± 1.6</td>
</tr>
<tr>
<td></td>
<td>Postsunset</td>
<td>Harpacticus sp.</td>
<td>5.4 ± 1.1</td>
<td>Harpacticus sp.</td>
<td>5.7 ± 1.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dactylopodia spp.</td>
<td>3.8 ± 0.5</td>
<td>Dactylopodia sp.</td>
<td>5.1 ± 1.3</td>
</tr>
</tbody>
</table>

Significantly greater numbers of copepods emerged ($F = 84.75; df = 1.8; p < 0.001$) during the April 1987 postsunset sampling period (Fig. 4), but only Ectinosomatidae spp. (38.0%) and Harpacticus sp. (12.0%) constituted >10% of all emerging copepods. Both Ectinosomatidae spp. ($\chi^2 = 6.86; df = 1, p < 0.01$) and Harpacticus sp. ($\chi^2 = 6.36; df = 1, p < 0.05$) emerged in greater numbers during the postsunset sampling period. Ectinosomatidae spp. were exclusively sediment-dwelling in April and constituted <3.0% of the copepods found on blades. Harpacticus sp. were found both in the sediments and on seagrass blades and constituted 47.3% of all blade copepods.

A substantial number of either benthic or phytal copepods that entered the water column in April were collected in horizontal traps (Fig. 4). The compass orientation of horizontal traps did not affect the number of copepods collected ($F = 0.20; df = 3.8; p > 0.5$), but significantly greater numbers were trapped during the postsunset period ($F = 11.86; df = 1.8; p < 0.01$). Trap exposure by the ebbing tide for ca 30 min likely contributed to the decrease in numbers sampled during the period ending at 24:00 h. Ectinosomatidae spp. (65.0%) and Harpacticus sp. (9.4%) were the predominant copepods collected in horizontal traps.

Consistently greater numbers of copepods were enumerated from petri compared to funnel traps (Fig. 5). Among sampling periods significantly different numbers of copepods settled into petri ($F = 4.14; df = 6.26; p < 0.005$) but not funnel traps ($F = 1.06; df = 6.26; p > 0.1$). Copepod densities in petri traps were significantly greater in postsunset and presunrise samples (Ryan's Q, $p < 0.05$). Ectinosomatidae spp. were 32.5% of all the resettled copepods and were the predominant taxa collected in either trap. Over twice the number of ectinosomatids settled into petri compared to funnel traps during the night (54 vs 21 ind. 10 cm²).

Harpacticus sp. was the only species found in greater
numbers within funnel traps, but mean densities were low (1 vs 2 ind. 10 cm^-2). Postsunset and presunrise petri traps collected 200% more ectinosomatids than other periods, but there were no significant differences among sampling periods in the numbers of ectinosomatids that settled in either petri ($\chi^2 = 10.99, df = 6, p > 0.05$) or funnel traps ($\chi^2 = 7.55, df = 6, p > 0.01$). Taxa that exhibited a difference in settlement times typically settled in greater numbers sometime between 02:00 and 07:00 h and included Laophontidae spp. ($\chi^2 = 15.07, df = 6, p < 0.05$) in petri traps and Harpacticus sp. ($\chi^2 = 23.03, df = 6, p < 0.001$). Dactylopodia spp. ($\chi^2 = 16.46, df = 6, p < 0.05$), and Tisbe furcata ($\chi^2 = 14.71, df = 6, p < 0.05$) in funnel traps. Enhydrosoma herrerai and Zausodes areniculus combined constituted only a small fraction of either the emerging (4.1%) or blade copepods (0.3%), but were an appreciable percentage of the copepods found in settlement traps (9.8%). Settlement of E. herrerai was significantly different between sampling periods in both petri ($\chi^2 = 16.37, df = 6, p < 0.05$) and funnel traps ($\chi^2 = 18.95, df = 6, p < 0.005$). Greater numbers of E. herrerai settled during presunset and postsunrise periods. Z. areniculus only exhibited settlement differences in petri traps ($\chi^2 = 14.16, df = 6, p < 0.05$) where greater numbers were collected during the presunrise period.

Copepod densities on natural and defaunated blades varied throughout the night (Fig. 6). On natural blades copepod densities between sampling periods were not significantly different ($F = 2.02; df = 6.27; p > 0.05$) and suggested limited movement onto or off of seagrass blades. A 32% difference in seagrass blade copepod densities could be detected with 80% power at the 0.05 level of significance. Sampling period differences in densities on defaunated blades were significant ($F = 3.50; df = 6.27; p < 0.05$) and greater numbers of copepods settled onto defaunated blades after 24:00 h (Ryan’s Q, p < 0.05). Densities of both Harpacticus sp. and Laophontidae spp., the predominant species on blades, were greater on natural blades. On defaunated blades Harpacticus sp. densities were significantly different between sampling periods ($\chi^2 = 19.88, df = 6, p < 0.005$), but Laophontidae spp. densities were not ($\chi^2 = 3.63, df = 6, p > 0.05$). Densities of Harpacticus sp. along with Dactylopodia spp. and Tisbe furcata increased on defaunated blades between 04:00 and 07:00 h.

A summary of species-specific copepod linkages between benthic, pelagic, and phythal habitats from various studies is presented in Table 2. Of the 21 species or species groups identified in Table 2, only 3 emerged from sediments or blades exclusively. Although there is an appreciable overlap in species composition between benthic and phythal habitats, 9 emerging species were not observed to resettle onto blades. Many species like Halicryptos hydrotuge that did not resettle onto blades also did not disperse through the water column. Only 3 species exhibited unequivocal resettlement to the sediments, but 100% of the species for which data were available resettled to the sediments in at least 1 study. A difficulty in evaluating resettlement to the sediments was the inability to link emergence and settlement statistically. Significant numbers of Laophontidae spp. emerged during the April postsunset period in the present study ($\chi^2 = 5.91, df = 1, p < 0.05$), but statistically significant settlement to the sediments above presunset levels only occurred during the presunrise period (Dunn’s, p < 0.05) and therefore can not be linked directly to increased postsunset emergence. The most striking observation from Table 2 is that 76% of the emerging species were found to disperse through the water column on at least 1 occasion.

**DISCUSSION**

The effects of sediment-dwelling metazoan emergence on linkages among benthic, pelagic, and phythal habitats will depend primarily on the numbers and fates of emerging individuals. For harpacticoid copepods in subtidal seagrass beds, postsunset emergence is a common behavior involving large numbers, $10^4$ to $10^5$ ind. m^-2, and a significant percentage of the benthic population, 7 to 31% (Walters 1991). Emerging copepods can experience a number of possible fates that include pelagic dispersal and resettlement to either benthic or phythal habitats. In the present study we found that copepod emergence from benthic habitats resulted in (1) rapid (within hours) recolonization of benthic habitats by a significant portion of the
emerging fauna, (2) active dispersal through the water column, (3) resettlement that is mediated by species behavior, and (4) nondetectable colonization of phytal habitats.

The inability to detect a postsunset decrease in sediment copepod densities or consistently increased densities on seagrass blades in March to August samples suggests that copepods rapidly, within 2 h, resettled back to the sediments. On the dates sampled, between 7 and 31% of the copepod assemblage emerged during postsunset periods, which resulted in decreased sediment copepod densities inside emergence traps where resettlement was prevented (Walters 1991). A significant decrease between presunset and postsunset copepod sediment densities outside of emergence traps should have been observed unless insignificant numbers of copepods emerged, the statistical power of tests was low, phytal or other migrations compensated for numbers emerging from the sediments, or emerging copepods rapidly resettled. The nonsignificant difference between presunset and postsunset copepod numbers emerging in August and September was the result of an atypical increase in presunset emergence and not unusually low copepod emergence (Walters 1991). Although tests were able to detect small, <10%, density differences with limited statistical power, the proportion of sediment-dwelling copepods that emerged during May, 22%, and August, 31%, postsunset periods (Walters 1991) should have produced statistically detectable results. Phytal copepod densities also did not decrease significantly on postsunset blades, which would have indicated possible movement to the sediments, for dates when sediment copepod densities were available. The most parsimonious
The settlement of copepods in seagrass beds is influenced by various factors. Our study suggests that the emergence of copepods during daylight is linked to flow. Copepods collected in traps with different compass orientations indicate that the emergence of ectinosomatids is rapid (Coull & Feller 1988, Feller et al. 1990, Hicks 1992).

That copepods settle quickly to the sediments is supported by April settlement trap results. Emerging copepods colonized petri traps in greater numbers just after sunset, typically the time of greatest emergence, and before sunrise. Copepod numbers emerging during the postsunset period (Fig. 4) were more than equaled by numbers that settled into traps (Fig. 5). Although postsunset data indicate a rapid resettlement of emerging copepods, the continued nighttime and increased presunrise settlement require explanation. Copepods typically emerge in greater numbers around sunset, but emergence can occur throughout the night (Walters & Bell 1986, Walters 1988, 1991). The increased percentage of the copepod assemblage that emerges in traps deployed for 6 h (Walters & Bell 1986) and the continued collection of emerging individuals from the same patch of benthos (Jacoby & Greenwood 1988, 1989) indicate that different individuals enter the water column during the night. Some copepod species also may remain in the water column after emerging (Bell et al. 1989) or exhibit differences in the timing of peak emergence (Walters 1988). Even after 9 h Bell et al. (1989) determined that >75% of emerging Zausodes arenicolus individuals remained in the water column within field aquaria. Walters (1988) determined that 25% of all copepods that emerged from a seagrass bed entered the water column during the 2 h after sunset while 55% emerged during the remainder of the night. Assuming similar percentages for our April data, a mean of 61 copepods 10 cm⁻² would have emerged during the hours between sunset and sunrise. During the same period, a total of 62 copepods 10 cm⁻² resettled onto petri traps. The protracted settlement of copepods in traps likely reflects the night-long emergence patterns typical in subtidal seagrass habitats.

Although settlement occurred rapidly, emerging copepods did disperse through the water column (Fig. 4). Tidal or wind driven currents had a minimal effect on the pelagic dispersal of emerging copepods. All horizontal traps were deployed during an easterly sea breeze on an ebbing tide when prevailing currents at the study site were towards the southeast. If emerging copepods were transported as passive particles through the water column, greater numbers would be expected to accumulate in north- and west-facing traps. However, there was no difference in the numbers collected in traps with different compass orientations. Equivalent numbers in traps facing north, east, south, and west suggest that emerging copepods haphazardly swam or moved through the water column. Our results differ from studies in hydrodynamically active sand habitats where the distribution of settling copepods is linked to flow (Fegley 1988, Savidge & Taghon 1988, Kern 1990).

The difference in numbers and settlement patterns between petri and funnel traps indicated that species behavior affected the resettlement of emerging copepods. Although design constraints may affect settlement trap results, biases typically are more pronounced in turbulent flows (Gardner 1980a, Baker et al. 1988). Current velocities in the seagrass bed were not measured directly, but evidence suggests flows were well below critical erosion values on 20–21 April. Field flows were not strong enough to influence the directionality of copepods collected in horizontal traps, and surface conditions in the seagrass bed after 22:00 h were dead calm (Walters pers. obs.). Differences in copepod abundance between petri and funnel traps also were attributable to only one species group, the ectinosomatids. A systematic bias in collection efficiencies should have affected all copepod species equally. Without evidence to suggest a sampling bias, differences between petri and funnel traps likely reflect either selective settlement or post-settlement behavior. The almost 2 to 1 difference in Ectinosomatidae spp. settlement into petri traps may be related to the species’ sediment-dwelling morphology (Bell et al. 1987). Although numerous in seagrass sediments throughout the year, ectinosomatids only emerge in any numbers from March to May (Walters 1991, this study) and only are abundant on seagrass blades between September and October (Hall & Bell 1993). Morphology, restricted emergence, and a limited presence on seagrass blades all suggest that ectinosomatids prefer a sedimentary environment and may avoid settlement onto the artificial surfaces in a funnel trap. Whatever the reason, ectinosomatids exhibited a uniquely different settlement behavior compared to other emerging copepod species.

Some copepod species exhibited a limited emergence into the water column in April but were collected in appreciable numbers in settlement traps. Enhydrosoma herrerai and Zausodes arenicolus constituted <5% of the emerging but often >10% of the settling copepods. Previous studies have documented that Z. arenicolus is an active migrant in seagrass habitats (Service & Walters 1991), but E. herrerai typically is not an emerging species (Bell et al. 1987, Walters 1991). The increased settlement in presunrise traps suggests that Z. arenicolus emerged later in the evening. Increased presunset and postsunrise numbers of E. herrerai settling suggest emergence during daylight.
hours, but *E. herrerai* also was collected in settlement traps throughout the night. The emergence behavior of *E. herrerai* may be limited to relatively short (<15 cm) distances off the sediment surface. Limited excursions would preclude collection in emergence traps, where vertical migrations of >15 cm are required, but permit collection in settlement traps that were only 10 cm above the sediment surface. Seagrass copepods appear to exhibit a range of emergence behaviors, from the major excursions into the water column typified by ectinosomatids to the short jaunts above the sediment surface exhibited by *E. herrerai*.

Emergence of sediment-dwelling copepods is not linked to a consistent settlement onto seagrass blades. Post-sunset emergence infrequently coincided with greater copepod numbers on blades. When blade copepod densities did increase in March and August the increase was not attributable to sediment-emerging species. Colonization of April defaunated blades also was not connected unequivocally to the emergence of sediment-dwelling copepods. Densities of *Laophontidae* spp., *Dactylopodia* spp., and *Harpacticus* sp., species that are both sediment (Walters et al., 1991) and seagrass blade residents (Kurdziel & Bell, 1992, Hall & Bell, 1993), did not increase on blades during peak periods of emergence from the sediments. The pattern of recruitment onto defaunated blades cannot be explained by the post-sunset increase in emergence unless *Harpacticus* sp., *Dactylopodia* sp., and *Tisbe furcata* remained in the water column for a protracted period of time, 22:00 h until 04:00 to 07:00 h. For individual copepod species within seagrass beds it is possible that sediments and blades represent distinct subpopulations only infrequently interconnected by emergence. Our inability to document an unambiguous exchange between sediment and seagrass populations of the same species suggests that not only species identity but habitat origin is critical to predicting the effects of emergence.

The interpretation of our blade colonization results may be complicated by differences in the availability of blade surface area, an inability to detect relatively small density differences between sampling periods, and/or the migration of blade-associated copepods. Differences in blade area influenced copepod density calculations and may have influenced settlement behavior, but March and August evidence for recruitment onto blades occurred on dates when the least and greatest blade area was sampled. If emerging copepods settled onto blades, numbers that would have produced density differences of from 50 to 93% could have been detected with reasonable statistical power. Given the range in detectable density differences, it is possible that settlement of a small percentage of the sediment-emerging copepods onto blades would go unnoticed. Copepod migrations off blades were not measured directly, but previous studies indicate phytal copepods are highly mobile (Gunnill, 1982, Bell & Hicks, 1991, Kurdziel & Bell, 1992). An accurate assessment of migration effects would require measurement of copepod turnover rates on individual blades. Short-term turnover rates for macrofaunal crustaceans on blades can exceed 50% of the assemblage (Howard, 1985) and are linked to the colonization of seagrass blades (Edgar, 1992), but turnover rates for meiofauna copepods remain unknown.

Synthesis of the information available to date (i.e. Table 2) reiterates that the linkages among benthic, pelagic, and phytal habitats in seagrass systems are often species- and habitat-specific. An Ectinosomatidae sp. emerging from the sediments may not resettle onto seagrass blades, but one migrating off of blades may resettle to the sediments. Each habitat probably offers a unique set of resources and ecological interactions, but a large number of harpacticoid copepod taxa appear to participate in a continual reassortment both within and between benthic, pelagic, and phytal habitats. Detailed autecological studies on seagrass system copepods are critical for interpreting the importance of linkages between habitats and the patterns of habitat utilization that are suggested from field sampling and experimentation.

Benthic copepod species have been identified in the pelagic environment since the turn of the century (Brady, 1899, Scott, 1909), but the possible connections between emergence and water column dispersal were recognized only recently (Palmer, 1988). Although sediment-emerging copepods appear to resettle rapidly, both water column dispersal (this study) and extended pelagic residence (Bell et al., 1989) can delay benthic reentry. Dispersal of migrating phytal copepods also results in prolonged water column habitation by normally benthic taxa (Bell & Hicks, 1991, Kurdziel & Bell, 1992). The pelagic dispersal of benthic copepods will increase the probability of interacting with both nektonic and planktonic assemblages. Consumption of emerging copepods by predatory nekton and zooplankton likely will increase with an increase in the time spent in the water column and may represent a significant trophic link between benthic and pelagic habitats. Fish in seagrass habitats can switch from preying on obligate zooplankton to preying on vertically migrating crustaceans (Robertson & Howard, 1978). Along with being consumed as prey, emerging copepods themselves may consume pelagic microbes. In nearshore coastal areas benthic copepods are associated with marine snow aggregates (Shanks & Edmondson, 1990) and can influence the dynamics of aggregate microbial assemblages (Walters unpubl.). Some intertidal sediment-dwelling copepod species
feed or plankton and preferentially ingest pelagic diatoms during periods of tidal inundation (Decho 1986, 1988). While the emergence of sediment-dwelling copepods may affect both benthic and pelagic food webs, evaluating the magnitude of the effects will be difficult. Our study indicates that movements between benthic, pelagic, and phytal habitats occur rapidly and are difficult to detect without an extensive sampling regime. Determining the effects such movements have in each environment may be problematic and require the use of new techniques or unique experimental approaches.

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Gardner, W. D. (1981). Hydrodynamic processes affecting pelagic food webs, evaluating the magnitude of the effects will be difficult. Our study indicates that movements between benthic, pelagic, and phytal habitats occur rapidly and are difficult to detect without an extensive sampling regime. Determining the effects such movements have in each environment may be problematic and require the use of new techniques or unique experimental approaches.
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