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Assessment of Seagrass Plant Demography Within and Among Beds of Turtle Grass (*Thalassia testudinum*), Manatee Grass (*Syringodium filiforme*), and Shoal Grass (*Halodule wrightii*) From the Northern Gulf of Mexico

PAUL A. X. BOLOGNA AND DOUG HAYWICK

Variability in the demographic patterns of seagrasses may affect a variety of physical and biological processes in aquatic communities. We investigated seasonal seagrass demography, sediment size structure, and sediment organic carbon from two sites (north and south) in St. Joseph Bay, Florida, for *Thalassia testudinum*, *Syringodium filiforme*, and *Halodule wrightii*. Geologically, sands dominated both sites, but the south site had significantly greater proportions of fine and very fine sands, whereas the north site was dominated by coarse and medium sands. Within each site, demographic parameters were assessed for edge (<1 m) and interior portions (>10 m) of grass beds. The objective was to quantify within-habitat demographic characteristics and assess whether differences in plant biomass and shoot density existed between edge and interior portions. Results for *T. testudinum* showed significantly greater shoot density (*P* < 0.05) and plant biomass (*P* < 0.001) from interior portions of the bed than from the edge for the north site, but there were no significant differences for the south site. *Syringodium filiforme* showed significantly greater shoot density from bed interiors than from edges for both the north and south sites (*P* < 0.001, *P* < 0.01, respectively) as well as plant biomass (*P* < 0.0001, *P* < 0.01, respectively). *Halodule wrightii* showed a similar response, with greater shoot densities from the north interior (*P* < 0.008) and significantly greater plant biomass from interior portions of beds for both the north and south sites (*P* < 0.003, *P* < 0.01, respectively). Within-bay comparisons showed that plant biomass and shoot density were generally greater from the north site than from the south site for *T. testudinum* and *H. wrightii*, but the reverse was true for *S. filiforme*. Although sands dominated both sites, the south site showed greater percentages of finer sediments, and *S. filiforme* biomass has been shown to be inversely related to sediment size. Although we were unable to resolve within-habitat differences in sediment size structure, which may suggest sediment baffling and water velocity changes, we surmise that physical and sedimentary differences exist between edge and interior portions of seagrass beds. These differences may have dramatic effects on processes such as larval settlement, predator–prey interactions, and per unit area primary production. As such, further investigations into within-habitat variability and edge effects in plant demography and associated biological and physical processes are needed.

The overall structure of seagrasses communities covers a broad spectrum of plant species, composition, and areal coverage. In general, seagrass habitats are often distributed as a mosaic of vegetated cover interspersed with varying degrees of unvegetated sediments (see Larkum and den Hartog, 1989; Robbins and Bell, 1994; Marba and Duarte, 1995). These habitat mosaics, coupled with succession, often create ecosystems of variable shoot density, species composition, canopy height, and biomass (Bell and Westoby, 1986; Irlandi, 1994). Therefore, seagrass habitat architecture can be defined at many spatial and temporal scales (Robbins and Bell, 1994), and defining the extent and physical arrangement of the landscape may be essential for addressing ecological questions (Holling, 1992; Levin, 1992).

Seagrass structure is important in coastal regions because they dampen wave energy and reduce water velocity (Fonseca et al., 1982; Gambi et al., 1990). The reduction of flow associated with grass beds increases particle deposition (Almasi et al., 1987), and the extensive root–rhizome mat may bind particles, thereby stabilizing sediments (Thayer et al., 1984; Fonseca and Fisher, 1986). Seagrass beds, therefore, act as sediment traps and may retain finer sediments than do the unvegetated regions around them (Orth, 1977). The change in physical regime is often most extreme at the edge of a given grass bed. Consequently, edges...
are where current baffling would first occur and where coarser-grained sediments should first be deposited. Ideally, finer-grained sediment would be carried under conditions of lower flow further into bed interiors. Particle size gradients should, therefore, occur from the edge into the interior of seagrass beds (Orth, 1977), thus creating settlement shadows (Bologna and Heck, 2000). Consequently, because of the potential differences in physical regime and sediment composition, two different subhabitats may exist within seagrass beds: "edge" and "interior." It might be surmised that physical and biological regimes would affect seagrass bed structure. For instance, sand dune migration can affect coverage and growth patterns of some species (Marba and Duarte, 1995), whereas consumption by herbivores can greatly reduce standing stock (Camp et al., 1973) and create grazing halos (Ogden et al., 1973). Bioturbation (e.g., stingray feeding) can also contribute significantly to the destruction of local grass beds (Orth, 1975), and in some areas, decapod burrowing creates favorable conditions for storm events to tear rhizome mats (Valentine et al., 1994). Consequently, the distribution of seagrass biomass within a region may vary greatly, and areas most prone to disturbance may be affected the greatest. Under these circumstances, edge habitats might be the most dynamic because of potential differences in physical regimes (e.g., water velocity, turbulence). Few detailed studies, however, have investigated edge effects in marine seagrass communities. As such, our intent was to examine both small-scale within-habitat variability in plant demography and sediment structure among seagrass species and, on a larger spatial scale, potential differences in plant and sediment characteristics within a region, varying in physical exposure. Specifically, the objectives of this study were to assess the seasonal and within-habitat differences in plant demography and biomass in Thalassia testudinum ex Banks König, Syringodium filiforme Kuetz, and Halodule wrightii Aschers. Additionally, we assessed the potential effect these seagrasses have on structuring sediment composition and organic carbon.

**STUDY SITE**

Research was conducted in St. Joseph Bay, Florida, which lies in the Northeastern Gulf of Mexico (29°N 85.5°W; Fig. 1). It is a shallow semienclosed lagoonal system open to the Gulf of Mexico at the north and has little freshwater input. Consequently, salinity is normally high and ranges from 25 to 35‰ annually. However, large storm events (e.g., Tropical Storm Alberto) can reduce salinity by as much as 20‰ during short periods of time (Bologna, 1998). Shallow portions of St. Joseph Bay are vegetated by a mosaic of seagrass species covering 2,300–2,400 ha of the benthos (Savastano et al., 1984). Two principal study sites in St. Joseph Bay were investigated: the north site was chosen to represent a more exposed, oceanic habitat, whereas the south site was chosen to represent a physically protected habitat (Koch and Gust, 1999). The southern portion of St. Joseph Bay was more heavily used for recreational activities (e.g., fishing, boating, snorkeling) than was the northern region (pers. obs.). Both sites were shallow (<1.2 m deep mean low water) and comprised a mosaic containing T. testudinum, S. filiforme, H. wrightii, and open sediment.

**METHODS**

Plant demography.—To determine seasonal and spatial differences in shoot density and plant biomass, 15.24-cm-diameter cores (0.01824 m²) were collected from edge and interior portions of T. testudinum, S. filiforme, and H. wrightii seagrass beds at each site. Edge was operationally defined as vegetation within 1 m of a seagrass-sand interface. Interior was defined to be at least 10 m from any seagrass-sand interface (Bologna, 1998). These distinctions were chosen to represent relative extremes within each habitat. Replicate cores (n = 2, sensu Livingstone et al., 1976) were collected in Oct. 1995 and in Jan., April, Aug., and Oct. 1996. Samples were processed in the field, and plant material was frozen and returned to the laboratory. In the laboratory, seagrasses were separated into shoots (aboveground), rhizomes, and roots. Shoots were separated and counted. Shoots, roots, and rhizomes were dried to constant weight at 80 C and then ashed at 500 C for 8–10 hr to determine ash-free dry weight (AFDW). Shoot density and total plant biomass were compared among seasons (e.g., dates of collection) and within habitats (e.g., edge vs interior) using two-way ANOVA. Data were square-root transformed before analysis to normalize data, and analyses were performed on individual seagrass species for a given location (i.e., north or south). Significance testing was performed with α = 0.05. Additionally, the individual plant biomass components (e.g., root, rhizome, and aboveground) were compared...
Fig. 1. St. Joseph Bay, Florida. Research was conducted on the western side of the bay in a shallow seagrass mosaic containing *Thalassia testudinum*, *Syringodium filiforme*, and *Halodule wrightii*. Specific study regions are indicated as north and south.

within habitats for each species at each site following the same protocol.

**Sediment size fractionation analysis.**—Sediment analysis was undertaken to assess potential differences in sediment type and organic content between sites and among habitats. Replicate 5.1-cm sediment cores were taken to a depth of 10 cm at edge and interior portions of *T. testudinum*, *S. filiforme*, and *H. wrightii* seagrass beds from both north and south sites. Sediment cores were also collected from unvegetated regions adjacent to the seagrass habitats. A total of 31 samples was collected and transported to the Sedimentology and Thin-Sectioning Laboratory at the University of South Alabama. Each sample was soaked overnight in distilled water to remove salts, decanted, and dried in a low-temperature oven (50 °C) until dry. Fractions from each sample (25–40 g) were extracted using a sediment splitter for grain size analysis. Percentages of gravel (particle size >2 mm; Folk, 1980), sand (particle size 2 mm to 63 μm), and “fines” (silt + clay) were determined by dry sieving. We opted not to distinguish between silt (particle size: 63–4 μm) and clay (particle size <4 μm) because initial analyses using the pipette and sieve method of Coventry and Fett (1979) demonstrated little clay-sized content within the samples (<0.1%). Sediment composition was compared between sites and among habitats using arcsin-transformed proportion data. Between-site comparisons were analyzed using an unpaired t-test (α = 0.05).

**Sediment organic carbon content.**—Sediment samples were collected before the onset of summer growth (May) and at the end of the growing season (Oct.) in 1996 to determine potential effects of seagrass habitat on organic carbon concentration. Sediment cores of 2.5-cm diameter (n = 58) were collected to a depth of 3 cm from unvegetated and edge and interior portions of *T. testudinum*, *S. filiforme*, and *H. wrightii* from each site. Samples were transported to the laboratory, dried to constant weight at 80 °C, and ashed at 500 °C for 8 hr. The difference in weight between dry weight and AFDW was calculated and used as a percentage of total weight to determine percent organic carbon. Data were arcsin transformed.
and compared by seagrass species and analyzed as a three-way ANOVA with date of collection, site, and habitat location as independent factors and organic carbon as the dependent variable ($\alpha = 0.05$).

**RESULTS**

Plant demography.—*Thalassia testudinum*: Analyses from square-root transformed data collected from the north site showed significantly greater shoot density from interior portions of the bed than from the edge ($F_{1,10} = 5.1; P < 0.05$; Fig. 2a). Additionally, there was a seasonal component with significantly fewer shoots from samples collected in winter ($F_{4,10} = 4.5; P < 0.02$; Fig. 2a). However, no spatial or temporal pattern was evident from samples collected at the south site (Fig. 2a). When biomass was analyzed, data collected from the north site showed consistent differences in plant biomass. Specifically, leaf ($F_{1,10} = 7.6; P < 0.02$), rhizome ($F = 37.1; P < 0.0001$), and root biomass ($F = 72.4; P < 0.0001$) were significantly greater from interior portions of the bed than from edges. The summed total biomass, consequently, was also significantly greater ($F_{1,10} = 62.0; P < 0.0001$; Fig. 3a). Biomass also showed significant seasonal variability, with reduced biomass in Jan. compared with other seasonal samples. When biomass was analyzed for samples collected at the south site, no significant biomass patterns were evident, but samples collected in 1996 showed a consistent pattern of greater total biomass from interior samples than from edges (Fig. 3b).

*Syringodium filiforme*: Results showed that shoot density was significantly greater from interior portions of the bed than from edges for both north ($F_{1,10} = 45.7; P < 0.0001$; Fig. 4a) and south sites ($F_{1,10} = 9.14; P < 0.018$; Fig. 4b). Additionally, the north site showed a temporal effect with significantly fewer shoots in Aug. compared with other sampling dates ($F_{4,10} = 11.1; P < 0.002$). When sites were compared, shoot density was greater from samples gathered from the south than from those from the north, with the exception of Jan. 1996, when very low shoot densities occurred from south interior samples (Fig. 4b). When biomass was analyzed for *S. filiforme*, data from the north site showed that leaf ($F_{1,10} = 65.6; P < 0.0001$), rhizome ($F = 20.1; P < 0.001$), and root biomass ($F = 28.7; P < 0.003$) were significantly greater from interior portions of the bed than from edges (Fig. 5a). Similar results were seen for data collected at the south site, with leaf ($F = 16.6; P < 0.002$) and rhizome biomass ($F = 12.3; P < 0.006$) significantly greater from interior sites than from edges (Fig. 5b), but there were no differences in root biomass ($F = 0.4; P > 0.5$). In both cases, the resultant total biomass was significantly greater from interior portions of the bed than from edges (north, $F = 48.8; P < 0.0001$; south, $F = 9.5; P < 0.01$; Fig. 5).

*Halodule wrightii*: Results showed significantly greater shoot density from interior north *H. wrightii* beds ($F_{1,10} = 10.8; P < 0.008$; Fig. 6a); however, no pattern was evident from samples gathered at the south site (Fig. 6b). When plant biomass was analyzed, north and south sites showed the same pattern for plant biomass with significantly greater leaf ($F_{1,10} = 6.8; P < 0.02$; $F = 4.5; P < 0.06$; north and south, respectively), root ($F = 21.3; P < 0.001$; $F = 24.9; P < 0.0005$, respectively), and total biomass ($F = 15.1; P < 0.003$; $F = 10.3; P < 0.01$, respectively) from samples gathered at interior portions of the bed than from edges (Fig. 7). However, both showed no significant differences in rhizome biomass, possibly because rhizomes comprised a relatively small portion of total biomass (Fig. 7).

Sediment size structure analysis.—Sediment analysis showed that samples were primarily composed of sand (mean 98.69%, range 90–100% sand by weight), with minimal amounts of fines and gravel (gravel components consisted of broken shell fragments). There was no significant variation in grain size among edge, interior, and unvegetated samples; however, there were significant differences in grain size between the two study sites. The north site was dominated by medium-grained sands (>50%, 250–500 µm), whereas the south site was dominated by fine sands (>70%, 125–250 µm; Fig. 8). Additionally, the north site had significantly greater proportions of coarse sand (500 µm to 1 mm) than the south ($t_{1,10} = 8.9; P < 0.0001$), whereas the south had a significantly greater proportion of very fine sand (63–125 µm, $t_{3,10} = 3.7; P < 0.001$; Fig. 8). There was generally no intrahabitat difference in sediment size distribution within the north sample site. Both edge and interior were dominated by medium to fine sand. However, the south site did show a difference in particle size from edge to interior. Both edge and interior were dominated by fine sand, but there were significantly greater percentages of very fine sands from interior *T. testudinum* ($t_{2} = 4.4; P < 0.05$).
Fig. 2. Comparison of *Thalassia testudinum* shoot density among dates of collection. Data presented represent mean shoot densities expressed as number of shoots per square meter ± SE for dates of collection, with the addition of the composite mean value. Asterisk above composite value represents significant differences in mean shoot densities between edge and interior portions of the habitat. (a) North *T. testudinum* site. (b) South *T. testudinum* site.

*S. filiforme* (*t*₂ = 4.6; *P* < 0.05), and *H. wrightii* (*t*₂ = 24.7; *P* < 0.002) than from edges.

**Organic carbon content.**—Generally, sediment organic carbon was greater from samples gathered in May than from samples gathered in Oct. (Table 1). Only the north *H. wrightii* interior site did not show this pattern. These results suggest that during winter, accumulation of detritus occurred, creating differences in organic carbon concentrations in the sediments. When percent carbon was compared between vegetated and unvegetated habitats (i.e., sand; Table 1), data showed greater content in veg-
Fig. 3. Comparison of Thalassia testudinum plant biomass among dates of collection. Data presented represent mean total biomass for dates of collection expressed as grams AFDW per square meter ± SE, with the addition of the composite mean value. Asterisk above composite value represents significant differences in mean plant biomass between edge and interior portions of the habitat. (a) North T. testudinum site. (b.) South T. testudinum site.

DISCUSSION

Understanding the distributional aspects of seagrass biomass may be essential for assessing processes within and among seagrass communities (Turner et al., 1999). Because these habitats are often vital to the survival of species (Rasmussen, 1973; Thayer and Stuart, 1974) and processes such as recruitment (Peterson, 1986), predation (Heck and Crowder, 1991), and growth (Bologna and Heck, 1999a) may
act differently between vegetated and unvegetated habitats, the essential structure of seagrass beds may, to a large degree, determine the productivity (Robertson, 1979; Edgar, 1990a, 1990b; Bologna and Heck, 2002) and trophic transfer within a system (Bologna, unpubl. data). Results from this research indicate that seagrass demography had not only strong seasonal variability, but also significant within-habitat spatial variability. Shoot density and seagrass biomass were greater from interior portions of continuous seagrass beds than from edges for *T. testudinum* (Figs. 2, 3), *S. filiforme* (Figs. 4, 5), and *H. wrightii* (Figs. 6, 7),

Fig. 4. Comparison of *Syringodium filiforme* shoot density among dates of collection. Data presented represent mean shoot densities expressed as number of shoots per square meter ± SE for dates of collection, with the addition of the composite mean value. Asterisk above composite value represents significant differences in mean shoot densities between edge and interior portions of the habitat. (a) North *S. filiforme* site. (b) South *S. filiforme* site.
Fig. 5. Comparison of *Syringodium filiforme* plant biomass among dates of collection. Data presented represent mean total biomass for dates of collection expressed as grams AFDW per square meter ± SE, with the addition of the composite mean value. Asterisk above composite value represents significant differences in mean plant biomass between edge and interior portions of the habitat. (a) North *S. filiforme* site. (b) South *S. filiforme* site.

and this response has also been reported for other systems as well (e.g., *T. testudinum*, Zie­
man, 1972; *Zostera marina*, Orth, 1977; *Cymodocea nodosa*, Duarte and Sand-Jensen 1990a, 1990b). However, other investigators have not identified significant edge effects or simply assumed that among-habitat differences were more important than within-habitat differenc­
es. Our results show that shoot density for all species showed high variability, with greater shoot density occasionally occurring at edges but averaged greater at interior sites (Figs. 2, 4, 6). On the other hand, only once was the biomass greater from an edge sampling date.
Fig. 6. Comparison of Halodule wrightii shoot density among dates of collection. Data presented represent mean shoot densities expressed as number of shoots per square meter ± SE for dates of collection, with the addition of the composite mean value. Asterisk above composite value represents significant differences in mean shoot densities between edge and interior portions of the habitat. (a) North H. wrightii site. (b) South H. wrightii site.

(Fig. 3b), and this corresponded to winter biomass minima. These results may have significant implications on our assessment of edge communities in marine systems.

Several factors may make edge habitats more productive. At edges, competition for light and nutrients may be reduced. Edges, with greater turbulent flow (Fonseca et al., 1982), may see increases in light through reflection from the sand, incident from the side, and increased flashing to the lower portions of the canopy. The greater turbulence may also decrease diffusive boundaries for plants (Koch, 1994), thereby increasing the transfer of nutrients
Fig. 7. Comparison of *Halodule wrightii* plant biomass among dates of collection. Data presented represent mean total biomass for dates of collection expressed as grams AFDW per square meter ± SE, with the addition of the composite mean value. Asterisk above composite value represents significant differences in mean plant biomass between edge and interior portions of the habitat. (a) North *H. wrightii* site. (b) South *H. wrightii* site.

and wastes (Wheeler, 1980), which may lead to increased primary production (Fonseca and Kenworthy, 1987). Because interior portions of the bed are relatively quiescent, nutrients and dissolved inorganic carbon may have to travel through relatively thick diffusive boundaries, possibly limiting primary production (Koch, 1994) compared with the more turbulent boundaries encountered at edges (Fonseca et al., 1982). Edges may also encounter greater water-column nutrient potential because water entering the meadow may become relatively depleted in nutrients as it continues to interior portions of a bed. Additionally, there may be
Fig. 8. Comparisons of sediment characteristics between north and south sites. Values expressed as percent total for each given size class ± SE. Abbreviations: G, gravel; VCS, very coarse sand; CS, coarse sand; MS, medium sand; FS, fine sand; VFS, very fine sand; FINES, fine particulate material (e.g., silt and clay combined < 63μm). Asterisks above bars indicate significant differences in percent abundance between north and south sites.

Reduced competition for sediment nutrients at seagrass bed edges because of the lower shoot density and plant biomass there, and this constitutes a major source of nutrients for submerged vegetation (see reviews by Short, 1987; Barko et al., 1991). This may be of great importance for relatively pristine regions such as St. Joseph Bay, Florida (Rutkowski et al., 1999), but less so for regions undergoing eutrophication (Lapointe et al., 1994; Short and Burdick, 1996).

Because there are intimate relationships between seagrass flora and fauna, understanding the distributional aspects of the flora provides essential evidence in the functioning of seagrasses as habitat. Clearly, our results show significant differences in shoot density (Figs. 2, 4, 6) and biomass (Figs. 3, 5, 7) between edge and interior regions for all species investigated. These differences in plant demography may have important effects on growth, predation, and recruitment of associated organisms (Edgar and Robertson, 1992). It has been shown that the density and species richness of seagrass-associated fauna are often related to the biomass and plant surface area (Stoner and Lewis, 1985). However, recent evidence has shown that although biomass and shoot density may be lower at edges, faunal density may be significantly greater (Bologna, 1998). In some cases, organisms may show a settlement shadow (sensu Orth, 1992) elevating densities at edges (Bologna and Heck, 2000), but evidence

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<td>G</td>
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Tidal height (m) Interior
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T. erectum

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T. sphaerocystis

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<td>T. sphaerocystis</td>
<td>0.45 ± 0.02</td>
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exists that for brooding organisms like amphipods, increased faunal density at edge habitat must confer some benefit to them (Bologna, 1998) and may be related to increases in secondary production (Bologna and Heck, 2002).

Although within-bay differences in plant demography existed in St. Joseph Bay, seagrass biomass in this region is among the highest seen in the Caribbean and Gulf of Mexico (Table 2). Regional comparisons indicate that plant biomass for all seagrass species from St. Joseph Bay more closely resembles biomass patterns from tropical regions rather than from subtropical ones (Table 2). In fact, biomass values for both \textit{S. filiforme} and \textit{H. wrightii} are among the highest values ever recorded for these species. These results are even more striking because our comparisons are made based on AFDW as opposed to dry weight alone (Table 2). It is interesting to note that only \textit{S. filiforme} had greater biomass at the south site than at the north site (Fig. 5). This may relate to the sediment size in the south (Fig. 8), as Iverson and Bittaker (1986) showed an inverse relationship between mean sediment size and \textit{S. filiforme} biomass and shoot density in the eastern Gulf of Mexico, and our results concur.

The differences in grain size characteristics between the north and south site could be attributed to energy regime. The north site is in a more open (higher energy) area of St. Joseph’s Bay and consequently, contained coarser sediment than did the south site (Fig. 8). It should be noted, however, that both sites were characterized by relatively well-sorted sand. The lack of fine sediment (e.g., silt and clay) within our study region is significant but not overly surprising. Because there is relatively little riverine input to the system (Wolfe et al., 1988), there may be limited amounts of fine material being deposited from terrestrial sources within St. Joseph Bay. Additionally, both sites may have been situated in relatively high-energy regions and consequently, any fine material that might have been carried over the seagrass beds may have stayed in suspension regardless of baffling effect provided by the seagrasses. Our south site did indicate that finer sediment occurred within the interior portion of beds for all seagrass species (Fig. 8), but as our data were not consistent across the study area, we are forced to conclude that if sediment baffling was occurring within these habitats, it was not resolvable using grain size criteria alone. Seagrass beds in St. Joseph Bay may show a slight positive relief (e.g., 10–30 cm; Bologna, 1998), which suggests that they are sites of increased sedimentation (cf. Almasi et al., 1987), or sediment retention (Fonseca and Fisher, 1986), and both are indicative of sediment baffling. However, it still may be possible to identify and quantify the importance of sediment baffling by seagrasses in sand-dominated environments through more intensive sampling. It is both desirable and ultimately necessary to assess the role of sediment baffling in seagrass beds because sediments may be a major source of nutrients (Short et al., 1990; Barko et al., 1991), and sediment baffling by various organisms (sea grasses, solitary corals, crinoids etc.) has been suggested as a means of stabilizing sediments before colonization by metazoans (e.g., Walker and Alberstadt, 1975; James and Bourque, 1992). Additionally, seagrasses provide surface area for coralline algal epiphytes (Bologna and Heck, 1999b), and this has been shown to increase the concentrations of \textit{CaCO}_3 in the sediments (Land, 1970). These changes in sediment composition in relatively pristine regions may then lead to phosphate limitation (Short et al., 1990; Duarte et al., 1995).

Although a great deal of information exists regarding seagrass demography and distribution, we have a very limited understanding of edge dynamics. Because edges are regions in which beds are either expanding or are being destroyed through mechanical disturbance (e.g., boat moorings, crab burrows), we need to understand their community dynamics. This is especially true in regards to plant–animal interactions occurring at edges because the distribution of fauna, as well as the processes affecting their distribution, may be substantially different compared with interior regions of vegetated habitat. Continuing research into edge effects is needed to assess habitat value, as well as per unit primary and secondary production in coastal ecosystems. This will become increasingly important because loss and fragmentation of seagrass beds continue worldwide.

\section*{Acknowledgments}

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<td>25–250*</td>
<td>10–50</td>
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<tr>
<td>NE Gulf of Mexico</td>
<td>50*</td>
<td></td>
<td>3–10</td>
<td></td>
</tr>
<tr>
<td>SE Gulf of Mexico</td>
<td>75–275*</td>
<td>25–250*</td>
<td>10–50</td>
<td></td>
</tr>
<tr>
<td>Pensacola, FL</td>
<td>20–130</td>
<td></td>
<td>3–10</td>
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<tr>
<td>Florida Keys</td>
<td>0–215</td>
<td>0–107</td>
<td>0–87.5</td>
<td></td>
</tr>
<tr>
<td>Mississippi Sound, MS</td>
<td>120–180</td>
<td></td>
<td>10–60</td>
<td></td>
</tr>
<tr>
<td>Chandelier Sound, LA</td>
<td>52.3</td>
<td>314.9</td>
<td>4–42</td>
<td>35–119</td>
</tr>
<tr>
<td>Guadeloupe Estuary, TX</td>
<td>42–166</td>
<td>4–42</td>
<td>17.5–29*</td>
<td></td>
</tr>
<tr>
<td>Laguna Madre, TX</td>
<td>77.8 ± 26.2*</td>
<td>37.8 ± 5.6*</td>
<td>20–70</td>
<td>50–210</td>
</tr>
<tr>
<td>St. Croix, US Virgin Islands</td>
<td>207 ± 52</td>
<td>1092 ± 209</td>
<td>24.5 ± 24.4</td>
<td>17.5–29*</td>
</tr>
<tr>
<td>Yucatan Peninsula, Mexico</td>
<td>222–693</td>
<td>261–467</td>
<td>57.7 ± 46.2</td>
<td></td>
</tr>
<tr>
<td>Cabelo Gordo de Dentro, Brazil</td>
<td>4–33</td>
<td>16–55</td>
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</table>

* Indicates values are Ash Free Dry Weight.
assistance in the laboratory. This is contribution \# 344 to the Dauphin Island Sea Lab.

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