Enhancement of sediment suspension and nutrient flux by benthic macrophytes at low biomass

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ABSTRACT: In shallow coastal ecosystems where most of the seafloor typically lies within the photic zone, benthic autotrophs dominate primary production and mediate nutrient cycling and sediment stability. Because of their different structure and metabolic rates, the 2 functional groups of benthic macrophytes (seagrasses, macroalgae) have distinct influences on benthic–pelagic coupling. Most research to date in these soft-bottomed systems has focused on mature seagrass meadows where shoot densities are high and on dense macroalgal mats that accumulate in response to eutrophication. Relatively little is known about the influence of low-biomass stands of seagrass and macroalgae on nutrient fluxes and sediment suspension. Using an erosion microcosm with controlled forcing conditions, we tested the effects of the eelgrass *Zostera marina* L. and the invasive macroalga *Gracilaria vermiculophylla* on sediment suspension and nutrient fluxes under high-flow conditions. At low densities, *G. vermiculophylla* increased sediment suspension and increased the nutrient flux from the sediment to the water column. For macroalgae, increased sediment suspension is likely due to dislodgement of sediment particles by bedload transport of the algae. In this case, the increase in sediment transport was reflected in an increase in nutrient flux from the sediment, showing that modification of physical forcing by benthic primary producers can also affect nutrient flux. The presence or absence of *Z. marina* did not have a significant effect on nutrient flux. However, the results suggest that there may be a range of low shoot densities for which storm-like flows increase sediment suspension to values higher than those expected for a bare sediment bed.

KEY WORDS: Benthic macrophytes · Sediment suspension · Nutrient flux · Seagrass · Benthic–pelagic coupling · Coastal bay · Lagoon

INTRODUCTION

In shallow coastal ecosystems where the seafloor typically lies within the photic zone, benthic autotrophs dominate primary production and mediate nutrient cycling and sediment stability (Sand-Jensen & Borum 1991, McGlathery et al. 2007). The strong benthic–pelagic coupling in these systems is influenced by both the distinct structure and the metabolic rates of the 2 functional groups of benthic macrophytes, viz. seagrasses and macroalgae (McGlathery et al. 2007). Almost all studies on the effects of seagrasses and macroalgae have been conducted on mature or high-density beds (e.g. Gambi et al. 1990, Fonseca & Cahalan 1992, Escartín & Aubrey 1995, Sfriso et al. 2005). Compared to these high-density populations, we know little about the effects of lower densities of benthic macrophytes on sediment and nutrient fluxes, even though it is common for seagrasses and macroalgae to occur at low densities. For example, multiple stressors, including sediment and nutrient runoff, algal blooms, and physical disturbances from storms, boat traffic, and some fishing practices, can cause thinning and/or patchiness in seagrass populations in addition to wholesale losses of seagrass habitat (Duarte 2002, Orth et al. 2006). Restoration of seagrass meadows, especially by seeding, also results initially in low-density populations.
that increase in density as the meadows develop over time (McGlathery et al. 2012, Orth et al. 2012, both in this Theme Section). Macroalgae can also be present at low densities in shallow coastal systems depending on availability of attachment structures or advection of ephemeral populations by wind-driven currents (Thomsen & McGlathery 2005).

The effect of benthic macrophytes on near-bed hydrodynamics and the resulting sediment/nutrient fluxes may be different at low density than at high density. Extensive research has shown that dense seagrass canopies reduce current velocity (Peterson et al. 2004) and dampen wave energy (Fonseca & Cahalan 1992), leading to decreased sediment resuspension and increased particle deposition (Heiss et al. 2000, Peterson et al. 2004). Observations of sediment stabilization in seagrass beds and increased turbidity following seagrass decline (e.g. Cottam & Munro 1954, Christiansen et al. 1981, Stumpf et al. 1999) have supported a conceptual model that sediment suspension increases when seagrass is replaced by macroalgae (e.g. Sand-Jensen & Borum 1991, Boynton et al. 1996). However, dense macroalgal mats can have similar stabilizing effects. Thick mats decrease sediment suspension (Sfriso & Marcomini 1997, Romano et al. 2003) and shear flow at the sediment surface (Escartin & Aubrey 1995). Both dense seagrass meadows (Gambi et al. 1990) and dense macroalgal mats (Escartin & Aubrey 1995) deflect flow around the bed, creating an area of low shear stress at the sediment surface within the bed. However, research on other types of emergent features in coastal systems, such as polychaete worm tubes, has shown that low-density stands typically displace velocity around individual features, rather than over or around the larger-scale bed/patch, leading to increased erosion around the features (Eckman et al. 1981, Friedrichs et al. 2000, Widdows et al. 2002). When flow is diverted around isolated low-density features, turbulent wake structures can cause areas of high bed shear stress to develop around the features, with the potential to cause scour (e.g. Sadeque et al. 2008), similar to that commonly observed around an isolated bridge pier. Macrophyte density, patch size, and flow velocity interact to determine whether flow, or some portion of the flow, is diverted around the entire bed/patch or individual features. Isolated areas of scour and decreased sediment accumulation have been documented in field, laboratory, and modeling studies of low-density stands of emergent features (Bouma et al. 2007).

Most of the experimental studies to date on the effects of macrophytes on nutrient fluxes across the sediment–water interface have been conducted in artificially-calm hydrodynamic conditions such as stirred laboratory cores or in situ chambers. It is well known that local hydrodynamics influence nutrient fluxes through porewater advection (Huettel & Webster 2001), desorption from suspended sediments (Morin & Morse 1999), and thinning of the diffusive boundary layer (Kelly-Gerreyn et al. 2005). Benthic macrophytes can have significant effects on these fluxes physically, by deflecting flow and changing the hydrodynamic conditions at the sediment surface, and biologically, by uptake and indirect effects on bacterial nutrient transformations in the sediment (McGlathery et al. 2007).

Here we report results from a study using controlled microcosm experiments to measure the effects of low-density seagrass and macroalgal populations on sediment suspension and nutrient fluxes. The motivation for this study was the local extinction and subsequent large-scale restoration of the seagrass Zostera marina (eelgrass) in the Virginia coastal bays that are part of the Virginia Coast Reserve Long Term Ecological Research (VCR LTER) site (McGlathery et al. 2012, Orth et al. 2012). Extensive seagrass populations were decimated in this region in the 1930s as a result of 2 natural events: a slime mold wasting disease and a severe hurricane. The loss of eelgrass in the mid-Atlantic region of the USA produced a state change in which the perennial, rooted eelgrass was replaced by ephemeral, attached and drift macroalgae, including the invasive species Gracilaria vermiculophylla. Fishery species, including scallops, declined and turbidity increased (Cottam & Munro 1954), similar to changes seen in other systems (Stumpf et al. 1999, Yamamuro et al. 2006).

Our experiments were carried out using a small-scale microcosm with controlled hydrodynamic conditions, allowing us to isolate the effects of the primary producers on sediment stability and nutrient fluxes. The use of microcosms with a controlled flow field permits both a mechanistic look at how flow conditions and primary producers interact and direct comparisons between 2 types of primary producers at varying densities. While this approach does not provide ecosystem-level results, microcosm-scale experiments are valuable to direct ecosystem-level studies and parameterize models.

**MATERIALS AND METHODS**

To determine the effects of Zostera marina and Gracilaria vermiculophylla on sediment suspension and nutrient flux, sediment cores with varying bio-
mass were exposed to flow conditions that generate a known bed shear stress in bare sediment cores. Five sampling sites with patchy distributions of eelgrass and macroalgae (attached and detached) were selected randomly within a region of restored eelgrass meadows in South Bay at the VCR LTER site (Fig. 1). At each site, 5 sediment cores (4 bare sediment and 1 with eelgrass) were taken. Each site was sampled on 1 day (18 to 22 June 2007), and bare sediment cores were randomly assigned to a control, low macroalgal biomass, medium macroalgal biomass, or high macroalgal biomass treatment (no medium algal biomass treatment was done on Day 3 due to the loss of 1 sediment core). The study was designed as a randomized complete block design, blocked by day and plot, with 5 cores planned per treatment (1 core was lost and 1 was re-classified resulting in 5 cores for all treatments except low algae). Sediment and water column samples were also taken at each site to characterize the sites for any differences in initial conditions.

**Shear stress experiments**

Sample collection, storage, and preparation

Sediment samples for the experiments were taken by hand on a falling tide (water depth approximately 0.8 to 1 m), within 1 h of predicted low tide, using polycarbonate core tubes with an internal diameter of 10.8 cm and a height of 22.5 cm, with a 10 cm water column preserved above the sediment surface. Core tubes were sealed using rubber plumbing end caps and kept in the dark at 4°C during transport back to the lab. Ambient water was collected in 20 l carboys to use as replacement water during the experiments and for core storage until analysis. Upon arrival at the lab, within 1 h of core collection, the cores were submerged in a cooler of ambient water oxygenated by an aquarium bubbler and maintained in low light at ambient temperatures. Macroalgae were collected in South Bay and maintained in a container of ambient water at 4°C for transport to the lab, then kept in oxygenated ambient water until use. Immediately prior to the experiment, the overlying water in each sediment core was gently replaced with the same water that would be used for replacement water during the experiment. No sediment suspension or disturbance was noted during this step.

Bare sediment cores were assigned randomly to 1 of 4 treatments (control, low macroalgae, medium macroalgae, and high macroalgae), and analysis order of the treatments was assigned randomly. Macroalgae treatments were defined as 2 (low), 4 (medium), or 6 (high) g wet weight (WW) of macroalgae, standardized by blotting excess water with a paper towel. Macroalgae were added to a bare sediment core immediately before the start of the experiment. At the end of the experiment, the macroalgae were collected and dried at 60°C to a constant weight to determine dry weight (DW) biomass. The macroalgae treatments, based on the average field biomass of 54 g DW m⁻² found in an adjacent bay (Thomsen et al. 2006), resulted in a range of dry biomass with averages ± SE of 18 ± 0.83 g DW m⁻² (low), 37 ± 2.1 g DW m⁻² (medium), and 66 ± 4.4 g DW m⁻² (high).

Height limitations of the microcosm required eelgrass to be trimmed to a maximum height of 8 cm prior to running the experiments. Trimming to this height allowed the blades to remain submerged without interfering with the functioning of the microcosm. While the trimming created shoots that were shorter than found in the field during June (the time of this experiment), the shorter length is representative of early spring and late fall shoots found.
in the nearby Chesapeake Bay (Orth & Moore 1986). The eelgrass cut at the beginning of the experiment was dried at 60°C and weighed. At the end of the experiment, all remaining eelgrass aboveground biomass was cut and also dried and weighed so that the biomass of the eelgrass cut before the experiment, the aboveground biomass in the cores during the experiment, and the total aboveground biomass could be calculated for each core. Total aboveground biomass of the eelgrass (before trimming) in the cores ranged from 64 to 470 g DW m⁻². This is comparable to 76 to 558 shoots m⁻², based on field measurements of densities and biomass per shoot from the study site (McGlathery et al. 2012). The aboveground biomass of the eelgrass in the cores during the experiments (i.e. the 8 cm tall blades that remained after trimming) ranged from 41 to 184 g DW m⁻².

Shear stress manipulation

The Gust microcosm used in these experiments consisted of an erosion head with a rotating plate and a push-through water system fitted over a polycarbonate core tube (Fig. 2; Thomsen & Gust 2000). With a level, bare sediment bed, the combination of the rotation of the plate and the suction of the water system in the center of the core generates a near-uniform bed shear stress and diffusive boundary layer thickness, though not a uniform pressure distribution (Tengberg et al. 2004). Because sediment conditions were similar between treatments and the hydrodynamic forcing was controlled, changes in sediment suspension should be a direct result of alterations to the near-bed hydrodynamics. Across treatments, any increase in the mass of sediment eroded from the bed should reflect localized increases in bed shear stress caused by the treatment. The use of Gust microcosms for studying benthic fluxes is further examined in the ‘Discussion’.

Immediately prior to the experiment, a water sample was pumped from the replacement water to provide background concentration values. The cores were then exposed to low forcing conditions (forcing that produces a low shear stress of 0.01 N m⁻² in bare sediment cores) for 20 min as a flushing step, followed by high forcing conditions (forcing that produces a shear stress of 0.32 N m⁻² in bare sediment cores) for 40 min. This shear stress is in the range of storm-generated bed shear stresses in the VCR (Lawson et al. 2007, Mariotti et al. 2010) and roughly corresponds to an average velocity of 0.25 m s⁻¹ in the microcosm (without obstructions), similar to the flow rates used in other studies of sediment suspension/stabilization in Zostera marina beds (e.g. Fonseca & Fisher 1986). Effluent water was collected in 1 l Nalgene bottles exchanged every 5 min for the higher shear stresses. All effluent water from the erosion tests was collected, subsampled, and filtered for total suspended solids (TSS), NH₄⁺, PO₄³⁻, and NO₃⁻. No fragmentation of macroalgae or eelgrass was noted during the experiments.

Site characterization

Bottom sediment samples were taken at each plot to characterize porewater nutrients, sediment grain size, exchangeable ammonium (NH₄⁺), and benthic chlorophyll. Five replicate samples for grain size and NH₄⁺ were taken to a depth of 2 cm with a modified 60 ml syringe, and 5 replicate samples for benthic chlorophyll were taken to 1 cm depth with a modified 5 ml syringe. Three replicate sediment cores were collected in polycarbonate tubes (8 cm internal diameter, 23 cm height) for laboratory sampling of porewater. All samples were kept at 4°C during transport to the lab. Samples for benthic chlorophyll and extractable NH₄⁺ were immediately frozen; grain size samples were maintained at 4°C until analysis. Porewater samples were extracted from the sediment cores at 2 cm depth intervals to 10 cm depth using a
stainless steel probe (Berg & McGlathery 2001) and were immediately filtered (0.45 µm) and frozen. Exchangeable NH$_4^+$ was extracted for 1 h with an equal volume of 1 N KCl solution. Porewater samples (for NH$_4^+$, phosphate PO$_4^{3-}$, and nitrate+nitrite NO$_3^-$), and exchangeable NH$_4^+$ samples were analyzed on a Lachat QuickChem 8500 using standard methods (Hach). Benthic chlorophyll was determined spectrophotometrically following extraction with a 45% acetone, 45% methanol, 10% deionized water solution using the equations of Lorenzen (1967). Sediment grain size was analyzed as 1 bulk sample per plot, from 5 pooled samples (0 to 2 cm depth), using a combination of wet sieving for sands and analysis on a Sedigraph 5100 particle size analyzer for silt and clay.

**Laboratory analyses**

All sediment and water samples were analyzed using standard methods. TSS (sample volumes 150 to 550 ml) was analyzed by weight difference after filtration onto pre-combusted, pre-weighed Whatmann GF/F filters (nominal particle retention 0.7 µm). Sub-samples for nutrients (20 ml) were filtered (0.45 µm) and frozen until analysis. NH$_4^+$, PO$_4^{3-}$, and NO$_3^-$ were analyzed using standard methods on a Lachat QuikChem 8500 (as described above). NO$_3^-$ and nitrite (NO$_2^-$) were not separated because NO$_2^-$ values were frequently below the detection level. The combined value is presented as NO$_3^-$ throughout this study.

To correct for the mass of measured components carried in the replacement water, all fluxes were corrected for background concentration. For the nutrient fluxes, the average concentration of all background samples for the day was used as the background concentration. For the mass of sediment eroded, the lowest measured concentration of TSS during the analysis of the core was used. Data from the flushing step were discarded, and only data from the higher shear stress step were analyzed. One macroalgal treatment was reclassified (from low to medium) at the end of the experiment based on the measured DW of the macroalgae (33 g DW m$^{-2}$).

**RESULTS**

**Site characterization**

We found no significant differences in the initial conditions at the 5 plots in South Bay (Table 1). Sediment extractable NH$_4^+$ ranged from 0.065 to 0.087 µmol g DW$^{-1}$ with an average ± SE of 0.08 ± 0.00 µmol g DW$^{-1}$. Porewater concentrations of NH$_4^+$ ranged from 0 to 152 µM, PO$_4^{3-}$ from 0 to 16 µM, and NO$_3^-$ from 1.2 to 6 µM. The sediment in all plots was very fine sand, with between 5 and 11% mud (average grain size = 71 µm). Water column nutrient concentrations, as measured from the replacement water, were 0.5 ± 0.1 µM NH$_4^+$, 0.4 ± 0.1 µM PO$_4^{3-}$, and 0.3 ± 0.1 µM NO$_3^-$, similar to values measured in neighboring Hog Island Bay (McGlathery et al. 2001).

**Sediment suspension**

During the 40 min high-forcing experiments, an average of 0.26 ± 0.03 g of sediment was eroded for all treatments, which is equivalent to an average erosion depth of 2.8 × 10$^{-3}$ m, assuming a bulk density (DW of sediment per unit volume of bed) of 1.0 g cm$^{-3}$, or 28.7 g m$^{-2}$ (see Figs. 3 & 4). There was no significant effect of eelgrass presence on sediment suspension compared to bare sediment for the flow conditions and biomass used in these experiments (paired t-test, p = 0.4, Fig. 3). However, closer analysis of the variation in eelgrass biomass within the replicate cores indicates that at low biomass (corresponding to shoot densities of approximately 76 to 263

<table>
<thead>
<tr>
<th>Day</th>
<th>Average grain size (µm)</th>
<th>Benthic chlorophyll (mg m$^{-2}$)</th>
<th>NH$_4^+$ (µM)</th>
<th>PO$_4^{3-}$ (µM)</th>
<th>NO$_3^-$ (µM)</th>
<th>Extractable NH$_4^+$ (µM)</th>
<th>NH$_4^+$ (µM)</th>
<th>PO$_4^{3-}$ (µM)</th>
<th>NO$_3^-$ (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1</td>
<td>76</td>
<td>4.7 (0.8)</td>
<td>0.4 (0.2)</td>
<td>0.3 (0.1)</td>
<td>0.5 (0.1)</td>
<td>0.06 (0.00)</td>
<td>62 (5)</td>
<td>4.3 (0.5)</td>
<td>2.2 (0.2)</td>
</tr>
<tr>
<td>Day 2</td>
<td>70</td>
<td>5.2 (0.2)</td>
<td>0.2 (0.1)</td>
<td>0.2 (0.0)</td>
<td>-</td>
<td>0.09 (0.00)</td>
<td>43 (5)</td>
<td>3.5 (0.6)</td>
<td>1.7 (0.3)</td>
</tr>
<tr>
<td>Day 3</td>
<td>76</td>
<td>5.6 (0.9)</td>
<td>0.6 (0.2)</td>
<td>0.4 (0.2)</td>
<td>0.2 (0.1)</td>
<td>0.08 (0.00)</td>
<td>82 (7)</td>
<td>5.7 (0.4)</td>
<td>2.0 (0.1)</td>
</tr>
<tr>
<td>Day 4</td>
<td>67</td>
<td>4.5 (0.6)</td>
<td>0.6 (0.1)</td>
<td>0.4 (0.0)</td>
<td>0.3 (0.1)</td>
<td>0.07 (0.00)</td>
<td>34 (9)</td>
<td>4.7 (0.2)</td>
<td>2.6 (0.3)</td>
</tr>
<tr>
<td>Day 5</td>
<td>66</td>
<td>6.1 (0.8)</td>
<td>0.7 (0.1)</td>
<td>0.5 (0.2)</td>
<td>0.3 (0.2)</td>
<td>0.08 (0.02)</td>
<td>42 (3)</td>
<td>3.4 (0.5)</td>
<td>2.6 (0.3)</td>
</tr>
</tbody>
</table>
239 shoots m$^{-2}$), sediment suspension appeared to increase with increasing biomass, whereas at higher biomass, sediment suspension dropped to almost 0. Moore (2004) found a lack of sediment trapping in seagrass beds of similar low biomass, but a mechanistic look at internal resuspension has not previously been conducted. A more complete understanding of the structure of this relationship and the biomass or density corresponding to peak suspension in eelgrass and control cores difficult and lead to no significant differences between sediment suspension in eelgrass and control cores (paired t-test, p = 0.4).

Macroalgal biomass had a significant effect on sediment suspension, with an initial increase above control values for the low biomass treatment, followed by a decrease for the medium and high biomass treatments (Fig. 4). Although high variability in nutrient fluxes within each treatment resulted in no significant differences between treatments for macroalgae (ANOVA, NH$_4^+$ p = 0.9, PO$_4^{3-}$ p = 0.4, NO$_3^-$ p = 0.2) or eelgrass (t-test, NH$_4^+$ p = 0.2, PO$_4^{3-}$ p = 0.08, NO$_3^-$ p = 0.4), some trends in the data are apparent. The sediment flux was related to the NH$_4^+$ flux ($R^2$ = 0.98, p = 0.007, Fig. 6) for the macroalgal treatments and the control. The close relationship between macroalgal biomass and mass of sediment eroded (Fig. 5) makes it difficult to distinguish effects of nutrient uptake by macroalgae from effects of hydrodynamic forcing.

**Nutrient flux**

For the macroalgae treatments, the NH$_4^+$ and PO$_4^{3-}$ fluxes showed a similar pattern to the sediment suspension, with an increase in flux between the control and low biomass treatment, followed by a decrease for the medium and high biomass treatments (Fig. 4). The mass of sediment eroded varied with macroalgal biomass (ANOVA, p = 0.04, n = 14, Fig. 5). The control treatment was excluded from this regression analysis because of the initial increase in sediment erosion from the control to low biomass treatments.
However, if nutrient fluxes were dependent on biomass, the control point would not fit the regression of mass of sediment eroded and biomass, because the control point is not part of the linear relationship between mass of sediment eroded and biomass (Fig. 4).

**DISCUSSION**

Our data provide evidence that low-biomass macroalgal populations can increase both sediment suspension and nutrient efflux under high-flow conditions. The data also suggest that at low densities, seagrass may not be effective at stabilizing sediment and may in fact enhance sediment suspension. Most previous research on high-biomass populations has shown that seagrass and macroalgal mats reduce sediment suspension (Sfriso & Marcomini 1997, Heiss et al. 2000, Romano et al. 2003, Peterson et al. 2004) and nutrient efflux (Thybo-Christesen et al. 1993, Sundback et al. 2000, McGlathery et al. 2004, 2007), although recent experiments on macroalgae by Canal-Vergés et al. (2010) are an exception. Because our results differ from those of most previous studies, these experiments give insight into conditions that may favor sediment destabilization by benthic primary producers.

**Applicability of microcosm studies to field conditions**

While microcosms can never exactly mimic field conditions, the Gust microcosm has been widely used to represent near-bed flows for the purpose of measuring benthic fluxes (Ziervogel & Bohling 2003, Porter et al. 2004a, Stevens et al. 2007, Dickhudt et al. 2009, Wiberg et al. in press). The Gust microcosm creates a relatively uniform bed shear stress through the action of a spinning plate just below the water surface and a push-through water system. With this forcing, flow paths in the Gust microcosm occur in a logarithmic spiral with an upward vortex in the mid-
dle of the core (Thomsen & Gust 2000). Because the flow path includes a significant vertical component, it more closely resembles shallow wave action or a combined wave-current flow than a simple uniform current, which is appropriate for shallow lagoons where wave forcing is dominant (Lawson et al. 2007, Mariotti et al. 2010). Gust microcosms have been used to measure benthic fluxes on sediment cores from a wide range of environments, including the Baltic Sea (Ziervogel & Bohling 2003), Adriatic shelf (Stevens et al. 2007), Chesapeake Bay tributaries (Porter et al. 2004a, Dickhudt et al. 2009), the northern Atlantic off the coast of the UK (Black et al. 2003), and tidal flats in southwest Washington State, USA (Wiberg et al. in revision). Sediment erodibility measured using Gust microcosms has been shown to compare well with measurements made using other techniques (e.g. in situ inverted flume; Dickhudt et al. 2009) and to yield calculated suspended sediment concentrations in good agreement with field measurements (Traykovski et al. 2007).

An extensive comparison of the Gust microcosm and other benthic chambers showed that the Gust microcosm produced the most uniform bed shear stress and diffusive boundary layer thickness of the chambers studied (Tengberg et al. 2004). This uniformity is important when introducing obstructions to the chamber, making the Gust microcosm an appropriate choice for this experiment. To confirm that the forcing conditions chosen for the experiments reasonably reflect the forcing conditions in benthic environments, the ratio of mean velocity (0.25 m s\(^{-1}\)) to the shear velocity (0.018 m s\(^{-1}\)), assuming a saltwater density of 1030 kg m\(^{-3}\)) in the chamber was compared with the ratios found in natural benthic environments (Porter et al. 2004b). The calculated value for the flow conditions without obstructions (14.1) is within the range seen in natural environments (13.9 to 22.8, Porter et al. 2004b). The presence of macrophytes is expected to decrease average bed shear stress because of drag on the macrophytes.

### Stabilizing versus destabilizing effects of seagrass

Sediment stabilization from seagrass beds typically occurs due to the high flow resistance offered by a dense seagrass bed (Gambi et al. 1990, Bouma et al. 2007). However, at low densities and high-flow rates, water predominantly moves through, rather than over or around a seagrass bed. In sparse seagrass beds, suspended sediment concentrations may be equal to those in unvegetated areas because the turbulent stresses are similar (Luhr et al. 2008). The interaction of flow with individual shoots causes wakes to form behind the shoots. At Reynolds numbers, Re = \(Usd/\nu \approx 200\), where \(U\) is velocity, \(d\) is shoot diameter, and \(\nu\) is kinematic viscosity, the wakes are turbulent (Nepf 1999). At relatively low shoot densities, increases in wake-generated turbulence production can be large enough relative to frictional decreases in velocity to cause an increase in turbulent kinetic energy (Nepf 1999) and local bed shear stresses (Sadeque et al. 2008) when vegetation present compared to unvegetated conditions. If the wake-generated shear stresses at the bed exceed the critical shear stress of the bed sediment, scour and increased sediment suspension can result, as seen with other isolated emergent features (e.g. Eckman et al. 1981, Friedrichs et al. 2000, Widdows et al. 2002). Assuming a representative blade width of 3 mm (Hansen & Reidenbach 2012, this Theme Section), Re = 600 for our experiments.

In addition to density and flow velocity, flow characteristics (i.e. wave or current dominated) may affect the ability of the seagrass to stabilize sediment. When seagrass is exposed to wave forcing, a mean current is generated within the canopy rather than a skimming flow developing over the top of the canopy (Luhr et al. 2010). Also, while seagrass blades in unidirectional flow bend and physically shield the sediment bed, seagrass blades in oscillatory flows move with the flow and can enhance the exchange between the sediment bed and the water column (Koch & Gust 1999). Taller seagrass in unidirectional flow may shield the bed (as noted by Fonseca & Fisher 1986); however, short seagrass shoots that are more representative of early spring and late fall conditions in the field do not provide bed shielding in wave-like flow conditions. Previous research has shown that these low-density and low-biomass beds are ineffective at trapping suspended sediment under riverine conditions (Moore 2004), but their influence on internal resuspension and wave-dominated flows has not been studied.

### Stabilizing versus destabilizing effects of macroalgae

Density and flow conditions may have similar effects on the ability of macroalgae to stabilize sediments. While dense macroalgae mat deflect flow around the mat (Gambi et al. 1990, Escartín & Aubrey 1995) similar to dense seagrass beds, isolated individual or individuals in patchy distributions are directly exposed to the undisturbed flow conditions.
Macroalgae such as *Gracilaria vermiculophylla*, which are frequently found detached, are often transported as bedload. Bedload transport of macroalgae is an important component of macroalgal transport (Flindt et al. 1997, 2004), and has been shown to significantly increase sediment suspension at densities of 17 to 33 g DW m⁻² (Canal-Vergés et al. 2010). Combining that study with the results of our experiment and the results seen in high-density mats (Escartín & Aubrey 1995, Sfriso & Marcomini 1997, Romano et al. 2003, Sfriso et al. 2005) shows that high-biomass populations of macroalgae stabilize sediment while low-biomass populations destabilize sediment. Given the high seasonal variability in macroalgal biomass (e.g. Thomsen et al. 2006), the stabilizing/destabilizing effect of macroalgae is also likely seasonal. A vertical component of flow, found in the microcosm and in wave-dominated flows, may make the macroalgae more likely to be transported because fronds are lifted away from the bed and into the faster-moving flow in the water column, whereas in unidirectional flows, the macroalgae likely lie closer to the sediment surface. The movement of these fronds in current-versus wave-dominated flows may be analogous to the differences seen in seagrass movement in unidirectional and oscillatory flow conditions (Koch & Gust 1999).

Bedload transport of macroalgae can destabilize sediments by dislodging particles in a process similar to the widely documented increase in sediment suspension due to saltating or abrading particles (e.g. Houser & Nickling 2001, Thompson & Amos 2002, 2004). In fine-grained beds, the critical shear stress required to initiate erosion is often greater than the stress required to maintain the sediment in suspension. Under these conditions, an object (e.g. macroalgae) that impacts or scrapes the bed while moving across it can dislodge particles and significantly increase sediment suspension/erosion. We observed higher sediment erosion in experiments in which the macroalgae were in motion compared to cases in which the macroalgae were stationary.

**Factors affecting nutrient fluxes**

The results of this study show that the effects of macrophytes on benthic nutrient fluxes are not limited to biotic effects. Modification of near-bed flow by benthic macrophytes can influence sediment–water column nutrient fluxes. Field and modeling studies have shown that increased hydrodynamic activity and sediment suspension influenced nutrient flux in areas such as Lake Okeechobee, Florida, USA (Chen & Sheng 2003, 2005), the Satilla River Estuary, Georgia, USA (Zheng et al. 2004), and the Mediterranean shelf (Gremare et al. 2003). Modification of near-bed flow by benthic primary producers can produce similar effects. Benthic primary producers are well known to indirectly affect nutrient cycling, including effects on nitrogen fixation, nitrification, and nitritification/denitrification (e.g. Viaroli et al. 1996, Hansen et al. 2000, McGlathery et al. 2004). Nutrient uptake by benthic primary producers can also significantly reduce the flux of remineralized nutrients to the water column (Thybo-Christesen et al. 1993, Sundbäck et al. 2000, McGlathery et al. 2004, 2007). The total impact of benthic primary producers on the flux of nutrients between the sediment and water column is thus the result of both physical modification and biotic influences, yet these have rarely been studied in tandem.

Our data suggest that the effect of primary producers on nutrient flux during a forcing event depends on the balance between uptake and physical modification of the flow near the sediment surface. In low-flow cores or microcosm incubations, it has been shown that benthic algae can prevent the efflux of nutrients from the sediment to the water column (e.g. Sundbäck et al. 2000, Tyler et al. 2001), with the balance between benthic autotrophy and heterotrophy having a significant influence on nutrient efflux (Eyre & Ferguson 2002). This uptake effectively decouples nutrient cycling in the sediment and the water column (Kristensen et al. 2000, Anderson et al. 2003). Our results show that when macroalgal biomass is low and flow conditions are relatively high, physical dynamics result in an enhancement of nutrient flux to the water column in addition to increased sediment suspension. This can be seen in the relationship between sediment suspension and NH₄⁺ fluxes (Fig. 6). Under high-flow conditions, processes such as porewater advection and desorption of nutrients from suspended particles increase nutrient fluxes above the levels supported by diffusion alone (Ward et al. 1984, Huettel et al. 1998, Morin & Morse 1999). The movement of algae caused by the increased shear stress likely enhances this effect. In a related study, we investigated the mechanisms accounting for the increased nutrient efflux from sediments in our microcosm experiments and showed that hydrodynamically-forced fluxes (advection, suspension/desorption) were similar in magnitude to diffusive fluxes, indicating that low-flow incubations may underestimate nutrient fluxes by 50% if higher flows are present (Lawson 2008).
In summary, our data show that seagrass and macroalgae can have both stabilizing and destabilizing effects on the sediment when exposed to forcing conditions that reflect the high flows representative of storm conditions in shallow bays. Low biomass, or densities, of macroalgae can destabilize sediments, leading to increased sediment suspension and nutrient efflux; the same may be true of low-density seagrass beds. The mechanisms responsible for the destabilization are different for the 2 macrophyte functional groups: increases in near-bed turbulence as flow moves through seagrass shoots versus dislodgement of sediment particles by bedload transport of macroalgae. The threshold between stabilizing and destabilizing effects is biomass-dependent, and suggests a ‘window of vulnerability’ to negative feedbacks when biomass or densities are low and flows are high. This is relevant to restored seagrass populations where seedling results in initially low-density populations that eventually increase in biomass and density as the meadow develops over time. It is also relevant to shallow coastal systems that have been impacted by nutrient over-enrichment where multiple stresses create patchiness and thinning of seagrass populations, and macroalgal biomass may accumulate on seasonal time scales.

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