INTRODUCTION

In shallow-water ecosystems, benthic and water column environments are closely coupled (Dame 1996). Bivalve filter feeders can play an important role in benthic-pelagic coupling by transferring particles and nutrients from the water column to the sediments in the form of biodeposits (Jordan 1987, Bayne & Hawkins 1992). The introduction of bivalves has been proposed as an additional method to reduce phytoplankton biomass (Dame et al. 1980), suggesting an increase in bivalve populations as a method to restore natural balance to improve water quality in these ecosystems (Cloern 1982, Officer et al. 1982, Cohen et al. 1984, Newell 1988). The use of Eastern oysters Crassostrea virginica (Gmelin) to mitigate eutrophication in the Chesapeake Bay has been suggested (Cerco & Noel 2007, Kellogg et al. 2014, Cerco 2015) and the abundance of oysters in the Chesapeake Bay is currently being increased through both
restoration (Schulte & Burke 2014) and aquaculture (Williamson et al. 2015, Ray et al. 2015) activities.

The euryhaline, epibenthic, bivalve filter feeder *C. virginica* filters large volumes of water and efficiently filters particles larger than 3 µm from the water column (Haven & Morales-Alamo 1970). Large amounts of feces and pseudofeces as biodeposits reflect a transport of particulate organic matter from the water column to the sediments (Jordan 1987), unless currents resuspend and transport the material (Lund 1957, Widdows et al. 1998). Recent studies have indicated that biodeposits can be resuspended (Golden et al. 2016) and in fact must be transported away from aquaculture sites to avoid deleterious sediment effects (Giles et al. 2009, Testa et al. 2015), such as ammonium effluxes and anaerobic sediments without macrofauna below oyster rafts. Hargrave et al. (2008) developed a nomogram of biogeochemical indicators over an organic enrichment gradient.

It has also been suggested that nutrient regeneration from bivalve biodeposits may counterbalance the removal of phytoplankton biomass by stimulating new phytoplankton blooms (Doering et al. 1986, Asmus & Asmus 1991, Soucull et al. 2001), at least in oligotrophic waters (Cranford et al. 2007). High rates of ammonium flux can occur in sediments underlying natural bivalve populations (Dame et al. 1989, 1991, Asmus & Asmus 1991, Dame & Libes 1993), but other factors such as microphytobenthos abundance also affect nutrient transformations and regeneration (Sundbäck & Granéli 1988, Sundbäck et al. 1991, 2000). Incubation experiments have shown that particulate organic matter, oxygen conditions, and microphytobenthos abundance affect nitrogen transformations and regeneration from the sediments (Caffrey et al. 1993, Enoksson 1993, Newell et al. 2002).

There have been no studies to date that specifically examine interactions between biodeposit resuspension and changes in water quality on oyster reefs or at aquaculture farms. It is generally assumed that oyster biodeposits stay where they fall (Newell et al. 2005, Kellogg et al. 2013). Existing studies of sediment resuspension and benthic processes have shown that resuspension not only enhances nutrient fluxes from sediments (Qin et al. 2004, Almroth et al. 2009, Corbett 2010, Porter et al. 2010, Almroth-Rosell et al. 2012) but also significantly affects microbial and planktonic communities, and thus ultimately biogeochemical cycles (Wainright 1987, 1990, Porter et al. 2010, Isobe & Ohte 2014). While the current focus is primarily on how oysters can fuel the biogeochemically driven nitrogen removal from sediments (Piehler & Smyth 2011, Kellogg et al. 2013, 2014, Smyth et al. 2013), the effect of biodeposit resuspension is less clear. The fraction of deposited material that is resuspended must be known to determine oyster environmental benefits (Cerco 2015).

To study the effects of oyster biodeposits and benthic boundary-layer flow on benthic-pelagic coupling processes in a controllable whole-ecosystem context, 1000 l shear turbulence resuspension mesocosms (STURM) were used. All of the tanks had the same realistic (Porter et al. 2018) volume-weighted water column turbulence levels (RMS turbulent velocity ~1.87 cm s⁻¹, energy dissipation rate ~0.3434 cm² s⁻³) and high area-weighted bottom shear stress (~0.62 Pa). Experiments were run without oysters and without a sediment bottom, and biodeposits collected from oysters (set up separately) were added daily to 3 of 6 tanks.

The specific questions addressed were: (1) How does oyster biodeposit resuspension affect nutrient transformations, nutrient regeneration, and the nitrogen budget? (2) How do resuspended biodeposits affect phytoplankton abundance and composition in whole-ecosystem experiments? (3) How does oyster biodeposit resuspension affect ecosystem processes such as the zooplankton community? Our experiments were designed to determine if the combined effect of high bottom shear stress and oyster biodeposits directly, indirectly, or non-linearly affect ecosystem processes and water quality. The experiment took place at the Morgan State University Patuxent Environmental and Aquatic Research Laboratory, where the outdoor STURM facility is located, from 6 July to 3 August 2016.

**MATERIALS AND METHODS**

**Setup of mesocosms, biodeposit resuspension, and mixing**

The experiments reported here primarily focused on water column processes as mediated by biodeposit resuspension and did not use oyster filtration and did not include a sediment bottom. Six cylindrical Shear TURbulence Resuspension Mesocosms (STURM) tanks (see Fig. S1 in the Supplement at www.int-res.com/articles/supp/m586p021_supp.pdf) were set up. Each tank contained a single paddle that produced high instantaneous bottom shear stress to resuspend biodeposits and create realistic water column turbulence levels, without overmixing the water column (Porter et al. submitted). The STURM
tanks are the successor design of large linked mesocosms reported by Porter et al. (2004a,b) and are further described by Sanford et al. (2009), Mason & Porter (2009), and Porter et al. (2010, 2013). The mixing goal for this experiment was to set up tidal resuspension where biodeposits would be resuspended during mixing ‘on’ phases and settling during mixing ‘off’ phases by keeping water column turbulence levels realistic. Moreover, the goal was to achieve a realistic profile of total suspended solids (TSS) over water column depth: lower near the surface and higher near the bottom.

A single paddle was used that slowly rotated in a forward-stop-backward-stop motion (9 s − 1.5 s − 8 s − 1.5 s) at 18.79 rpm to avoid the entire water column rotating evenly (thus not mixing) during the mixing ‘on’ phase. Mixing was turned on for 4 h and off for 2 h in all systems to simulate tidal cycles throughout a 4 wk experiment. The tanks had a water volume of 1000 l, a water column depth of 1 m, and a sediment surface area of 1 m², but contained no sediment for this experiment. Three tanks each received a daily addition of oyster biodeposits (5.8 ± 3.3 mg TSS daily) starting in the afternoon of Day 2 of the experiment. Biodeposits were generated in an oyster holding area separate from the experiment using natural plankton in flow-through conditions.

Direct flow and turbulence measurements were made using a Sontek 10 Mhz Acoustic Doppler Velocimeter to determine water column RMS turbulent velocity and energy dissipation rates at predefined locations throughout the tank, and at different mixing speeds. RMS turbulent velocity is defined in Tennekes & Lumley (1972) as:

\[ q = \frac{1}{3}(\langle u^2 \rangle + \langle v^2 \rangle + \langle w^2 \rangle) \]  

(1)

where \( \langle u^2 \rangle \), \( \langle v^2 \rangle \), and \( \langle w^2 \rangle \) are the variances of their respective velocity components. Energy dissipation rates were determined following Sanford (1997). Volume-weighted average RMS turbulent velocities as determined using Surfer (Golden Software) were ~1.87 cm s⁻¹ (Fig. 1a), which are at the low end of values measured in continental shelf benthic boundary layers (Heathershaw 1976). Volume weighted energy dissipation rates were ~0.343 cm² s⁻³ during mixing (Fig. 1b), similar to values found in continental shelf benthic boundary layers (Gross et al. 1994) or near the surface at breaking waves (Terry et al. 1996). Energy dissipation rate was about
half as high as in the high mixing treatment of Petersen et al. (1998). These turbulence levels are slightly higher than used in previous experiments comparing resuspension vs. non-resuspension systems (Porter et al. 2010) and in linked mesocosms (Porter et al. 2004a,b), but are in the realistic range (Porter et al. 2018, Table 1 in Sanford 1997). This mixing setting kept energy dissipation rates, which are non-linearly related to RMS turbulent velocity (Sanford 1997), at realistic levels (Porter et al. 2018).

Shear stress was determined at the bottom of the tank by using hot film anemometry (Gust 1988). Shear (or ‘friction’) velocity ($u_*$ in cm s$^{-1}$), defined by:

$$u_* = \sqrt{\frac{\tau_b}{\rho}}$$  \hspace{1cm} (2)

where $\tau_b$ is bottom shear stress in dynes cm$^{-2}$, and $\rho$ is the density of water in g cm$^{-3}$, was measured at 5 locations across the tank radius. Bottom shear stress [Pa] was calculated as:

$$\tau_b = \frac{u_*^2 \rho}{10}$$  \hspace{1cm} (3)

At the chosen mixing setting, area-weighted maximum instantaneous bottom shear stress reached 0.62 Pa in all tanks (Fig. 2) and produced biodeposit resuspension in the R_BD tanks during the mixing ‘on’ phases, while settling occurred during the mixing ‘off’ phases (Fig. 3) over a 4 wk period. These shear stress levels are found in continental shelf and microtidal estuarine benthic boundary layers (Grant et al. 1984, Sanford & Halka 1993, Gross et al. 1994) and macrotidal estuarine boundary layers (Wright et al. 1992, Johnson et al. 1994). In addition, a vertical profile of TSS concentration over water column depth was measured to be similar to profiles in estuaries (e.g. Liu et al. 2014) when shear stress is high enough to resuspend sediments off the bottom and move them up into the water column (Fig. 4).

The tanks were wrapped in aluminum-coated reflective bubble wrap (Shelter Institute) to reduce overheating of the tank water during high outdoor summer temperatures (up to 38°C). In addition, 2 layers of window screen mesh were placed over the superstructure to reduce insolation, ~1.5 m above the tanks.

Fig. 2. Bottom shear stress during mixing ‘on’ phases in all systems. Shown is data from 1 of 6 shear stress sensors distributed radially across the bottom of a STURM tank. Data from all sensors were area-weighted.

Fig. 3. Tidal resuspension of biodeposits with resuspension and settling of total suspended solids (TSS) in the 3 R_BD tanks (T3, T4, and T5) from 31 July to 1 August 2016 as measured with OBS-3 turbidity meters.
The tanks were slowly and evenly filled with unfiltered 12.9 g kg\(^{-1}\) salinity water containing the resident plankton community from the Patuxent River, a tributary of Chesapeake Bay. Only megazooplankton were excluded. Mixing began with the programmed tidal cycles and all tanks were synchronized. The experiment took place over 28 d, from 6 July 2016 to 3 August 2016.

Ten percent of the water in each tank was exchanged daily at the end of the mid-day mixing ‘off’ phase to mimic tidal exchange and replaced with 0.5 µm absolute filtered Patuxent River water. Filtered water was used for the exchange (e.g. Porter et al. 2004a,b, 2010, 2013). Any nutrients added with the fill water were accounted for in the nitrogen budget. The tank walls were cleaned of periphyton daily to minimize wall growth (Chen et al. 1997) and the wall material was left in the tank to avoid affecting any mass balances (Porter et al. 2004b, 2010, 2013).

**Biodeposit additions**

Starting on Day 2, equal amounts of oyster *Crassostrea virginica* biodeposits were added to each of 3 R_BD tanks during the late afternoon mixing ‘on’ phase. The sampling of particulates and nutrients on Day 2 was done in the morning before any biodeposits had been added to the tanks and reflects all tanks without any biodeposit additions. Oyster biodeposits were collected daily from 218 oysters (10.4 ± 1.7 cm long) held in flow-through conditions of ambient Patuxent River water in an indoor raceway. Thus, biodeposits from 72.7 oysters m\(^{-2}\) were added to each tank daily. Total suspended solids (TSS), particulate inorganic matter (PIM), particulate organic matter (POM), and quality (ratio POM/PIM) of the biodeposits added were determined daily. Particulate carbon (PC), particulate nitrogen (PN), chlorophyll \(a\) (chl \(a\)), and phaeophytin concentrations were determined every 2 d. On average 5.8 ± 3.3 g TSS, 632 ± 365 mg PC, 76.5 ± 50.1 mg PN, 14.9 ± 9.2 mg particulate phosphorus (PP), 1.42 ± 0.83 mg chl \(a\), and 6.3 ± 3.5 mg phaeophytin in the biodeposits were added daily to each tank, i.e. per m\(^{-2}\), with a ratio of POM to PIM of 0.6 ± 0.2. Newell et al. (2002) define low, medium, and high biodeposit additions as 0.25, 2.5, and 5 g C m\(^{-2}\) d\(^{-1}\), respectively, and our biodeposit additions were in the medium range.

**Temperature, salinity and dissolved oxygen**

Dissolved oxygen concentrations were measured daily during mixing ‘on’ phases in the afternoon at mid depth using a YSI 95 sonde. Water temperature was measured at 10 min intervals in all tanks over the experiment using Campbell T107 temperature probes connected to a Campbell CR10x data logger (Campbell Scientific). The 6 tanks tracked each other closely, and the Campbell temperature data were analyzed in detail to determine the temperature variability in the systems. During Week 3, during a heat wave, temperatures in the tanks were compared daily to water temperatures in a shallow local Patuxent River cove. The temperatures were similar, and no cooling intervention took place to further cool the tank water. Silver bubble wrap and 2 layers of window screen above the tanks were sufficient to shield the tanks from excessive heat. Salinity was measured weekly using a refractometer.

**TSS, particle settling, particulate and dissolved nutrients**

Geochemical variables measured included TSS concentrations, particulate nitrogen (PN), particulate carbon (PC) and particulate phosphorus (PP) concentrations, as well as water column dissolved nutrient concentrations (ammonium, nitrate plus nitrite, dissolved inorganic nitrogen [DIN], phosphate, total nitrogen [TN], and total phosphorus [TP]) measured biweekly during morning mixing ‘on’. Particulates were also measured during the mixing ‘off’ that followed on Days 13, 20 and 27. Particle bulk settling speeds were measured during the mid-day mixing ‘off’ phases in the R_BD tanks using optical backscatter (OBS-3) turbidity data.
Turbidity was measured continuously at 1 s intervals in each tank with optical backscatter (OBS-3, D&A Instrument Company) turbidity sensors located at mid-depth. Turbidity was calibrated with concurrently collected mid-depth TSS samples, analyzed by filtration and weighing as described below. In post-processing, OBS data were averaged over 66 s intervals.

Water was filtered through pre-ashed, pre-weighed 47 mm Whatman GF/F filters (0.7 μm nominal pore size) and rinsed with an isotonic solution of ammonium formate to remove salts (Berg & Newell 1986). To determine TSS mass, filters were weighed on a Mettler AE 240 microbalance after drying for 24 h to constant weight at 60°C. Then the filters were ashed at 450°C for 4 h and re-weighed to determine particulate inorganic matter (PIM) concentrations. Particulate organic matter (POM) was determined by the difference between TSS and PIM (Berg & Newell 1986).

Known volumes of water were filtered through 47 mm Whatman GFF filters for particulate phosphorus (PP). In addition, water was filtered through pre-ashed 25 mm Whatman GFF filters to measure particulate nitrogen (PN), and particulate carbon (PC) concentrations. Nutrients and water column chl a and phaeophytin were analyzed by the Analytical Services Laboratory, University of Maryland Center for Environmental Science (Chesapeake Biological Laboratory Analytical Services Laboratory 2017a,b). Water samples were analyzed for particulate phosphorus (PP) following Environmental Protection Agency (EPA) Method 365.1 ‘ASPILA’ (1979). Particulate carbon (PC) and nitrogen (PN) samples were analyzed on a Model CE-440 Exeter Analytical (CHN) analyzer using EPA Method 440.0 (Zimmerman et al. 1997). The solute was captured and frozen in individual vials until it was analyzed for dissolved nutrients including ammonium (NH₄⁺), nitrate + nitrite (NO₃⁻ + NO₂⁻), dissolved phosphate (PO₄³⁻), dissolved silicate (Si), total dissolved nitrogen (TDN) concentrations, and total dissolved phosphorus concentrations. Ammonium (SM 4500-NH3 G-1997), nitrite plus nitrate (ASTM D-7781, EPA 352.2 Cadmium), and phosphate were analyzed on a Konelab Aquakem 250 multi-wavelength automated discrete photometric analyzer, and total dissolved nitrogen and phosphorus were analyzed using EPA Method 353.2 and EPA Method 365.1, respectively. Dissolved organic phosphorus (DOP) was calculated by the difference between total dissolved phosphorus and phosphate (PO₄³⁻). Dissolved organic nitrogen (DON) was calculated by the difference between TDN and dissolved inorganic nitrogen (DIN). Dissolved silicate was analyzed using a Konelab Aquakem 250 multi-wavelength automated discrete photometric analyzer with oxalic acid added to minimize interference from phosphates (EPA Method 366.0, 1997). For quality control of all variables, each sixth sample was analyzed in duplicate.

Exchange water was also sampled for dissolved nutrients to track any nutrient inputs through a 10% daily water exchange. In addition, the fill water was sampled for particulates to confirm that their abundance in the fill water was low (8.3 ± 4.7 mg l⁻¹). The same techniques were used to analyze particulates in the biodeposits. Duplicate subsamples were analyzed randomly for quality control during each sampling.

During each water sampling, the exact times of sampling for each OBS-3 turbidity meter were recorded to establish linear calibration curves of TSS versus OBS volts. The linear calibrations for each OBS-3 turbidity meter (TSS = 135.5 × OBS_V + 13.941, R² = 0.94; TSS = 1122.87 × OBS_V + 15.486, R² = 0.93; TSS = 60.53 × OBS_V + 19.324, R² = 0.92) were used to determine TSS concentrations in the tanks from continuous OBS turbidity measurements.

Bivalves repackage organic matter into biodeposits, and we expected the size structure and settling of the particles to be different in the R_BD compared to R tanks. Bulk particle settling speed (mm s⁻¹) was measured in the 3 R_BD tanks using the settling profiles generated by the OBS-3 turbidity meters during mixing-off phases. We used the distance from the surface to the turbidity sensor (0.5 m depth) divided by the time to reduce the initial TSS concentration by 50% of the range to its steady state value. TSS concentrations measured 3 times during the mixing ‘on’ and mixing ‘off’ phases during the experiment were not significantly different in the R tanks, particles did not settle, and, thus, no bulk settling speeds were determined in the R tanks. More frequent samples of TSS, PIM, POM, POM:PIM, PN, PP, PC, and chl a were collected during 3 mixing off periods on Days 13, 20, and 27 to examine changes in the particulate properties over time during settling.

Profiles of TSS concentrations over tank depth were determined 6 times over the experiment during the mixing ‘on’ phases. OBS readings, calibrated for TSS, were taken for 5 min each at 5, 10, 20, 30, 40, 50, 60, 70, and 80 cm from the surface. This was as close to the bottom as possible without interfering with the paddle. TSS concentrations over depth were graphed
and increased with increasing depth until ~50 cm and then stayed constant to 80 cm.

**Phytoplankton, light, microphytobenthos**

Biological variables measured included water column chl \( a \), phaeophytin, phytoplankton identification, and cell count measured biweekly and during 3 mixing ‘off’ times on Day 13, 20, and 27. In addition, light profiles, Secchi depth, and sediment chl \( a \) and microphytobenthos pigments were measured. Known volumes of water were filtered through 47 mm Whatman GF/F filters for chl \( a \) and phaeophytin concentrations. Chl \( a \) concentrations were measured using fluorometric techniques without acidification (Welschmeyer 1994) after extraction with 90% acetone to provide estimates of phytoplankton biomass. Phaeophytin was measured following EPA Method 445. In addition, water subsamples were preserved biweekly in Lugol’s iodine solution for later microscopic phytoplankton identification and cell counts. Phytoplankton cells were counted using Utermöhl procedures (Marshall & Alden 1990, Lacouture 2001). Phytoplankton biomass was determined by converting mean cell volumes of individual taxa to cell carbon according to Smayda (1978) and Strathmann (1967). To determine if the light regime in the R and R_BD tanks affected phytoplankton, the ratio of chl \( a \)-phytoplankton carbon was determined.

For the first 4 d of the experiment, an intensive study of surface irradiances was carried out at each of 4 quadrants in each tank to determine if the mesh cover was limiting light and to determine the most representative location in each tank to measure light profiles. Light levels of ~255 µmol photons m\(^{-2}\) s\(^{-1}\) were measured at the water surface of the R tanks and R_BD tanks, using a LI-192 Underwater Quantum sensor (LI-COR Biosciences) attached to a model LI-250 readout. In previous experiments, it was found that light levels of ~160 µmol photons m\(^{-2}\) s\(^{-1}\) are required at the water surface to prevent light limitation (Porter et al. 2004b). Therefore, any light limitation within the tanks was due to the impact of biodeposit resuspension and/or the density of phytoplankton that resulted.

Subsequently, light profiles were measured twice a week during mixing ‘on’ and ‘off’ phases measuring the total downwelling photosynthetically active radiation (PAR, in the 400 to 700 nm waveband) at 0, 10, 25, 50, and 60 cm from the water surface. Light attenuation coefficients were determined and irradiance levels at the tank bottom calculated. In addition, the mean geometric irradiance was calculated in the water column as:

\[
\text{Mean geometric irradiance} = \exp[0.5 \times (\ln(E_0) + \ln(E_{\text{Sed}}))] 
\]

where \( E_0 \) and \( E_{\text{Sed}} \) are irradiances at the surface of the water column and the bottom, respectively. The values obtained for mean geometric irradiance were similar to irradiance values measured at 50 cm depth. Secchi depth was measured almost daily during the mixing ‘on’ and ‘off’ phases using a PVC cap painted black and white, similar to a Secchi disk, mounted onto a PVC rod and lowered into the water column until it disappeared from view.

Sediment chl \( a \) and microphytobenthos pigment concentrations were measured at the end of the experiment. Using razor blades, microphytobenthos was scraped off duplicate 4 cm × 15 cm and 10 cm × 4 cm fiberglass plates that were affixed with aquarium sealant to the tank bottoms at 0–10 cm from the outer wall and 20–35 cm across the radius of the tanks before the experiment to allow easy removal of microphytobenthos. Sediment pigment samples were frozen at −70°C pending analysis with high performance liquid chromatography (HPLC; Van Heuvelen & Thomas 2001). Microphytobenthos pigments were extracted by adding acetone to the sample containers, transferring the sample to sample tubes, and sonicating each tube at 0°C. The supernatant was decanted after centrifugation and subsequently analyzed by HPLC. HPLC was used to prevent interference from degradation products such as chlorophyllides and phaeophorbides that interfere with fluorescence measurements (MacIntyre et al. 1996) and affect chl \( a \) readings. Chlorophyllides and phaeophorbides can be abundant in areas with bivalves (e.g. Karakassis et al. 1998). The level of chl \( a \) without degradation products was determined by subtracting phaeophytin and phaeophrbide from total chl \( a \). The ratio of 19’ Butanoyloxyfucoxanthin, 19’ Hexanoyloxyfucoxanthin, Alloxanthin, Diadinoxanthin, Diatoxanthin, Fucoxanthin, Peridinin, Zeaxanthin, Lutein, and Violaxanthin to chl \( a \) was determined to see if there were differences between the microphytobenthos community in the R and R_BD tanks at the end of the experiment. Due to the difficulty in the transfer of solute from the original sample containers of R tanks to the sonications, it is not clear if all pigment was conveyed, and the microphytobenthos biomass is likely underestimated in the R tanks.
Mesozooplankton and macrofauna

Biological variables measured included mesozooplankton abundance and species composition as well as macrofauna abundance. Mesozooplankton were sampled twice a week during mixing ‘off’ by pumping 40 l tank\(^{-1}\) at 27 l min\(^{-1}\) through a 63 µm Nitex screen, using a diaphragm pump. The samples were washed into bottles and preserved with buffered formaldehyde. The dominant taxa in the mesozooplankton and age groups were determined on a dissecting microscope using direct counts. To estimate the dry weights of the different taxa, the number of individuals was multiplied by their respective weight characteristic (Table 2 in White & Roman 1992; W, µg ind\(^{-1}\)). Thus, the number of adult *Acartia tonsa* for each sampling day was multiplied by 8, the number of polychaete larvae by 1.3, the number of copepods by 2.7, the number of copepod and barnacle nauplii by 0.31, the number of barnacle cyprids by 2.8, crab zoea by 20 (Torres et al. 2002) and veliger larvae by 0.1342 (Sprung 1984). The dominant taxa were copepod nauplii, *Acartia* sp. adults, polychaete larvae, and copepodes.

Zooplankton densities (ind. l\(^{-1}\)) were converted to carbon (µg l\(^{-1}\)) for each taxon, and the taxa were combined for an estimate of combined mesozooplankton biomass to compare their relative biomass to phytoplankton abundance (in a common carbon unit) following White & Roman (1992, their Table 1: ‘Carbon [µg C ind\(^{-1}\)] = 0.32 W’). In addition, the nitrogen content of the zooplankton community was estimated for an overall nitrogen budget as described with the nitrogen budget below.

Unexpectedly, adult polychaetes had built burrows in the microphytobenthos mat at the end of the experiment in the R tanks. In addition, tunicates *Molgula manhattensis* were found in the R\(_{BD}\) tanks. Thus, to determine the abundance of macrofauna that may have settled or grown up over the 4 wk period of the experiment, a diaphragm pump was used to suction up the bottom half diameter of each tank after emptying the tank to a water depth of ca. 2 cm. The contents were washed through a 0.5 mm diameter mesh and the macrofauna preserved with buffered formaldehyde. *Molgula manhattensis* has been reported to filter 540 ml h\(^{-1}\) per g wet weight (Millar 1971). The wet weight of 10 tunicates was determined and averaged to result in 0.152 g ind\(^{-1}\) wet weight. The amount of water filtered per day by the tunicates was determined.

Nitrogen budget

In estimating nitrogen budgets in the R and R\(_{BD}\) tanks, water-column phytoplankton direct cell counts were converted to phytoplankton carbon after Smayda (1978) and Strathmann (1967), and the Redfield ratio of C:N of 106:16 was used to convert phytoplankton to nitrogen equivalents.

Average sediment chl \(a\), indicative of microphytobenthos abundance, was converted to nitrogen equivalents (g N per tank) after converting mg chl \(a\) m\(^{-2}\) to µg chl \(a\) m\(^{-2}\). To convert chl \(a\) biomass, a POC to chl \(a\) ratio of 51.68 was used (Porter et al. 2010). In addition, the measured carbon to nitrogen ratio of 5.138 from Porter et al. (2010) was used. The nitrogen estimate of microphytobenthos is likely an underestimate in the R tanks because microphytobenthos was probably underestimated. To estimate the nitrogen content of the zooplankton community, we determined the dry weights of the individuals of the different taxa following White & Roman (1992), Torres et al. (2002) and Sprung (1984), then the numbers were scaled up to total dry weight in the tanks using measured abundances. The nitrogen content of the mesozooplankton stock in the tanks was considered to be 7.5% of the mesozooplankton dry weight (5 to 10% of mesozooplankton dry weight is nitrogen; Parsons et al. 1984). Water-column concentrations of TDN were corrected for the daily water exchange and scaled up to the whole tank.

Statistical analyses

After Porter et al. (2010, 2013), chl \(a\), phaeophytin, total suspended solids, the dissolved inorganic and organic nutrients, and particulate nitrogen, phosphorus, and carbon were each averaged from Day 2 to Day 27 of the experiment for each tank (8 measurements over a 4 wk period) and also analyzed without Day 2 as noted below, because no biodeposits had been added to the tank yet at the Day 2 sampling. The mesozooplankton abundance was averaged for each tank from Day 3, the first time of zooplankton measurement, to Day 28. Only data from the mixing ‘on’ phases were included in statistical comparisons. Statistical t-tests were used for water column chl \(a\), phaeophytin, PN, PC, the dissolved nutrient data, the dissolved oxygen, zooplankton abundance, and the microphytobenthos accessory pigment to chl \(a\) ratio. We used linear regression analyses of mesozooplankton biomass and phytoplankton biomass (estimated from direct cell counts) to determine the relationship between the
mesozooplankton community and phytoplankton. Statistical analyses were performed using SAS (SAS Institute), and the *t*-tests using the Microsoft Excel Analysis tool pack. Significances of all analyses were defined at the $p \leq 0.05$ level. Results over the whole datasets are reported here, including sampling Day 2 before biodeposits were added, except where noted.

**RESULTS**

**Temperature, salinity, and dissolved oxygen**

Water temperatures ranged from 25.8 to 30.8°C in the R tanks (average 28.09 ± 0.11°C) and from 25.4 to 30.5°C (average 27.88 ± 0.189°C) in the R_BD tanks, and Days 19 to 23 experienced a heat wave. Dissolved oxygen concentrations were significantly lower in the R_BD tanks compared to the R tanks (Fig. 5). Salinity ranged from 11 to 12 ppt over the course of the experiment.

**TSS, particle settling, particulate and dissolved nutrients**

Total suspended solids (TSS) concentrations were significantly higher in the tanks with oyster *Crassostrea virginica* biodeposits (R_BD; ~95 mg l$^{-1}$ in Week 3; Fig. 6a) than in the resuspension tanks without added biodeposits (~15 mg l$^{-1}$; Fig. 6a, Table 1). In the morning of Day 2 of the experiment, before biodeposits were added, TSS concentrations were 13.94 ± 0.70 mg l$^{-1}$ and 13.39 ± 4.15 mg l$^{-1}$ in the R tanks and R_BD tanks, respectively, and not significantly different ($p = 0.841$). Throughout the experiment, TSS concentrations continuously increased in the R_BD tanks to ~95 mg l$^{-1}$, until they started to decrease somewhat towards the end of the experiment (Fig. 6a). During mixing ‘off’ phases, TSS concentrations were about 15 to 20 mg l$^{-1}$ in both the R_BD and the R tanks (Fig. 6a,b).

TSS concentrations were the same between mixing ‘on’ and ‘off’ phases in the R tanks (Fig. 6a,b), and particles did not settle. Bulk settling speeds changed in the R_BD tanks over the course of the experiment (Fig. 7). On Day 2, bulk settling speed was only approximately 0.07 mm s$^{-1}$ and hardly any particle settling occurred; however, between Day 3 and 13, bulk settling speeds rose to 0.44 ± 0.17 mm s$^{-1}$, and after Day 15, it greatly increased to 1.30 ± 0.09 mm s$^{-1}$ for the remainder of the experiment in the R_BD tanks (Fig. 7).

Particulate carbon (PC), nitrogen (PN), and phosphorus (PP) concentrations were linearly related to TSS concentrations: 

- PC (mg l$^{-1}$) = 0.1166 × TSS (mg l$^{-1}$) − 0.7515, $R^2 = 0.97$; 
- PN (mg l$^{-1}$) = 0.0158 × TSS (mg l$^{-1}$) − 0.0876, $R^2 = 0.97$; 
- PP (mg l$^{-1}$) = 0.0028 × TSS (mg l$^{-1}$) − 0.0225, $R^2 = 0.96$.

PC and PN concentrations were significantly enhanced in the R_BD tanks (Fig. S2a−f, Table 1). Much of the PN, PC, and PP settled out during mixing ‘off’ phases in the R_BD tanks (Fig. S2b,d,f in the Supplement).

Defining nutrient limitation as dissolved silicate <5 µmol l$^{-1}$, DIN <2 µmol l$^{-1}$, and phosphate <0.1 µmol l$^{-1}$ (Fisher et al. 1992, T. Fisher pers. comm.), all nutrients except silicate became limiting in the R tanks after Day 4 (Fig. 8c,e,g), and silicate became limiting after Day 16. In the R_BD tanks, silicate was never limiting and DIN and phosphate were limiting from Day 4 to 14. However, nutrient addition bioassays such as used by Fisher et al. (1999) would be needed to determine the limiting nutrient. Dissolved organic phosphorus concentrations were significantly higher in the R_BD tanks (Fig. 8h, Table 1).

While total nitrogen (NH$_4^+$ + NO$_2^-$ + NO$_3^-$ + DON + PN) concentrations were about the same in all tanks at the beginning of the experiment and stayed at ~40 µmol l$^{-1}$ in the R tanks for the duration of the experiment, they increased steadily in the R_BD tanks until Day 22 after which they decreased (Fig. 8f, Table 1e). Total phosphorus concentrations (PO$_4^{3-}$ + DOP + PP) were ~1.5 µmol l$^{-1}$ in all tanks at the beginning of the experiment and stayed the same in the R tanks over the experiment. In the R_BD tanks, total phosphorus concentrations increased until about Day 22 and then decreased (Fig. 8i). The ratio of TN:TP was significantly lower in the R_BD tanks than in the R tanks (Fig. 8l, Table 1). The patterns of TP and TN progressions were similar to the patterns
of the TSS concentrations over time, suggesting that resuspended biodeposits may have been a source of additional nutrients. Most of this increase in TP was due to resuspended PP, which was related linearly to TSS levels. Most of the increase in TN was due to resuspended PN, which was related linearly to TSS levels.

The switch to increased bulk settling speeds after Day 15 coincided with the start of an increase in phosphate, DIN, and nitrate + nitrite concentrations that lasted for the remainder of the experiment (Fig. 8). In addition, this increase in nutrient concentrations overcame nutrient limitation in the R_BD systems (Fig. 8) and increased as chl a concentration decreased (Fig. 6c).

Nutrient concentrations were consistent with the differences for chl a (Fig. 6c, Table 1). Nitrate plus nitrite concentrations (Fig. 8d), dissolved phosphate (PO$_4^{3-}$) concentrations (Fig. 8e), total dissolved nitrogen, and phosphorus concentrations (Fig. 8f,i) were significantly higher in the R_BD tanks than in the R tanks (Fig. 8, Table 1), and DON concentrations were similar (Table 1, Fig. 8j). As chl a increased in the R tanks (Fig. 6a), NO$_2^-$ + NO$_3^-$ (Fig. 8b), NH$_4^+$, and dissolved PO$_4^{3-}$ (Fig. 8a,e) decreased. However, midway in the experiment, chl a concentrations decreased as nutrient concentrations increased. While nutrient concentrations were related to the chl a concentrations in the R and R_BD tanks, they did not relate to phytoplankton direct cell counts.

Fig. 6. Left panels: time series of 4 variables; right panels: data from R and R_BD tanks for mixing ‘on’ and mixing ‘off’ phases averaged over 3 of the days, as indicated by ellipses in panel (a). (a) Total suspended solids (TSS), (b) chlorophyll a concentrations, (c) phaeophytin concentrations, (d) ratio of chl a:phaeophytin over the experiment. (e) TSS, (f) chl a, (g) phaeophytin and (h) ratio of chl a:phaeophytin during mixing ‘on’ and mixing ‘off’ phases. Means and SD (n = 3)
Phytoplankton, light, microphytobenthos

Water column chl a levels (10 to 30 µg l−1) were significantly higher in the R_BD tanks than in the R tanks (8–12 µg l−1) but decreased in the R_BD tanks after Day 15 to the end of the experiment (Fig. 6f, Table 1). Chl a levels decreased, in particular, in the R_BD tanks that had generated high concentrations of tunicates. Chl a did not substantially resuspend and settle in the R_BD and R tanks (Fig. 6f) as chl a concentrations were the same during the mixing ‘on’ and ‘off’ phases. Phaeophytin concentrations differed significantly between the mixing ‘on’ and the mixing ‘off’ phases as degraded material was resuspended and deposited in the R_BD tanks but not in the R tanks (Fig. 6c,g). The ratio of chl a to phaeophytin

Table 1. Summary of statistical results for the resuspension (R) and resuspension with biodeposit addition (R_BD) systems, mixing ‘on’ phases. Statistical t-tests used in all analyses. Significance was defined as p ≤ 0.05. Means ± SD, n = 3. Significant differences in bold

<table>
<thead>
<tr>
<th>(a) Seston</th>
<th>R</th>
<th>R_BD</th>
<th>p-value</th>
<th>Day</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total suspended solids (mg l−1)</td>
<td>14.63 ± 0.88</td>
<td>62.02 ± 0.65</td>
<td>0.0012</td>
<td>2−27</td>
</tr>
<tr>
<td>Particulate inorganic matter (mg l−1)</td>
<td>8.59 ± 0.74</td>
<td>40.91 ± 0.89</td>
<td>0.0018</td>
<td>2−27</td>
</tr>
<tr>
<td>Particulate organic matter (mg l−1)</td>
<td>6.11 ± 0.15</td>
<td>21.11 ± 0.26</td>
<td>0.0009</td>
<td>2−27</td>
</tr>
<tr>
<td>Ratio particulate organic matter:particulate inorganic matter</td>
<td>0.88 ± 0.17</td>
<td>0.61 ± 0.06</td>
<td>0.0994</td>
<td>2−27</td>
</tr>
<tr>
<td>Percent particulate inorganic matter</td>
<td>57.55 ± 2.35</td>
<td>63.21 ± 1.67</td>
<td>0.1561</td>
<td>2−27</td>
</tr>
<tr>
<td>(b) Chl a, phaeophytin</td>
<td>8.53 ± 0.50</td>
<td>17.73 ± 2.22</td>
<td>0.0094</td>
<td>2−27</td>
</tr>
<tr>
<td>Chlorophyll a (µg l−1)</td>
<td>3.20 ± 0.34</td>
<td>29.59 ± 1.40</td>
<td>&lt;0.001</td>
<td>2−27</td>
</tr>
<tr>
<td>Phaeophytin (µg l−1)</td>
<td>4.05 ± 0.42</td>
<td>0.60 ± 0.06</td>
<td>0.0045</td>
<td>2−27</td>
</tr>
<tr>
<td>Ratio chl a:phaeophytin</td>
<td>0.0107 ± 0.003</td>
<td>0.0277 ± 0.0062</td>
<td>0.0239</td>
<td>3−27</td>
</tr>
<tr>
<td>(c) Particulates</td>
<td>0.55 ± 0.04</td>
<td>4.94 ± 0.18</td>
<td>0.0015</td>
<td>2−27</td>
</tr>
<tr>
<td>Particulate phosphorus (µmol l−1)</td>
<td>80.12 ± 12.01</td>
<td>545.80 ± 35.06</td>
<td>0.014</td>
<td>2−27</td>
</tr>
<tr>
<td>Particulate nitrogen (µmol l−1)</td>
<td>10.12 ± 0.71</td>
<td>65.20 ± 3.65</td>
<td>&lt;0.001</td>
<td>2−27</td>
</tr>
<tr>
<td>Ratio particulate carbon:particulate nitrogen</td>
<td>8.25 ± 0.71</td>
<td>8.04 ± 0.11</td>
<td>0.8126</td>
<td>2−27</td>
</tr>
<tr>
<td>(d) Dissolved nutrients, dissolved organic carbon, dissolved oxygen</td>
<td>0.41 ± 0.07</td>
<td>0.54 ± 0.07</td>
<td>0.2108</td>
<td>2−27</td>
</tr>
<tr>
<td>Ammonium</td>
<td>1.15 ± 0.22</td>
<td>5.33 ± 2.73</td>
<td>0.0488</td>
<td>2−27</td>
</tr>
<tr>
<td>Total dissolved nitrogen (µmol l−1)</td>
<td>23.44 ± 0.72</td>
<td>28.52 ± 3.57</td>
<td>0.0214</td>
<td>2−27</td>
</tr>
<tr>
<td>Dissolved inorganic nitrogen (µmol l−1)</td>
<td>1.55 ± 0.18</td>
<td>5.89 ± 2.69</td>
<td>0.0455</td>
<td>2−27</td>
</tr>
<tr>
<td>Dissolved organic nitrogen (µmol l−1)</td>
<td>21.88 ± 0.59</td>
<td>22.63 ± 1.43</td>
<td>0.5117</td>
<td>2−27</td>
</tr>
<tr>
<td>Total dissolved phosphorus (µmol l−1)</td>
<td>0.37 ± 0.006</td>
<td>0.87 ± 0.18</td>
<td>0.0021</td>
<td>2−27</td>
</tr>
<tr>
<td>Phosphate (µmol l−1)</td>
<td>0.08 ± 0.003</td>
<td>0.45 ± 0.14</td>
<td>0.0685</td>
<td>2−27</td>
</tr>
<tr>
<td>Dissolved organic phosphorus (µmol l−1)</td>
<td>0.29 ± 0.006</td>
<td>0.42 ± 0.05</td>
<td>0.0124</td>
<td>2−27</td>
</tr>
<tr>
<td>Dissolved organic carbon (µmol l−1)</td>
<td>180.65 ± 11.85</td>
<td>181.34 ± 8.78</td>
<td>0.9587</td>
<td>2−27</td>
</tr>
<tr>
<td>Dissolved silica (µmol l−1)</td>
<td>19.76 ± 3.15</td>
<td>26.57 ± 6.87</td>
<td>0.4439</td>
<td>2−27</td>
</tr>
<tr>
<td>Ratio dissolved inorganic nitrogen:dissolved inorganic phosphate</td>
<td>17.90 ± 0.89</td>
<td>15.31 ± 0.91</td>
<td>0.6388</td>
<td>2−27</td>
</tr>
<tr>
<td>Ratio dissolved silicate:phosphate</td>
<td>244.06 ± 40.02</td>
<td>186.96 ± 44.58</td>
<td>0.5652</td>
<td>2−27</td>
</tr>
<tr>
<td>Dissolved oxygen (mg l−1)</td>
<td>7.15 ± 0.20</td>
<td>4.73 ± 0.14</td>
<td>0.0009</td>
<td>1–28</td>
</tr>
<tr>
<td>(e) Total nutrients</td>
<td>33.56 ± 1.40</td>
<td>93.72 ± 1.83</td>
<td>&lt;0.001</td>
<td>2−27</td>
</tr>
<tr>
<td>Total nitrogen (µmol l−1)</td>
<td>0.92 ± 0.04</td>
<td>5.80 ± 0.02</td>
<td>0.0016</td>
<td>2−27</td>
</tr>
<tr>
<td>Ratio total nitrogen:total phosphorus</td>
<td>36.74 ± 1.09</td>
<td>18.32 ± 0.35</td>
<td>&lt;0.0001</td>
<td>2−27</td>
</tr>
<tr>
<td>(f) Irradiance</td>
<td>154.85 ± 9.67</td>
<td>3.68 ± 2.09</td>
<td>0.0003</td>
<td>2−27</td>
</tr>
<tr>
<td>Irradiance at sediment surface (µmol photons m−2 s−1)</td>
<td>255.84 ± 2.91</td>
<td>26.94 ± 7.34</td>
<td>0.0002</td>
<td>2−27</td>
</tr>
<tr>
<td>Geometric mean irradiance (µmol photons m−2 s−1)</td>
<td>255.84 ± 2.91</td>
<td>26.94 ± 7.34</td>
<td>0.0002</td>
<td>2−27</td>
</tr>
<tr>
<td>(g) Zooplankton</td>
<td>3.70 ± 2.69</td>
<td>12.40 ± 7.17</td>
<td>0.0247</td>
<td>3–28</td>
</tr>
<tr>
<td>Adult Acartia sp. (ind. l−1)</td>
<td>3.70 ± 2.69</td>
<td>12.40 ± 7.17</td>
<td>0.0247</td>
<td>3–28</td>
</tr>
<tr>
<td>Copepodites (ind. l−1)</td>
<td>3.73 ± 1.61</td>
<td>7.24 ± 3.50</td>
<td>0.1501</td>
<td>3–28</td>
</tr>
<tr>
<td>Copepod nauplii (ind. l−1)</td>
<td>52.90 ± 11.63</td>
<td>82.21 ± 23.67</td>
<td>0.2338</td>
<td>3–28</td>
</tr>
<tr>
<td>Polychaete larvae (ind. l−1)</td>
<td>14.27 ± 9.64</td>
<td>1.91 ± 1.18</td>
<td>0.0793</td>
<td>3–28</td>
</tr>
<tr>
<td>Zooplankton carbon</td>
<td>5.11 ± 2.58</td>
<td>10.88 ± 5.22</td>
<td>0.0277</td>
<td>3–28</td>
</tr>
</tbody>
</table>
increased linearly in the R_BD tanks and was ~8-fold higher in the R_BD compared to the R tanks towards the end of the experiment (Fig. 6d,h, Table 1).

Phytoflagellates, cyanobacteria, dinoflagellates, and diatoms were present in all tanks. The tanks initially were dominated by phytoflagellates in low-moderate densities, but as the experiment progressed, diatoms became the dominant group in moderate-high densities. Many of the phytoflagellates were unidentifiable taxa but included Cryptomonas and Pyramimonas. The diatom taxa were dominated by the genus Cyclotella and included moderate numbers of Thalassionema nitzschiodes, Cylindrotheca closterium, and Skeletonema costatum. The densities of dinoflagellates were very low throughout the experiment, and there were moderate numbers of an unidentified filamentous cyanobacteria (likely Oscillatoria).

While chl a concentrations were significantly higher in the R_BD tanks than in the R tanks (Fig. 5c), phytoplankton biomass as determined by direct cell counts converted to carbon was not significantly different between tanks (Fig. 9a). In addition, dinoflagellate (Fig. 9b), diatom (Fig. 9c) cyanobacteria (Fig. 9d), and phytoflagellate (Fig. 9e) abundances were similar, although they diverged after Day 20 with phytoflagellates, cyanobacteria, and dinoflagel-
lates more abundant in the R_BD tanks. Diatoms decreased after Day 15 in the R_BD tanks until the end of the experiment (Fig. 9c), especially in the tanks where a high density of tunicates was found at the end of the experiment. Interestingly, if data from Day 2 (up to which point no biodeposits had yet been added) is excluded, the ratio of chl : carbon is significantly higher in the R_BD tanks (Fig. 9f).

Light, as measured by a home-made Secchi disk, penetrated ~30 cm into the R_BD tanks during resuspension (Fig. 10b). During the off phases, Secchi depth in the R_BD tanks reached to 60 cm from Day 5 to 12 and then increased to 90 cm, occasionally reaching the bottom during the second phase of the experiment (Fig. 10b) which coincided with an increased bulk settling speed (Fig. 7). During that time, the water column became visually clearer in the R_BD tanks shortly after mixing ‘off’ (Fig. 10b). In contrast, in the R tanks, Secchi depth reached ~90 cm into the water column from Day 2 to 12 and reached to the bottom during the remainder of the experiment. At all times, Secchi depths reached the bottom during the mixing ‘off’ phases in the R tanks (Fig. 10b).

Measured bottom irradiance levels during the resuspension phase were low in the R_BD tanks that had high TSS concentrations (Fig. 10a). Irradiance at the sediment surface with ~150 µmol m$^{-2}$ s$^{-1}$ was higher in the R than in the R_BD systems (Fig. 10a, Table 1), and geometric mean irradiance in the water column was higher in the R tanks with ~250 µmol m$^{-2}$ s$^{-1}$ than in the R_BD tanks with ~10 µmol m$^{-2}$ s$^{-1}$ (Fig. S3 in the Supplement, Table 1).

Microphytobenthos grew on the fiberglass tank bottoms of the R tanks even though bottom shear stress was high with ~0.62 Pa. Qualitatively, microphytobenthos biomass was larger in the R tanks, which had significantly more light at the bottom, with high variability amongst the R tanks (Fig. 11a). Problems with the sampling container for microphytobenthos likely led to an underestimate of microphytobenthos chl a in the R tanks and the high variability. Percent degradation was significantly higher in R_BD tanks (Fig. 11b). The ratios of Fucoxanthin:chl a, Peridini:chl a and Lutein:chl a of the microphytobenthos were significantly larger in the R_BD tanks that had lower light at the bottom (Fig. S4 in the Supplement). The levels of 19’ Butanoyloxyfucoxanthin

![Fig. 9. Phytoplankton carbon (C) determined from direct cell counts, means ± SD (n = 3) for (a) total phytoplankton, (b) dino (dinoflagellates), (c) diatoms, (d) cyano (cyanobacteria), and (e) phyllo. (phytoflagellates). (f) Ratio of chlorophyll a to total phytoplankton C.](image-url)
and Alloxanthin showed trends of being higher in the R_BD tanks, and Violaxanthin showed a trend of being higher in the R tanks \((0.1 > p > 0.05)\). None of the other pigment ratios were significantly different between the R and R_BD tanks.

**Mesozooplankton and macrofauna**

At the end of the experiment, 20 ± 5 m\(^{-2}\) adult polychaetes were found in the R tanks, some with established burrows in the microphytobenthos mat, and none in the R_BD tanks. Two R_BD tanks had 228 and 222 m\(^{-2}\) tunicates *Molgula manhattensis*, that filtered 449 and 437 l d\(^{-1}\), respectively, in the second half of the experiment, however, 1 R_BD tank had only 2 tunicates in it that filtered 4 l d\(^{-1}\). In the R tanks, 4 ± 7 m\(^{-2}\) tunicates were found that filtered 8 ± 14 l d\(^{-1}\) in the second half of the experiment. Moreover, 8 ± 7 m\(^{-2}\) small crabs were discovered in the R_BD tanks and none in the R tanks. Chl \(a\) concentrations and diatom biomass decreased greatly after Day 15 in the 2 R_BD tanks with a high density of tunicates.

Dominant mesozooplankton taxa found in the water column were adult *Acartia* sp. copepods (Fig. 12a), copepodites (Fig. 12b), copepod nauplii (Fig. 12c), and polychaete larvae (Fig. 12d). The pumps likely destroyed the adult mesozooplankton stages (Adey & Loveland 1998) during the initial raw water fill of the tanks at the start of the experiment, and it took ~1 wk until adult copepods and 16 d until polychaete larvae were found (Fig. 12a,d). The R_BD tanks contained ~15 adult *Acartia* sp. l\(^{-1}\) in the middle of the experiment and 25 at the end, while the R tanks contained 5 and 1, respectively. After 3 wk, *Acartia* sp. abundance decreased in all systems during a heat wave when copepods potentially migrated towards the bottom of the tank, but increased again on Day 24 in the R_BD tanks. Starting on Day 16 to the end, polychaete abundance was significantly \((p = 0.045)\) higher in the R tanks (40 l\(^{-1}\)) than in the R_BD tanks (4 l\(^{-1}\)) and showed a trend \((p = 0.0793)\) even when the first part of the experiment was included in the statistical analyses. *Acartia* sp. abundances were significantly higher in the R_BD system than in the R systems as was zooplankton carbon of all mesozooplankton combined. Barnacle nauplii, barnacle cyprid, crab zoea, and bivalve veliger larvae abundances were not significantly different between the R and R_BD tanks. Linear regression analysis showed no significant relation between phytoplankton biomass (from phytoplankton identifications) and zooplankton biomass (both converted to carbon units).
Nitrogen budget

Zooplankton, phytoplankton biomass, microphytobenthos, and TDN were included in the nitrogen budget percentage estimations. An additional 0.77 g nitrogen per tank was found in the biodeposits in the R_BD tanks. The data in Table 2 suggest that biodeposit addition and tidal resuspension significantly affected nitrogen partitioning within the system. The R_BD system accumulated more nitrogen with 0.57 g N per tank for zooplankton nitrogen, phytoplankton nitrogen, microphytobenthos nitrogen and TDN; there was also an additional 0.77 g biodeposit particulate nitrogen (making a total of 1.34 g N per R_BD tank as compared to the R tanks with 1.02 g N per R tank). In the R tanks, all particulate nitrogen was composed of zooplankton nitrogen and phytoplankton nitrogen, while in the R_BD tanks, 0.77 g nitrogen was due to other sources, i.e. the biodeposit additions. In the R tanks, microphytobenthos dominated the system with 51.5% of the nitrogen portion, and in the R_BD tanks, TDN dominated the system with 72.4% of the nitrogen pool of zooplankton, phytoplankton, microphytobenthos and TDN due to biodeposit addition and resuspension. Phytoplankton nitrogen was 21% in the R_BD tanks and 13.8% in the R tanks of the zooplankton, phytoplankton, microphytoplankton and TDN nitrogen pool. The zooplankton nitrogen portion was small with 2% in the R_BD tanks and 1% in the R tanks. A major nitrogen input was also due to water exchanges, and the daily water exchange inputs were identical between the R and the R_BD tanks. Some phytoplankton nitrogen was exported during the daily exchange in the R and R_BD tanks. As we did not have bottom sediments in the tanks, no nitrogen is partitioned to nutrient fluxes from the sediments and to porewater resuspension. While for the R tanks microphytobenthos dominated the nitrogen pool, in the R_BD tanks particulate nitrogen and total dissolved nitrogen dominated the nitrogen pool in addition to the particulate nitrogen in biodeposits.

Table 2. Nitrogen budget (g N per tank). All particulate nitrogen (PN) in the R tanks is zooplankton and phytoplankton. TDN: total dissolved nitrogen

<table>
<thead>
<tr>
<th></th>
<th>R (g N/tank)</th>
<th>R_BD (g N/tank)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zooplankton</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Phytoplankton</td>
<td>0.14</td>
<td>0.12</td>
</tr>
<tr>
<td>Microphytobenthos</td>
<td>0.53</td>
<td>0.03</td>
</tr>
<tr>
<td>TDN</td>
<td>0.35</td>
<td>0.41</td>
</tr>
<tr>
<td>Biodeposit PN</td>
<td>0.00</td>
<td>0.77</td>
</tr>
<tr>
<td>Total (g N per tank)</td>
<td>1.02</td>
<td>1.34</td>
</tr>
</tbody>
</table>
DISCUSSION

Oyster *Crassostrea virginica* biodeposit additions and high bottom shear stress affected TSS concentrations, particle settling, light, nutrient dynamics, oxygen concentrations, and zooplankton community structure and abundance through direct and indirect interactions. Temperatures were high in this outdoor mesocosm experiment, especially during the heat wave from Days 19 to 22, but temperatures measured in an adjacent shallow Patuxent River cove were similarly high.

It has been suggested that biodeposition may promote phytoplankton growth during periods of nitrogen limitation (Cranford et al. 2007). However, resuspended biodeposits and nutrients regenerated from it did not affect phytoplankton abundance and composition in this whole-ecosystem experiment as measured by direct cell counts. Phytoflagellates, cyanobacteria, dinoflagellates, and diatom cell counts were not significantly different between systems even though differences were found during Day 22 to the end of the experiment. Phytoplankton carbon concentrations were similar in both systems. However, chl a concentrations were significantly higher in the R_BD tanks than the R tanks, and the increase and decrease in chl a concentrations corresponded to regenerated nutrients. At the end of the experiment, 2, 228, and 222 m$^{-2}$ tunicates *Molgula manhattensis* were found in the 3 R_BD tanks, respectively, and led to decreased chl a and diatom concentrations. Tunicates are efficient filter feeders, and Comeau et al. (2015) reported that in the presence of the solitary tunicate *Ciona intestinalis* or *Styela clava* on mussel farms, the demand for phytoplankton increases by 30 to 47% compared to non-tunicate scenarios. *M. manhattensis* is an even more efficient filter feeder than *C. intestinalis* (Randløv & Riisgård 1979).

The chl a:carbon ratio was significantly higher in R_BD tanks than in the R tanks (if the first day is excluded), which suggests an adaptation of the phytoplankton to the low light regime in the R_BD tanks by increasing their chl a content in the cells (Buchanan et al. 2005). In addition, collected biodeposits contained a small amount of chl a added to the tanks daily that needed correction.

Resuspension of biodeposits increased TSS in the water column of the R_BD tanks, which was also found by Hildreth (1980) and Colden et al. (2016). In addition, biodeposit resuspension increased PN and PP levels in the water column of the R_BD tanks, reduced light penetration to the bottom, and resulted in lower microphytobenthos biomass in the R_BD tanks compared to the R tanks. While the experiment was run without a sediment bottom, microphytobenthos formed in the R tanks even though shear stress was high and polychaetes built burrow structures in it. While microphytobenthos abundance was higher in the R tanks that had more light at the bottom, a high Fucoxanthin:chl a ratio in R_BD tanks indicated that the microphytobenthos mat was adapted to low light (Cibic et al. 2007).

Oyster biodeposit resuspension affected nutrient transformations, nutrient regeneration, and the nitrogen budget. While tunicate biodeposition can be significant (Haven & Morales-Alamo 1966, McKindsey et al. 2009, Qi et al. 2015) and vertical material fluxes and nutrient regeneration from it important (Lee et al. 2012), nutrient regeneration was similar between the 3 R_BD tanks that had 2, 228, and 222 tunicates, respectively, at the end of the experiment, and tunicate biodeposition is thought to not be the driving factor here.

While nutrients did not get regenerated right away, after ~2 wk nitrate + nitrite, phosphate, and DOP increased strongly in the R_BD tanks, where desorption from particles may have played a role, whereas nutrient concentrations remained low in the R tanks. High temperatures may have mediated nutrient regeneration. Sufficient biodeposits may have accumulated by 2 wk into the experiment to see an effect. The increase in nutrient regeneration coincided with a change in bulk settling speeds in the R_BD. While particles did not settle in the R tanks, bulk settling speed increased in the R_BD tanks after Day 16, which suggests both that biodeposit particles aggregate and that, as they become bigger and denser, their net sedimentation rate increases. As feces and pseudofeces aggregate, bivalves are known to increase material sinking rates (Kautsky & Evans 1987). Tunicates are also thought to have a strong influence on vertical material fluxes (Lee et al. 2012), but all 3 R_BD tanks with 2, 222 and 228 tunicates, respectively, had similar bulk settling speeds and the tunicates are thought to not have had a dominant effect here. TSS concentrations were higher during the second half of the experiment which could have contributed to increased aggregation and settling.

The nitrogen budget partitioned differently in the R compared to the R_BD tanks. While the largest portion was partitioned in microphytobenthos (51.5%) in the R tanks, the largest portion of non-biodeposit nitrogen was partitioned in total dissolved nutrients in the R_BD tanks (72.4%). Resuspended biodeposits increased nutrient concentrations and reduced light penetration to the bottom where microphytobenthos
could form. Some of the regenerated nutrients likely came from desorption of nutrients from nutrient rich particles.

Oyster biodeposit resuspension affected ecosystem processes such as the zooplankton community composition and abundance. Zooplankton carbon concentrations were significantly higher in R_BD tanks compared to R tanks, and *Acartia* sp. were significantly more abundant in the R_BD tanks. However, polychaete larvae were more abundant in the second half of the experiment in the R tanks, and adult polychaetes had built burrows in the the microphytobenthos mat in the R tanks by the end of the experiment. Many tunicates were found in 2 of the 3 R_BD tanks. However, polychaete larvae were more abundant in the second half of the experiment in the R tanks, and adult polychaetes had built burrows in the the microphytobenthos mat in the R tanks by the end of the experiment. Many tunicates were found in 2 of the 3 R_BD tanks and decreased chl a concentrations and diatom biomass but did not affect nutrient regeneration and particle aggregation and settling significantly.

No study has evaluated the effect of continuous levels of biodeposition to simulate conditions in bivalve culture. Biodepositions are typically assumed to fall to the bottom and stay at the bottom with the sediment matrix (Newell et al. 2005, Kellogg et al. 2013). Enhanced sedimentation under shellfish aquaculture is well documented (Cranford et al. 2003). However, transport from the reef or aquaculture facilities may have far-reaching consequences for the nutrient and particle dynamics and the light regime. Chamberlain et al. (2001) concluded that variations in the dispersion of biodeposits caused by local currents affected differences in the organic enrichment of sediments and macrobenthic infaunal diversity at 2 aquaculture farms. Testa et al. (2015) concluded, using a sediment model, that it is important to consider biodeposit resuspension and transport in effectively removing oyster biodeposits from an aquaculture farm to prevent negative effects. Giles et al. (2009) found that it was important to simulate biodeposit erosion and decay where changes in the decay rate, a complex process, greatly affected fecal pellet density on the bottom as well as dispersal distances. Bivalve biodeposit erosion, settling velocity, and disaggregation rate and any effects on the ecosystem are poorly understood.

Nutrient budgets need to include biodeposit resuspension when evaluating the impact of oysters on eutrophication mitigation. For the calculation of oyster environmental benefits, the fraction of deposited material that is resuspended and the fraction of nitrogen diagenesis that undergoes subsequent denitrification must be known (Cerco 2015). Moreover, nutrient transformations in the water column from resuspended biodeposits must also be known. In our experiments, nitrate + nitrite were regenerated, which are needed for denitrification, but we did not measure denitrification. It is possible denitrification took place in the anaerobic environment of resuspended biodeposits. A current focus in the field is on the effect of biodeposits fueling biogeochemically driven nitrogen removal from sediments (Newell et al. 2002, Higgins et al. 2013, Kellogg et al. 2014); however, effects on the biogeochemistry due to biodeposit resuspension must be considered. Furthermore, it has been suggested that strategies to maximize tissue-based nitrogen removal will be particularly important where environmental conditions do not support substantial nitrogen removal through biogeochemical sediment processes (Dalrymple & Carmichael 2015), but these studies do not include the effects of biodeposit resuspension.

Impacts of biodeposit resuspension may not be immediate: it took ~2 wk for effects to appear in this experiment, and short-term experiments will not capture these effects. Thus, experiments must have long enough duration to make sure that the system is in equilibrium, including biodeposit resuspension. The experiments reported here did not include a sediment bottom and primarily focused on water column processes as mediated by biodeposit resuspension. Further changes can be expected when a muddy sediment bottom is included and thus sediment resuspension and interaction with biodeposits are included as well as sediment nutrient and gas fluxes. Existing studies of sediment resuspension and benthic processes have shown that sediment resuspension not only enhances nutrient fluxes from sediments (Qin et al. 2004, Almroth et al. 2009, Corbett 2010, Porter et al. 2010, Almroth-Rosell et al. 2012) but also significantly affects microbial and planktonic communities, and thus ultimately biogeochemical cycles (Porter et al. 2010, Wainright 1987, 1990, Isebe & Ohre 2014). Despite this limitation, this study contributed to the understanding of the effect of biodeposit addition and resuspension on the nutrient dynamics and ecosystem processes. Future studies will examine further effects of the interaction of resuspended biodeposits and sediments.

This study did not include actual oysters in the tanks. Additional well-known effects of oyster filtration include changes to phytoplankton biomass, seston concentration, and light penetration to the sediments (Porter et al. 2004b, Newell & Koch 2004). However, the focus in this experiment was on the less studied resuspended biodeposits that are difficult to track in nature (Testa et al. 2015) and cannot be resuspended in typical experimental ecosystem experiments with low bottom shear stress (Doering et al.
The STURM facility allows high bottom shear stress for the tidal resuspension of biodeposits with realistic water column turbulence levels for benthic-pelagic coupling experiments (Porter et al. 2010, 2013), so that the focus could be on biodeposits, exclusively. Addition and regular tidal resuspension of oyster biodeposits profoundly affected nutrient dynamics, nitrogen partitioning, and zooplankton dynamics in this experiment.

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