Significance of euphotic, subtidal sediments to oxygen and nutrient cycling in a temperate estuary

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ABSTRACT: Sediment-water nutrient and oxygen exchanges were measured under light and dark conditions at 1 oligohaline (Stn D) and 1 mesohaline (Stn A) shallow (1 m) site in the Neuse River estuary, North Carolina, USA. Mean respiration rates were very similar between sites (11 to 12 mg O₂ m⁻² h⁻¹), but maximum net productivity at the mesohaline site (74 mg O₂ m⁻² h⁻¹) was nearly twice that of the oligohaline site (40 mg O₂ m⁻² h⁻¹). NH₄ fluxes were also significantly different. On average, releases of NH₄ from sediments at both sites occurred in the dark (13 to 22 pmol N h⁻¹ for Stns D and A, respectively) and, slightly, at the lowest irradiance (67 E m⁻² s⁻¹, 0.3 to 5 pmol N m⁻² h⁻¹). NH₄ was taken up at average rates between 3 and 12 pmol N m⁻² h⁻¹ at higher irradiances. Mean NO₃ (nitrate + nitrite) fluxes were very low (<10 pmol N m⁻² h⁻¹), nearly always directed out of the sediment, and not significantly different. Filterable reactive phosphorus fluxes also did not follow any consistent response to changes in irradiance. Flux vs irradiance curves were used in conjunction with estimates of in situ light availability to the benthos to compute ecosystem-level influence of autotrophy on material exchanges. The autotrophic-heterotrophic transitions were 1 to 2 m (oligohaline) and 3 to 4 m (mesohaline). When integrated over depth and area the mesohaline sediments were always net NH₄ sources to the water column, net O₂ sources in fall and winter and net O₂ sinks in spring and summer. Oligohaline sediments were O₂ sinks except during fall, and NH₄ sources only in winter and summer. Cultural eutrophication has the potential to alter the balance of benthic autotrophy and heterotrophy which in turn may foster phytoplanktonic productivity.

INTRODUCTION


Our main goal was to determine the contribution of sediment-water exchanges to oxygen and nutrient cycling within the Neuse River estuary, North Carolina, USA. Due to its shallowness, light input to the benthos may act as a negative feedback control on
sediment nutrient release and subsequently on phytoplankton production. This control may be rapidly eroded by increased light attenuation. We compared heterotrophically dominated (dark-incubated) sediment-water oxygen and nutrient exchanges with autotrophically dominated (light-incubated) exchanges as functions of varying light intensity. We then calculated area- and depth-integrated fluxes for 2 salinity zones of the estuary to evaluate sediment contributions to material exchanges and the importance of euphotic sediments to those contributions.

STUDY AREA

The Neuse River estuary is a drowned river estuary beginning near New Bern, North Carolina, and emptying into Pamlico Sound 65 km downstream (Fig. 1). The estuary is subject to high rates of nutrient loading, principally from the Neuse River (Christian et al. 1989, 1991). Nutrient concentrations, particularly NO$_3$ (nitrate + nitrite), become rapidly depleted through the oligohaline zone such that nutrient concentrations at the mouth are often near detection limits (Christian et al. 1984, 1989, 1991). The zone of nutrient depletion also corresponds with the zone of maximum phytoplankton and bacterial production (Christian et al. 1984, 1991).

Water depth is greatest at the mouth of the estuary, but averages only 4.5 m there. The tide range at the head is less than 0.3 m (Giese et al. 1979), and tidal mixing is slight compared to wind mixing throughout the estuary. Stratification and hypoxia occur in deeper waters (>2 m), particularly in the oligohaline zone, but due to the shallow depths and frequent wind mixing these events are generally periodic rather than prolonged.

Study sites were located in water depths of about 1 m in both the mesohaline (Stn A) and oligohaline (Stn D) salinity zones (Fig. 1). Sediments at both stations were >95 % sand, and low in organic matter (mean total carbon ≈ 0.21 %; total nitrogen = 0.03 %). Other enumerated sites were sampled only for heterotrophically mediated exchanges as part of another study, but are shown to illustrate the areas included in the integration procedures.

METHODS

Sampling. Sampling was done at 6 wk intervals between July 1988 and July 1989 at both sites. Sediments were collected from a boat using plexiglass core tubes (3.7 cm I.D.) and a coring device. The device was a 1.25 m galvanized pipe threaded onto a one-way brass valve. PVC connectors were used to achieve the appropriate connections to the pipe and to our corers. When lowered to the sediment, water flowed through the one-way valve into the pipe. The hydrostatic pressure of the water in the pipe against the brass valve provided the seal that allowed the core to be lifted to the surface. The sediment core with the overlying water was stoppered at both ends before removal from the water, and placed in a cooler filled with water near ambient temperature. Ambient surface water was collected in Nalgene bottles and also placed in the coolers. On each sampling date, profiles of photosynthetically active radiation (PAR, \( \mu \text{E} \text{m}^{-2} \text{s}^{-1} \); LiCor Model 185A) were determined at 0.2 m intervals. Water column temperature, salinity (YSI model 33) and dissolved oxygen concentration (YSI model 57) were determined at the surface and within 10 cm of the bottom.

Sample analysis. Core incubations were begun 2 to 6 h after collection. Water samples were double-filtered through 1.5 \( \mu \)m glass-fiber filters (Whatman 934-AH), and then double-filtered through 0.45 \( \mu \)m membrane filters (Gelman GN-6). The filtering procedure reduced water column metabolism to undetectable levels (final oxygen concentrations not significantly different from initial concentrations) over the time period of the incubations so that changes in concentrations within the water were attributed to sediment metabolism. The filtered water was shaken and sampled for determination of dissolved oxygen, \( \text{NH}_4 \) (ammonium), \( \text{NO}_3 \) (principally nitrate) and FRP (filterable reactive phosphorus). The ambient water overlying each core was then withdrawn by syringe and gently replaced with the filtered water. Sediments
were exposed to the air for <1 min. Usually 2, but occasionally 4, cores were then randomly assigned to each of 5 PAR treatments: 0, 67, 107, 220, or 362 \( \mu \text{E m}^{-2} \text{s}^{-1} \). The highest PAR treatment was based on maximum light-saturation values for benthic microalgae (Colijn & van Buurt 1975, Sundbäck & Jonsson 1988). Cores to be incubated in the dark were stoppered and double-wrapped in aluminum foil. Cores for incubation in the light were sealed by covering the water surface with a layer of paraffin oil ca 5 mm thick. A strip of black rubber was wrapped around these cores at the level of the sediment-water interface and secured with a rubber band to prevent light from reaching the sides of the cores. PAR reductions were achieved by wrapping cores in neutral density screening. Light was supplied by 8 General Electric cool-white (F40CW) fluorescent lights alternating with 8 wide-spectrum (F40PL/AQ) lights. All cores were incubated in an ambient temperature water bath for 4 to 6 h. Positive fluxes represent an increase in the concentration of the overlying water during the incubation, i.e. net release from sediments. ‘Sediment respiration’ includes both aerobic respiration and chemical demand.

At the end of the incubation period the paraffin oil was removed from light-incubated cores with a 60 ml syringe. The entire volume of incubation water was then removed with a second syringe. Two subsamples were taken immediately from the large syringe with 10 ml glass syringes and fixed for dissolved oxygen determination by using the azide modification of the Winkler method scaled to the appropriate sample volume (Rizzo & Wetzel 1985). Two 12 ml subsamples were then withdrawn and fixed immediately for \( \text{NH}_4 \) determination (Solorzano 1969). The remainder of the sample was stored in a polypropylene bottle and frozen for later automated determinations of FRP and \( \text{NO}_3 \) (SJC 1981).

Stirring may (Boyn et al. 1981) or may not (Pamatmat 1977, Hall et al. 1979, Hargrave et al. 1983) affect sediment oxygen metabolism. Our cores were unstirred because we deemed potential stirring effects on respiration to be slight, while the potential overestimation of nutrient fluxes could be substantial. Stirring can stimulate aerobic respiration by removal of low oxygen tensions (Boyn et al. 1981). However, oxygen concentration changes in our experiments were small (\( \pm 2 \text{ mg l}^{-1} \)) and final oxygen concentrations were relatively high, \( \geq 2.5 \text{ mg l}^{-1} \) (vs \( \geq 3.0 \) in Boyn et al. 1981). Since oxygen tensions were never low stirring effects on respiration would probably have been minimal. However, stirring may increase solute release especially when erosion velocities are reached (Boyn et al. 1981). The potential resuspension of surface sediments or surface flocculations could result in overestimation of nutrient fluxes. Any potential errors which could have arisen from discrete sampling of possibly stratified water in the unstirred cores were avoided by sampling the entire overlying water column at the end of an experiment.

**Data analysis.** Student’s \( t \)-test (Sokal & Rohlf 1969) was used to compare hourly flux rates in dark-incubated cores with the maximum flux rate obtained from the light incubations for each material flux (e.g. respiration vs maximum net productivity, \( NP_{\text{max}} \)). Results were considered significant at \( p \leq 0.05 \).

Material fluxes which were significantly different between PAR treatments were depth- and area-integrated to evaluate effects due to euphotic sediments, from a system-wide perspective. Hourly oxygen and \( \text{NH}_4 \) fluxes were calculated for sediments at depth intervals of 0–1, 1–2, 2–3, 3–4 and >4 m by using a hyperbolic tangent model (Jassby & Platt 1976) and daily insolation data collected by the Physics Dept of East Carolina Univ. (Greenville, NC). The average daily insolation for each month of 1988 was determined from the continuous record. Light at depth was calculated from the expression \( I_z = I_0 e^{-kz} \), where \( I_0 = \text{PAR just under the surface, } I_z = \text{PAR at depth } z (\text{m}); \) and \( k \) is the mean vertical attenuation coefficient \( (\text{m}^{-1}) \), calculated from all values of \( k \) for a given month, determined between 1985 and 1989 (5 to 9 samples per month; Christian, Stanley & Boyer, East Carolina Univ., unpubl.; this study). The hourly production estimates were summed to give the total production during the photoperiod. Total nighttime fluxes were combined with the photoperiod fluxes to estimate daily rates for each depth interval. Average seasonal fluxes for the mesohaline zone [Stns A, B and C (lower half); Fig. 1] and oligohaline zone Stns C (upper half), D and E (lower half); Fig. 1) were computed by averaging daily rates for each season and depth interval and weighting by the area within each depth interval. We used the data from Stn A for the mesohaline zone and data from Stn D for the oligohaline zone. Areas of each depth interval were derived from nautical charts and sounding measurements. Daily rates for depths >4 m were calculated using only dark-incubation data (hourly rate \( \times 24 \text{ h} \)). Daily averages for each season and depth interval were multiplied by number of days per season and then summed to give annual rates.

**RESULTS**

**Material exchanges in cores**

Mean hourly rates (\( \pm \text{SD} \)) of nutrient and oxygen exchanges for treatments over the entire study are shown in Table 1. Autotrophism was significant at both
Table 1. Mean (± SD) nutrient (μmol N or P m⁻² h⁻¹) and oxygen (mg O₂ m⁻² h⁻¹) exchanges by PAR treatment (μE m⁻² s⁻¹). Positive values denote release from sediments.

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>PAR treatment</th>
<th>0</th>
<th>67</th>
<th>107</th>
<th>220</th>
<th>362</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stn A</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NH₄</td>
<td></td>
<td>22.1</td>
<td>±32.8</td>
<td>±10.9</td>
<td>±18.8</td>
<td>±5.6</td>
</tr>
<tr>
<td>NO₃</td>
<td></td>
<td>7.4</td>
<td>±4.3</td>
<td>±3.8</td>
<td>±2.3</td>
<td>±10.4</td>
</tr>
<tr>
<td>FRP</td>
<td></td>
<td>±15.0</td>
<td>±9.5</td>
<td>±9.3</td>
<td>±10.2</td>
<td>±16.7</td>
</tr>
<tr>
<td>O₂</td>
<td></td>
<td>±24.3</td>
<td>±15.1</td>
<td>±7.1</td>
<td>±19.2</td>
<td>±10.0</td>
</tr>
<tr>
<td>Stn D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NH₄</td>
<td></td>
<td>13.1</td>
<td>±64.0</td>
<td>±64.4</td>
<td>±33.0</td>
<td>±17.2</td>
</tr>
<tr>
<td>NO₃</td>
<td></td>
<td>9.1</td>
<td>±0.7</td>
<td>±4.7</td>
<td>±7.9</td>
<td>±5.7</td>
</tr>
<tr>
<td>FRP</td>
<td></td>
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<td>±39.6</td>
</tr>
<tr>
<td>O₂</td>
<td></td>
<td>±4.9</td>
<td>±11.6</td>
<td>±1.6</td>
<td>±8.2</td>
<td>±22.9</td>
</tr>
</tbody>
</table>

stations in that oxygen flux (uptake) rates in the dark were statistically different from the rates of oxygen flux (release) at NP_max. Mean NP_max was 40 mg O₂ m⁻² h⁻¹ at Stn D and 74 mg O₂ m⁻² h⁻¹ at Stn A. Maximum rates occurred during summer or early fall, and spring minimums approaching zero occurred at both sites (Fig. 2). NP_max occurred at 362 μE m⁻² s⁻¹ on 7 of 9 dates at Stn A, but occurred at 220 μE m⁻² s⁻¹ on 4 of 7 dates at Stn D.

Respiration was typically low in winter but varied much less seasonally than did NP_max (Fig. 2). Also, respiration was not consistently related to temperature. At Stn A both maximum and minimum rates occurred in summer at temperatures of 24 to 25 °C, whereas at Stn D peak respiration occurred in March (15 °C) and minimum respiration in November (18 °C).

Of the nutrients compared over the entire study, only NH₄ exchanges were significantly different between dark and light treatments. Mean hourly rates of NH₄ release in the dark were <25 μmol m⁻² h⁻¹ at both stations. The mean rates were greatly affected by very high releases in late summer, particularly at Stn D (Fig. 3), where NH₄ was often taken up in the dark. At both sites mean NH₄ release was negligible at 67 μE m⁻² s⁻¹ and uptake occurred at greater irradiances. Maximum NH₄ flux in the light was always negative (uptake) at Stn A, and on all but 2 occasions at Stn D (Fig. 3). In most samples (63 % for Stn A and 57 % for Stn D) maximum NH₄ uptake occurred at irradiances lower than those giving NP_max. This was especially pronounced at Stn D, where maximum NH₄ uptake in April and November occurred at 67 μE m⁻² s⁻¹ while NP_max occurred at 362 μE m⁻² s⁻¹.

Mean FRP exchanges were also low (≤25 μmol P m⁻² h⁻¹) at both stations. FRP was released from the sediments under all light treatments at Stn A. At Stn D, FRP was taken up in the dark, and means representing both uptake and release occurred under positive PAR treatments. On a few occasions at both stations, FRP release in the light was significantly lower than dark release rates, and decreased with increasing irradiance in a hyperbolic pattern similar to the NH₄ fluxes. FRP releases at Stn A were consistently much lower in the light than in the dark. At this site, (considering all PAR treatments) light-incubation release averaged only 40% of dark release (range 15 to 61 %).

There were no significant differences in mean NO₃ exchanges resulting from PAR treatment at either site. Mean exchanges were very low (≤10 μmol m⁻² h⁻¹) and generally directed out of the sediments.
**Integrated material fluxes**

Daily and annual integrated exchanges of oxygen are given in Table 2, and NH$_4^+$ fluxes in Table 3. Over the entire salinity zone, mesohaline sediments were net daily (24 h) sources of oxygen to the water column during fall and winter. Oligohaline zone sediments were net sources in the fall. Mesohaline sediments ≤2 m deep were net oxygen sources year round, and were sources to 3 m depth in fall and winter. In the oligohaline zone, the source-sink transition occurred at much shallower depths: 1 to 2 m in summer and fall and <1 m in winter and spring.

The daily ratios of gross production to respiration (GP:R, where GP = NP + R) for the first 4 depth intervals indicated substantially greater autotrophy in the mesohaline zone than the oligohaline zone for each depth (Table 2B). As a result, mesohaline sediments consumed only 33% of the total estuarine benthic oxygen consumption, though representing 70% of the total area.

All mesohaline sediments were year-round NH$_4^+$ sources except during spring for depths ≤2.0 m (Table 3B). The dominance of release at the systems level resulted from the high fluxes in dark incubations relative to the smaller fluxes of opposite direction in the light (Table 1). Nevertheless, autotrophic metabolism in the shallow sediments had a substantial impact on NH$_4^+$ exchange. Euphotic sediments (<4 m deep) comprise 56% of the total area but contributed only 22% of the total annual NH$_4^+$ release in the mesohaline zone (Table 3B). Seasonally, most of the total annual NH$_4^+$ release (76%) took place in summer. Summer fluxes also contributed most of the total annual NH$_4^+$ flux (63%) in the oligohaline zone, disregarding flux direction. But, in contrast to the mesohaline zone, sediments were sinks at all depths during both spring and fall (Table 3B), reflecting NH$_4^+$ uptake in dark-incubated cores.

**DISCUSSION**

**Influence of light on material exchanges**

The GP:R ratio has long been used as an indicator of the trophic status of ecosystems (Odum & Wilson 1962). Values >1 indicate autotrophic dominance of metabolism and values <1 indicate heterotrophic dominance. There are strong longitudinal and vertical variations.

<table>
<thead>
<tr>
<th>Site</th>
<th>Depth (m)</th>
<th>Area (km$^2$)</th>
<th>Oxygen flux (mg O$_2$ m$^{-2}$)</th>
<th>Oxygen flux (t O$_2$ zone$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Winter</td>
<td>Spring</td>
<td>Summer</td>
</tr>
<tr>
<td>M</td>
<td>0-1</td>
<td>48</td>
<td>434</td>
<td>388</td>
</tr>
<tr>
<td></td>
<td>1-2</td>
<td>29</td>
<td>180</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>2-3</td>
<td>27</td>
<td>8</td>
<td>-35</td>
</tr>
<tr>
<td></td>
<td>3-4</td>
<td>45</td>
<td>-57</td>
<td>-81</td>
</tr>
<tr>
<td></td>
<td>&gt;4</td>
<td>117</td>
<td>-182</td>
<td>-275</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>265</td>
<td>-15.2</td>
<td>-18.9</td>
</tr>
</tbody>
</table>

| O    | 0-1       | 22     | -58    | -43    | 87    | 165    | -1.3   | -0.9   | 0.9   | 3.6   |
|      | 1-2       | 19     | -157   | -176   | -79   | -5     | -3.1   | -3.4   | -1.5  | -0.1  |
|      | 2-3       | 26     | -163   | -200   | -91   | -30    | -4.3   | -5.3   | -2.4  | -0.8  |
|      | 3-4       | 33     | -163   | -202   | -92   | -33    | -4.3   | -6.3   | -2.8  | -0.8  |
|      | >4        | 14     | -394   | -488   | -222  | -80    | -5.4   | -6.7   | -3.1  | -1.1  |
|      | Total     | 115    | -18.4  | -22.6  | -8.9  | 0.8    |        |        |       |

Table 2. Integrated rates of oxygen flux by depth strata and season for the mesohaline (M) and oligohaline (O) zones of the Neuse River estuary. (A) Daily rates; (B) annual rates, and mean ratios of daily gross production to daily respiration (GP:R). Fluxes for areas >4 m deep are calculated from dark-incubation rates per 24 h. Positive values denote release from the sediment.
changes in trophic status within the Neuse River estuary. $GP:R$ declines from the mouth to the head of the estuary and, as expected, declines rapidly with depth.

To facilitate comparison with other studies we converted our data (from Table 2) to carbon units using a photosynthetic quotient of 1.0 (Pomeroy 1959). Annual $GP$ in the sediments for the first 4 depth intervals of Stn A were 110, 64, 37 and 26 g C m$^{-1}$ yr$^{-1}$. At Stn D, the annual rates were 43 g C m$^{-1}$ between 0 and 1 m, and between 24 and 26 g C m$^{-2}·$yr$^{-1}$ for 1 to 4 m depths. Other estimates of annual benthic microalgal production from subtidal areas <2 m deep range from 75 to 253 g C m$^{-2}·$yr$^{-1}$ for 1 to 4 m depths. Other estimates of annual benthic microalgal production from subtidal areas <2 m deep range from 75 to 253 g C m$^{-2}·$yr$^{-1}$ (Marshall et al. 1971, Davis & McIntire 1983, Nowicki & Nixon 1985a, Rizzo & Wetzel 1986, Wasmund 1986, Murray & Wetzel 1987, Fielding et al. 1988). Thus annual rates at Stn A span the lower limit of reported values while rates at Stn D are lower at all depths.

The difference in $GP$ and $GP:R$ between Stns A and D must arise from differences in $NP_{\text{max}}$ since respiration rates were nearly identical for these stations. Lower $NP_{\text{max}}$ at Stn D most probably results from reduced light penetration. Nutrient limitation would be unlikely since nutrient concentrations at Stn D are much higher. The onset of light saturation $I_{\text{L}}$ (after Talling 1957) was similar at both sites: 297 ± 78 µE m$^{-2}·$s$^{-1}$ at Stn A and 274 ± 88 µE m$^{-2}·$s$^{-1}$ at Stn D (Lackey 1992). Over the photoperiod at 1 m depth mean PAR was 220 µE m$^{-2}·$s$^{-1}$ at Stn A (74 % of mean $I_{\text{L}}$) but only 68 µE m$^{-2}·$s$^{-1}$ at Stn D (25 % of mean $I_{\text{L}}$). The larger difference between average in situ PAR and light-saturating PAR at Stn D indicates a much higher potential for light-limitation.

In estuarine sediments, NH$_4$ exchanges have been invariably affected by benthic microalgal metabolism during incubation in the light. NH$_4$ release has either been greatly reduced compared to dark-incubation rates (Nowlcki & Nixon 1985b, Rizzo 1990) or flux direction has been reversed, with sediments becoming NH$_4$ sinks (Sundbäck & Granéli 1988, Nilsson et al. 1991, Sundbäck et al. 1991). The earlier study by Rizzo (1990) for a site in the York River estuary of Chesapeake Bay (USA) is of particular interest. Except for the tide range (0.8 m), his site is extremely similar to Neuse River Stn A regarding depth, sediment type, sediment organic concentrations, light penetration, annual gross productivity and annual ranges of temperature, salinity, and water column nutrient concentrations (Rizzo 1986, 1990, Rizzo & Wetzel 1986). In situ NH$_4$ release by York River sediments in the light was
75% less than within dark domes, but uptake in the light occurred only once - on the date when minimal sediment respiration rate occurred during the study. In contrast, cores from Stn A virtually always took up NH₄ even during incubations at low light intensities and low NP. The difference in effect of light incubation on NH₄ flux between these 2 very similar systems probably results from differences in trophic state arising from different rates of sediment respiration. Mean respiration rate at the York River site was more than 3-fold higher than at the Neuse River site. This greater heterotrophic activity is reflected in differences in mean GP:R between these sites. Mean GP for the York River site was 1.6 compared to 3.2 for Stn A (on a daily basis). Also, values <1 were recorded on most sample dates for the York River site but were never found at Stn A. Heterotrophic activity at the York River site apparently supplied the nitrogen requirements of the benthic microalgal community resulting in low but consistent NH₄ release during light incubations. Lower respiration rates at Stn A apparently did not satisfy microalgal requirements which were met by removal of NH₄ from the water column. Since sediment NP occurs in surface micro-layers only about 1 mm thick (Revsbech & Jorgensen 1983) micro-zones of nutrient limitation may occur in sediments which otherwise show high concentrations of NH₄ in coarse (e.g. cm section) vertical profiles. Benthic microalgae may be more easily released from nutrient limitation by removing NH₄ from the water column rather than by diffusion from deeper sediments.

At Stn D rates of NH₄ release might be expected to be greater than at Stn A since sediment respiration is the same while autotrophic demand is less. However, the frequent dark-NH₄ uptake at Stn D suggests a substantial involvement of other pathways in sediment-water NH₄ exchange at this site. Dark uptake of NH₄ at Stn D could arise from the following mechanisms: (1) diffusive fluxes may occasionally dominate exchanges during periods of low respiration and high concentrations in the water column, and/or (2) dark uptake of NH₄ could result from either heterotrophic or chemoautotrophic metabolism. Heterotrophs using organic matter with a relatively high C:N ratio may take up NH₄ whereas nitrifiers can consume NH₄ oxidatively to generate energy.

The lack of impact of light-incubation on rates of NO₃ exchange is probably due to relatively high NH₄ concentrations and a presumed preference of the microbiota for NH₄. We would expect that NH₄ is preferred by benthic microalgae as it is for phytoplankton (McCarthy et al. 1977), although there is presently no experimental evidence for this. If this is the case, then little autotrophic impact on NO₃ fluxes would be expected at Stn A since NH₄ concentrations exceed NO₃ concentrations (Christian et al. 1991). At Stn D, NO₃ concentrations are typically much higher than NH₄ concentrations but concentrations of both nutrients are much greater than at Stn A (Christian et al. 1991). Since NP at Stn D is much lower than Stn A, there is less reason to suspect greater NH₄ limitation of primary production or autotrophic impact on NO₃ exchanges. In general, sediment-water exchanges of NO₃ in areas with benthic microalgal communities are low (<20 µmol m⁻² h⁻¹) and erratic in direction (Nowicki & Nixon 1985b, Sundbäck & Graneli 1988, Rizzo 1990, Sundbäck et al. 1991), although in sandy sediments light incubations have resulted in limiting NO₃ release or NO₃ uptake compared to dark releases (Nowicki & Nixon 1985b, Sundbäck et al. 1991).

The results for NH₄ fluxes are not surprising since nitrogen limitation is common in marine systems. However, freshwater ecosystems, particularly lakes, often have phosphorus-limited primary production (Schindler 1977). In freshwater systems benthic microalgae typically reduce phosphorus releases (Carlton & Wetzel 1988, Kelderman et al. 1988, Forsberg 1989, Hansson 1989) in a manner similar to our results for NH₄ fluxes. Several estuarine and coastal researchers also report benthic microalgal effects on phosphorus fluxes (Sundbäck & Graneli 1988, Nilsson et al. 1991, Sundbäck et al. 1991) although Rizzo (1990) found no impacts of light-incubation for the York River sand shoal. In the Neuse River light-incubation FRP releases are much reduced compared to dark releases, albeit without statistical significance, but on individual dates FRP can be affected by benthic microalgal metabolism in a manner similar to effects on NH₄. FRP may intermittently limit or co-limit benthic microalgal production in the Neuse, requiring uptake from the water column. However, phytoplankton production in the lower Neuse River estuary is generally nitrogen limited rather than phosphorus limited (Paerl et al. 1990, Mallin et al. 1991). Because oxic sediments are effective phosphorus traps, phosphorus limitation of benthic microalgal production would seem even less likely than for phytoplankton, but the availability of sediment-bound phosphorus to microalgae is unknown. Sporadic FRP limitation of benthic microalgal production remains a possibility. Also, other mechanisms, some indirectly linked to NP such as redox potential changes, may be significant controls on FRP flux rates, possibly masking biological effects.

**Sediment contributions to system oxygen and nutrient cycles**

As in any extrapolation, embedded scales of spatial-temporal variability should be borne in mind when considering the results. Extrapolations of hourly rates,
determined monthly, is common practice in calculating annual rates of both phytoplankton and benthic microalgal productivity. Nevertheless, such averaging may be affected by embedded temporal variability. For instance, benthic microalgal productivity can vary significantly over diel, day-to-day, tidal and seasonal scales (Rizzo & Wetzel 1986). A relatively few measurements will produce an annual mean in general agreement with the literature (Rizzo & Wetzel 1985), but the essentially random selection of sampling dates and incubation periods can result in very different apparent seasonality (Rizzo & Wetzel 1986). While our data are replicated within seasons, except for winter, and not subject to tidal influences, other scales of variability (e.g. diel or day-to-day) may have affected the results.

Extrapolation of data from limited spatial coverage to large areas is also frequent and useful in providing different perspectives (Baird & Ulanowicz 1989, Seitzinger 1991). Nevertheless, extrapolation of spatial scales also poses difficulties. Although we found no significant longitudinal differences in sediment respiration for 5 sandy shallow areas (A to D in Fig. 1), the associated nutrient fluxes appear to differ among stations (Rizzo & Christian unpubl., this study). Preliminary analyses indicate potentially significant differences in heterotrophically dominated exchanges associated with macrofauna, sediment type and depth.

In contrast to the view of sediments as oxygen sinks, sediments in the Neuse River estuary often function as net sources of oxygen. During seasons when these sediments did function as oxygen sinks uptake was only 19 to 21% (mesohaline) and 34 to 53% (oligohaline) of the uptake expected had all sediment metabolism been heterotrophically dominated (i.e. based only on the dark incubations). In the mesohaline zone net daily oxygen production exceeds consumption over 29% of the total benthic area, and the percentage increases to 39% in fall and winter. In contrast oligohaline sediments at most depths are oxygen sinks. Very shallow areas (<1 m) are sinks in winter and spring, and net daily oxygen production during the most active seasons (summer and fall) involves only 19% of the total area. Regions of autotrophic sediments can help maintain oxygen conditions within the water column and may offer important refuges to animals during the ephemeral warm-season hypoxic or anoxic events characteristic of the Neuse River estuary (Hobbie & Smith 1975).

Mesohaline sediments are net year-round NH₄ sources for the water column while oligohaline sediments provide a net release of NH₄ only during winter and summer. Aphotic sediments, 41 to 44% of the total area, dominate total annual fluxes in each zone (78% of the total NH₄ release for the mesohaline zone; 66% of total flux regardless of direction for the oligohaline zone).

Sediment release (internal loading) of NH₄ to the water column is much less than external dissolved inorganic nitrogen (DIN) loading. Annual external loadings of DIN from tributaries and major point sources for 2 yr spanning March 1985 to February 1987 (Christian et al. 1989) ranged from 111 to 196 Mmol (20 to 26 Mmol as NH₄). Total system internal loading is only 30 Mmol yr⁻¹, comprising only 13 to 21% of the total DIN input to the water column. If based strictly on dark-incubation data, internal loading would nearly double (59 Mmol).

Based on average annual phytoplankton production for the lower estuary (343 g C m⁻²; Mallin et al. 1991) and Redfield ratios (Redfield et al. 1963), the phytoplankton nitrogen requirement is ca 4 to 5 mol N m⁻² yr⁻¹. Benthic NH₄ release in this zone (0.09 mol N m⁻² yr⁻¹) thus supplied only about 2% of this demand. Since measured NH₄ uptake is about twice the calculated phytoplankton demand (Boyer et al. 1988), the benthic contribution to total integrated water column NH₄ uptake is negligible. Most of the phytoplankton nitrogen requirement appears to be supplied by pelagic recycling, a view supported by the results of network analyses of static multicomartment models of nitrogen cycling in the Neuse River estuary (Christian et al. in press). These analyses also showed little sediment contribution to planktonic nitrogen requirements. However, they also showed a greater potential for contributions if more organic nitrogen was made available to the sediments.

**Influence of light attenuation**

Light penetration to the sediments is obviously an important control on material exchange. In the Neuse River estuary light attenuation varies considerably over short-term and seasonal scales, especially in the oligohaline zone. There may be both natural and direct and indirect anthropogenic causes for the reduction in water transparency in the oligohaline zone. Oligohaline zones are classically associated with sediment trapping and turbidity maxima (Postma 1967). This is also true for the Neuse River estuary oligohaline zone (Wells & Kim 1991). However, discharge of dissolved organic matter from a paper pulp mill effluent (between Stns G and F, Fig. 1) contributes to increased oligohaline water-column light absorption. Also, direct loading of both particulate and dissolved material to this zone may have increased in recent years for the Neuse River estuary...
as the basin has undergone cultural eutrophication (Stanley 1988).

The increased dissolved inorganic nutrient loadings to the estuary may indirectly reduce light transparency by causing the oligohaline maximum that occurs in phytoplankton biomass in the Neuse River (Christian et al. 1991). In turn, reduced light penetration to the benthos may create a positive feedback situation whereby light reduction results in increased sediment inputs of NH₄ to the water column. Continued eutrophication may radically change the character of the benthos of the Neuse River estuary from one in which substantial in situ primary productivity supports a grazed food web and oxic conditions, to one similar to that proposed by Webb (1981) in which the benthos is increasing reliant on sedimentation and subject to anoxia from increased oxygen consumption (due both to decreased autotrophy and increased heterotrophy).

Since the estuarine benthos is often primarily aphytonic (Nixon & Pinson 1983), many studies have considered material cycling only from the standpoint of heterotrophic metabolism (Nixon et al. 1976, Boynton et al. 1980, Fisher et al. 1982, Boynton & Kemp 1985). However, in 2 large East Coast estuaries, the York River estuary (Rizzo 1990) and the Neuse River estuary, 30 to 40 % of the benthos supports some benthic microalgal metabolism. Based on bathymetry we propose a similar role for benthic microalgal communities in other tributary estuaries of Chesapeake Bay and Albemarle-Pamlico Sound assuming similar light attenuation characteristics. Benthic microalgal communities may also be quantitatively important within both Chesapeake Bay and Albemarle-Pamlico Sound, as suggested by high rates of benthic microalgal production within Chesapeake Bay seagrass beds (Murray & Wetzel 1987), and the overall shallowness and distance from riverine turbidity inputs of Albemarle-Pamlico Sound. Benthic microalgal communities may be important in other estuaries as well. Nevertheless, the ecological role of these communities and their response to anthropogenic impacts remain poorly understood.

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