Reference No. [UNCEES]CBL 90-107

Sediment-water oxygen and nutrient exchanges along a depth gradient in the Baltic Sea

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ABSTRACT: Bottom water and sediment characteristics and net sediment-water fluxes of oxygen, nitrogen, phosphorus and hydrogen sulphide were measured under oxic and experimentally induced anoxic conditions along a depth gradient (47 to 130 m) during summer in the Baltic proper. Temperature, salinity and dissolved nutrient concentrations (particularly phosphorus and nitrate) in bottom waters increased with depth while oxygen concentrations decreased sharply. Sediment organic content was much higher in sediments located beneath the permanent pychocline (ca 65 m). Sediments at all stations were somewhat depleted in total N relative to total C (C:N \approx 10). Sediments at the shallow station (47 m) were highly enriched with total P relative to C or N (C:N:P \approx 18:2:1) but were somewhat depleted in P at the deeper stations (C:N:P \approx 120:12:1). Under oxic conditions oxygen fluxes ranged from 214 to 777 μ mol O m⁻² h⁻¹ and decreased with depth. Phosphorus and nitrite fluxes were always very small and were directed either into or out of the sediments. Ammonium fluxes were small (1 to 30 μ mol N m⁻² h⁻¹) at all stations and did not exhibit a clear pattern with depth; O:N-NH₄ flux ratios were close to expected 'Redfield ratios' at the deep station (16:1) but were much higher (>55:1) at the shallower sites indicating that processes other than only ammonification were taking place in sediments. Nitrate was always consumed by sediments (ca 1 to 16 μ mol N m⁻² h⁻¹) and fluxes were proportional to nitrate concentrations in the overlying water. Under anoxic conditions there was a dramatic increase in P-PO₄ fluxes (2 to 40 μ mol P m⁻² h⁻¹) and a smaller increase in N-NH₄ fluxes (14 to 35 μ mol N m⁻² h⁻¹). Large hydrogen sulphide fluxes (>40 μ mol S m⁻² h⁻¹) were observed in sediments from the deepest station only. Under oxic conditions sediment recycling of N and particularly of P were small compared to estimated rates of burial of P and burial plus denitrification of N. However, under anoxic conditions, sediment recycling of both N and P were similar to or much greater than sediment loss terms of burial and denitrification. Sediment regeneration under oxic conditions could supply 1 to 8 % and 0 to 2 % of estimated phytoplankton demand of N and P. respectively; under anoxic conditions 12 % of N and up to 200 % of P demand could be met via sediment recycling.

INTRODUCTION

In many shallow coastal and estuarine systems, sediment nutrient and oxygen dynamics have been found to exert an important influence on conditions in overlying waters and sediments (Zeitzschel 1980, Nixon & Pilson 1983). Under various circumstances, sediments have been found to be an important temporary source of some compounds, a terminal sink for others and an interface in the ecosystem where a diversity of biogeochemical transformations occur (e.g. Blackburn & Henriksen 1983). In many cases the magnitude of these sediment-water processes is sufficient to be a major component in oxygen or nutrient budget calculations and hence also of importance in formulating estuarine management strategies. Concurrent with these observations, there has been an increased interest in understanding the regulation of these processes. Several environmental factors have emerged as often being significant, including the quantity and quality of organic matter reaching the sediment surface (Wassmann 1984, Jensen et al. 1988), infaunal community characteristics (Blackburn & Henriksen 1983) and the chemical status of surficial sediments and overlying waters (Henriksen & Kemp 1988). The interactive nature of such variables, however, often makes it difficult to identify cause-effect relationships.

We undertook a study of sediment-water nutrient

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0171-8630/90/0063/0065/\$ 03.00

Contribution No. 2139, Center for Environmental and Estuarine Studies of the University of Maryland.

exchanges in the central region of the Baltic Sea, for which there is a good record of traditional water quality measurements which suggest that the system has been undergoing cultural eutrophication for the past 100 yr (Larsson et al. 1985). Nutrient budget calculations (Larsson et al. 1985, Rönner 1985) and recent numerical simulation model results (Stigebrandt & Wulff 1987) suggest that sediments in this system play an important role in both nutrient and oxygen dynamics and hence have a particular relevance to the eutrophication issue. However, there are no direct measurements of sediment-water exchange from the Baltic proper from which to judge the accuracy of overall nutrient budget calculations and to calibrate and verify simulation modelling results. Some direct measurements exist from the shallow (<20 m) Kiel Bight (Ruhmor et al. 1987) and the Bothnian Bay (Nedwell et al. 1983).

In this study we address the question of how sediment-water fluxes vary along a depth gradient in the Baltic proper and assess potential environmental controls on these fluxes. Specifically, we measured ambient sediment-water fluxes from 3 distinctive depth zones in the northern Baltic proper using intact sediment cores collected about 3 mo after the spring bloom period. In addition, we subjected these cores to experimentally induced anoxic conditions which occur periodically at each of these depth zones.

SAMPLING LOCATIONS

Three sampling locations were chosen in the northern Baltic proper, each representing different environmental conditions common in the Baltic (Table 1). The

Table 1. Bottom water and surface sediment (0 to 2 mm and 0 to 20 mm) characteristics at the 3 sampling stations along the depth gradient in the Baltic Proper. Numbers in parentheses are from cores sampled at the end of the experiment

Measure		Station	
	1	2	3
Latitude	N 58° 40′ 88″	N 58° 31′ 19″	N 58° 29′ 73″
Longitude	E 17° 50' 38"	E 18° 02' 60"	E 18° 06' 77"
Depth (m)	47	82	130
Bottom water			
Temp. (°C)	4.1	4.8	4.9
Salinity (units)	6.9	8.2	9.4
Oxygen (mg l^{-1})	13.7	6.3	1.8
Nitrate (μ mol 1^{-1})	1.9	8.9	10.5
Nitrite (μ mol l ⁻¹)	0.2	0.3	<0.1
Ammonia (μ mol l ⁻¹)	0.6	0.5	0.8
Phosphate (μ mol l ⁻¹)	0.9	2.7	3.7
N:P (by atoms) ^a	3.0	3.6	3.1
Sediment (0-2 mm)			
Total carbon (%)	3.6 (1.4)	10.2 (7.8)	7.1 (7.5)
Total nitrogen (%)	0.35 (0.16)	1.31 (0.93)	0.83 (0.88)
C:N (by atoms)	12.0 (10.2)	9.1 (9.8)	10.0 (10.0)
Sediment (0-20 mm)			
Sediment H ₂ O (%)	28.6	81.1	91.1
Bulk density (g cm^{-3})	1.75	1.11	1.05
Total carbon (%)	0.53 (0.63)	5.93 (5.88)	6.74 (7.27)
Total C (g m^{-2})	133	249	126
Total nitrogen (%)	0.076 (0.076)	0.73 (0.72)	0.79 (0.84)
Total N (g m^{-2})	19	31	15
Total phosphorus (%)	0.076	0.150	0.130
Total P (g m ⁻²)	19	6	2
C:N:P (by atoms)	18:2:1	103:11:1	134:13:1
C:N (by atoms)	8.2 (9.8)	9.6 (9.6)	10.1 (10.1)
Sediment Eh (mV)			
Overlying water	425	420	370
Sediment surface	434	430	380
2 mm	384	346	254
10 mm	271	232	-18
20 mm	224	178	-84
200 mV at depth (mm)	26	13	2.5
^a Calculated as $(N-NO_2 + N-NO_3)$	+ N-NH ₄)/(P-PO ₄)		

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first station was at 47 m water depth which is above the permanent halocline (65 m), rarely experiences severe hypoxia, has a well-established macrofaunal community and relatively low organic matter content (Ankar & Elmgren 1976). The second station was in 82 m of water and may be subjected to seasonal hypoxia and anoxia. The infaunal community here is less well developed, sediments have a higher organic matter content and tend to have some degree of lamination. The third station was at 130 m, a depth at which hypoxic and anoxic conditions commonly occur, and the macrofaunal community is poorly developed or absent for much of the year; sediment organic matter content is high and sediments are clearly laminated to depths of 4 to 5 cm. Depths below 40 m represent about 70 % of the total Baltic area but only 5 % lies deeper than 150 m (Stigebrandt & Wulff 1987). Temperatures at all stations were similar in early July (4.5 \pm 0.4 °C). The stations were also outside the influence of the coastal archipelago zone so that primary production, nutrient conditions in euphotic waters and direct deposition of organic material to the sediments could be assumed to be similar among stations. Major differences among stations were largely related to differences in the sediment characteristics and those of the overlying waters.

MATERIALS AND METHODS

At each station 50 l of near-bottom water was collected for later use in the laboratory measurements. Four intact sediment cores were obtained using a modified Kajak corer (Kajak et al. 1965, Blomqvist & Abrahamsson 1985) equipped with an acrylic liner (8 cm diameter, 50 cm length, 2.5 mm wall thickness). In most cases the cores penetrated the sediment to a depth of about 20 cm, the remaining core volume being filled with overlying water. Immediately after sampling, 4 of the cores were placed in a darkened, insulated chamber at ambient water temperature for transport to the laboratory. A vertical Eh-profile was taken on one core from each station using a platinum electrode slowly inserted into the sediment with a micromanipulator. A surficial sediment sample (top 2 mm) was taken and triplicate samples analysed for particulate carbon (PC) and nitrogen (PN) content using an elemental analyser (Perkin-Elmer Model 240 B). Particulate phosphorous (PP) content was measured using persulphate oxidation followed by standard phosphate procedures (Parsons et al. 1984). The top 2 cm of sediments were also analysed for percent water, PC, PN and PP. These measurements were repeated on all cores at the end of the measurement period. Results from these analyses varied by less than 10 % in all cases.

Three cores from each station were brought to the laboratory and maintained in the dark at 5 \pm 1°C for the duration of the experimental period (33 d). Larger animals, mainly amphipods, were removed and each core was fitted with water-tight plastic bottom and top stoppers, the latter with ports for sampling and addition of new water. A gentle circulation was maintained using magnetic stirring bars suspended within the cores. Water samples were withdrawn from the cores every 72 h and analysed for dissolved oxygen, phosphate, ammonium, nitrite, nitrate, and hydrogen sulphide following standard analytical methods (Parsons et al. 1984). Following each sampling, water was added to each core from reservoirs containing near-bottom water from the appropriate station. Net sediment-water fluxes were then calculated based on concentration differences between successive measurements, corrected for concentration changes due to the replacement water added after each sampling period and the surface-volume characteristics of each core. Concentration changes associated with the overlying water measured in control cores without sediments were alwavs small.

During the first 3 flux measurements (9 d) the sediment surface and overlying water in each core was maintained in an oxidised condition by the addition of oxygen-saturated water following each measurement period. Prior to making the fourth flux measurement about 75 % of the overlying water was removed and replaced with water from the appropriate reservoir which had been bubbled with N2 gas to remove dissolved oxygen. After 3 d sampling was again resumed under anoxic conditions. All water added to the cores until Day 27 was stripped of oxygen prior to being used. We refer to this second phase of the experiment as the 'anoxic' phase although, after addition of new water, there were always small amounts of oxygen and nitrate present in the overlying water. This was always metabolised during the following incubation and the sediments themselves were anoxic, as evidenced by the production of hydrogen sulphide.

RESULTS AND DISCUSSION

Characteristics of bottom waters and sediments along the depth gradient

Measurements of bottom water and sediment characteristics from the 3 sampling stations are shown in Table 1. Bottom water temperature and salinity ranged from 4.1 to 4.9 °C and 6.9 to 9.4 units respectively, increasing slightly from the shallow to the deep station in both cases. Dissolved oxygen concentrations decreased sharply with depth from 13.7 mg l^{-1} at Stn 1 to 1.8 mg l⁻¹ at Stn 3. Nitrate was the dominant form of dissolved inorganic nitrogen at all stations, ranging from 1.9 μ mol N l⁻¹ at Stn 1 to 10.5 μ mol N l⁻¹ at Stn 3. Dissolved inorganic phosphorus concentrations also increased sharply along the depth gradient (0.9 to 3.7 μ mol P l⁻¹). The N:P ratio of dissolved inorganic N and P compounds ((N-NH₄ + N-NO₃ + N-NO₂)/P-PO₄) in bottom waters was well below that expected based on simple oxidation of organic matter (16:1), indicating a preferential loss of N relative to P.

Redox conditions in surface sediments (Table 1) at Stn 1 were relatively oxidised (redox > 200 mV) down to about 25 mm depth, whereas sediments from beneath the halocline (Stns 2 and 3) had shallower redoxclines, at 13 and 3 mm, respectively. This is reflected in the physical appearances of the cores: the shallow cores had uniform light brown sediments with little evidence of black sulphide layers or zones. Cores from Stn 2 were light brown to gray in color with small zones of black sediment. The deep cores had distinct striations with layers 0.5 to 1.0 mm thick alternating between light gray and black down to at least 4 cm (Fig. 1). The surface of these cores had a growth of white filamentous bacteria, probably sulphide-oxidising *Beggiatoa*.

There was a distinct gradient in total carbon content of the top 2 cm of the sediments with a range of 0.5 to 6.7 % from the shallowest to the deepest station. Nitrogen content was similar at Stns 2 and 3 but an order of magnitude lower at Stn 1 (Table 1). C:N ratios showed an increase with depth, probably reflecting a change in the nature of the organic matter, with more refractory material in the deeper surface sediments.

Sediment characteristics data from the cores analysed after termination of the experiment showed little deviation from those at the beginning for most of the measurements. One notable exception was the change in both carbon and nitrogen content in the top 2 mm at the shallow station. Here total carbon content dropped by some 60 % and total nitrogen by 55 % during the 1 mo study resulting in a drop of the C:N ratio from 12.0 to 10.2. There is no evidence, however, that cores from the shallow station were substrate-limited towards the end of the experiment because fluxes remained about equal during the whole study period (Table 2).

Fluxes of oxygen, nitrogen, phosphorus and sulphur along the depth gradient

Nutrient fluxes (mean \pm SD of 3 replicate cores) recorded during the 27 d experiment are shown in Table 2, where positive values represent a loss from the sediment (i.e. a flux into the water) and negative values a flux from the overlying water into the sediment.



Station 1



Station 2



Fig. 1. Appearance of cores from 3 depths in the northern Baltic proper. Stn 1, 47 m: sediments appeared to be well mixed and light brown to well below 20 mm. Stn 2, 82 m: sediments were gray with some black zones apparent. Stn 3, 130 m: sediments were obviously undisturbed by bioturbating organisms, laminations alternating between gray and black were obvious down to 40 or 50 mm

Oxygen fluxes were always from the water into sediments but are reported as positive values to follow general conventions. The first 3 values (Days 3 through 9) were calculated during the oxic phase, the following 5 during the anoxic phase of the experiment (Days 15 through 27).

Oxygen fluxes

There were clear patterns of sediment oxygen consumption (SOC) along the depth gradient during the oxic phase of the experiment. Stn 1 consistently had the highest oxygen consumption rates; fluxes at Stn 2 were generally between 76 and 84 % of those recorded from the shallowest station whereas those from the deep

Koop et al.: Sediment-water fluxes in the Baltic

Table 2. Fluxes of oxygen and inorganic nutrients at 3 stations along a depth gradient in the Baltic proper measured under summer oxic and anoxic conditions in incubated sediment cores. All numbers are means of 3 measurements ± standard deviation (SD). Where n < 3 no SD is given. Positive numbers indicate fluxes out of the sediment and negative ones fluxes into the sediment; oxygen was always consumed by sediments and is given as a positive value to follow convention. -: no data available

Stn	Flux		Oxic period Day				Anoxic period Day		
		3	6	9	15	18	21	24	27
1 2 3	Oxygen ^a (μM O m ⁻² h ⁻¹) Sulphide ^b (μM S m ⁻² h ⁻¹)	777.1 ± 93.4 635.4 ± 29.7 214.9 ± 47.6	574.2 ± 57.0 479.8 ± 92.3 378.4 ± 122.0	789.9 ± 55.0 602.8 ± 265.0 -	2.3 ± 1.3 2.3 ± 0.2 49.5 ± 18.1	0.0 ± 0.0 0.0 ± 0.0 3.4 ± 6.7	1.6 ± 0.2 1.8 ± 2.1 43.1 ± 5.1	8.1 ± 5.5 12.8 ± 14.0 647.6 ± 187.0	5.4 ± 4.7 11.6 ± 13.5 79.9 ± 13.9
1 2 3	Phosphate (μ M P m ⁻² h ⁻¹)	-0.9 ± 0.6 1.4 ± 0.6 0.9 ± 0.7	-0.5 ± 0.3 -1.7 ± 0.9 0.1 ± 2.6	0.0 ± 0.1 0.5 ± 2.7 -0.3 ± 3.4	9.2 ± 7.4 20.0 2.1	19.0 ± 7.6 37.0 2.7	23.0 ± 10.0 36.9 ± 4.2 4.8 ± 0.8	$\begin{array}{c} 13.7 \pm 6.4 \\ 21.8 \pm 3.0 \\ 2.9 \pm 0.2 \end{array}$	15.4 ± 6.7 19.4 ± 6.4 7.3 ± 0.6
1 2 3	Ammonium (μ M N m ⁻² h ⁻¹)	$\begin{array}{c} 11.1 \pm 3.7 \\ 3.7 \pm 1.4 \\ 10.7 \pm 3.5 \end{array}$	$\begin{array}{c} 10.4 \pm 5.3 \\ 0.7 \pm 0.8 \\ 28.9 \pm 12.4 \end{array}$	17.3 ± 3.9 1.4 ± 2.3 14.0	24.0 ± 10.6 26.2 ± 7.0 26.9 ± 5.1	$\begin{array}{c} 22.7 \pm 10.5 \\ 35.5 \\ 28.5 \pm 5.5 \end{array}$	$\begin{array}{c} 13.9 \pm 6.3 \\ 28.0 \pm 5.7 \\ 30.5 \pm 3.5 \end{array}$	17.9 ± 3.6 31.0 ± 2.4 34.4 ± 5.9	$\begin{array}{c} 14.1 \pm 2.8 \\ 18.8 \pm 0.6 \\ 25.2 \pm 2.3 \end{array}$
1 2 3	Nitrate (μ M N m ⁻² h ⁻¹)	-8.0 ± 6.8 -13.9 ± 4.0 -16.2 ± 0.9	-3.7 ± 4.6 -5.0 ± 3.7 -11.7 ± 1.8	-0.5 ± 0.2 -7.3 ± 2.6 -7.6 ± 2.7	-1.3 ± 0.7 -15.4 ± 1.3 -23.8 ± 3.2	-1.4 ± 0.2 -7.7 ± 1.0 -2.7 ± 0.1	-0.7 ± 0.5 -6.2 ± 0.4 -6.7 ± 0.1	-1.5 ± 0.2 -8.4 ± 0.0 -8.9 ± 0.1	-0.7 ± 0.5 -10.8 ± 0.0 -11.4 ± 0.1
1 2 3	Nitrite (μ M N m ⁻² h ⁻¹)	-0.1 ± 0.2 -0.2 ± 0.1 -0.1 ± 0.2	-0.0 ± 0.0 -0.4 ± 0.3 -0.1 ± 0.4	-0.2 ± 0.0 0.2 ± 0.7 0.9 ± 1.0	-0.2 ± 0.1 -0.1 ± 0.6 -0.6 ± 0.1	0.3 ± 0.4 0.8 ± 1.6 -0.4 ± 0.1	-0.1 ± 0.2 -1.5 ± 0.8 -1.3 ± 0.2	0.2 ± 0.2 0.4 ± 0.1 0.1 ± 0.4	$\begin{array}{c} -0.4 \pm 0.1 \\ 0.7 \pm 0.6 \\ -0.6 \pm 0.0 \end{array}$
a Oxyge ^b Sulph	en fluxes Days 3 through 9 ide fluxes Days 15 through 27								

station were much lower, only 30 to 66 % of those from Stn 1. During the anoxic phase, any oxygen added to the cores after sampling was always consumed during the 3 d incubation.

SOC values, although low when compared to measurements from other parts of the world, and particularly from highly productive ecosystems (Seiki et al. 1989), are within the range of measurements recorded from other parts of the Baltic Sea and the west coast of Sweden and Norway which characteristically lie between 500 and 1000 μ mol O m⁻² h⁻¹ (Table 3). Low SOC rates are directly related to the low temperatures which prevail in the Baltic Sea for most of the year, particularly in the deeper water, and to the supply of organic matter to sediments which is also low in our area. Primary production is about $150 \text{ g C} \text{ m}^{-2} \text{ yr}^{-1}$ (Elmgren 1984) of which 25 to 30 % reaches the bottom, by which time it has also been partially degraded in the water (Table 1). Our estimates of SOC fit the empirical relationship between primary production plus organic input and benthic carbon mineralisation developed by Nixon & Pilson (1983). Their relation indicates that a total annual organic input equal to 150 g C M⁻² yr⁻¹ yields a benthic carbon mineralisation rate of 1070 μ mol O m⁻² h⁻¹, close to our measured values.

Sulphide fluxes

Environmentally significant sulphide fluxes were found only at Stn 3 during the anoxic phase of the experiment and these were quite variable (Table 2). Fluxes at Stn 3 were of the same order of magnitude as those recorded by Tuttle in portions of Chesapeake Bay (pers. comm.). Production of sulphide during anoxic conditions is respiration based on the use of S-SO₄ as a terminal electron acceptor, and we can calculate an oxygen-based respiration rate equivalent to this production of sulphide assuming that sulphate is directly reduced to sulphide and that no other geochemical processes are involved (e.g. production of solid phase S compounds). Our measured sulphide fluxes are equivalent to an SOC of between 14 and 2590 μ mol O m⁻² h^{-1} , with a mean value of 659 µmol $m^{-2} h^{-1}$ at the 130 m station. These values are about equivalent to or higher than the SOC rates we recorded under oxic conditions.

The sulphide fluxes we observed on Day 18 were uncharacteristically low at all stations; we attribute this to a possible error in the measurements. Also, the reading for the deep station on Day 24 was much higher than any other measurement. This may be due to the fact that bottom waters at Stn 3 are frequently subjected to anoxic conditions and there may have been a build-up of H_2S beneath the surficial sediments. What we measured on that day may have been a pulse of sulphide which had diffused to the surface from deeper layers. Thus, the high flux may not represent metabolic processes occurring at the time of measurement. If Day 18 and 24 data are omitted (Table 2) mean sulphide fluxes during the anoxic period at Stn 3 amounted to 58 μ mol S m⁻² h⁻¹ which is equivalent to a respiration rate of 230 μ mol O m⁻² h⁻¹, similar to the respiration rates measured under oxic conditions.

The 2 shallower stations had much lower fluxes of sulphide, equivalent to means of 14 and 23 μ mol O m⁻² h^{-1} at Stns 1 and 2, respectively. Fluxes increased slowly during the experimental period. Several explanations are possible. First, there could have been oxygen leaks in all the cores from Stns 1 and 2. This would have resulted in erroneously low fluxes as we suspected happened on Day 18. However, this seems unlikely because cores from all stations were treated alike and coefficients of variation were similar among all stations. A more likely explanation for these low fluxes is that a large fraction of the S-SO4 reduced went to solid phase sulphide (e.g. FeS and FeS₂) rather than H₂S. Tuttle et al. (pers. comm.) recently found that about 70 % of the S-SO4 reduced became monosulphides and pyrite in bioturbated, iron-rich sediments in Chesapeake Bay (USA). If processes of similar magnitude occurred at the 2 shallow stations in our experiments then much higher sulphide fluxes, equivalent to between 32 and 53 μ umol O m⁻² h⁻¹ can be calculated for these sediments.

Phosphate fluxes

Phosphate fluxes showed distinct patterns along the depth gradient. Fluxes during the oxic phase of the experiment were consistently low at all stations (Table 2) and were either positive or negative. During the anoxic phase phosphate fluxes were always positive, between 10 and 40 μ mol P m⁻² h⁻¹ at Stns 1 and 2, but much lower (2 to 7 μ mol P m⁻² h⁻¹) at the deep station.

A plot of phosphate fluxes versus ambient oxygen concentration in the overlying water is shown in Fig. 2. Oxygen concentrations above 1 g O_2 m⁻³ led to small and variable phosphate fluxes. Below this concentration fluxes were generally positive and much higher. In contrast to most other fluxes measured in this study, phosphate fluxes recorded from our 3 stations are in the same range as those reported from other productive ecosystems (Seiki et al. 1989). Whereas carbon, oxygen and nitrogen transformations are controlled largely by biological processes, those of phosphorus are driven mostly by redox-dependent chemical reactions (Krom & Berner 1980, Klump & Martens 1981). Thus, these fluxes are likely to be very similar over a wide range of habitats if redox conditions favour phosphorus release

Location	Sampling Scheme	Sedime Type	ent characteristi % Org. C	¢	Depth (m)	Temp. (°C)	0	Fluxes (μN N-NH₄	[m ⁻² h ⁻¹) N-NO ₃ + N-NO ₂	P-PO4	Comments	Source
Fanafjorden, Norway	Annual	Silt/sand	1-35	54-73	6090	1-5	442963	I	1	I	Max. rates in May–Jun following spring bloom	Wassmann (1984)
Kiel Bight, W Germany	Annual	Silt/sand	1.0–2.25 ^b (0.45)	37	20	2-18	341–855 (661) ^c	5.6–27.8 (18.3) ^c (110) ^d	5.6-10.0	0.7–2.6 (2.2) ^c (31) ^d	Occasional periods of anoxia encountered du- ring summer	Balzer (1984)
Baltic proper	Annual	1	1	1	65	5	1	18	1	I	Calculated from nutrient budget model	Wulff & Stige- brandt (1989)
Gulf of Bothnia	Apr/Jul	Soft silt	Detritus layer present in summer	L.	22	0-15	1000–2400	0.2–20.0 (40.0) ^d	2.8–8.5	1	NH4 and O fluxes greatly enhanced by presence of or- ganic detritus on sediment surface and tempature, respectively	Nedwell et al. (1983)
Öresund, S Sweden	Annual	Sandy mud	í .	1	27	3-10	(1005) ^c	I	I	Ì	Oxygen flux cor- related with temperature, well-developed macrofauna	Kanneworff & Christensen (1986)
Laholm Bay, W Sweden	Sep	Silt	3.4	80	20	6	1563	50 (200) ^d	0 to -5 (-20 to -30)	0 to -5 (0 10) ^d	Organic-rich sediments	Enoksson (1987)
Gullmar Fjord, W Sweden	Sep	Sand/silt	1–3	t	30	10	1644	-4 to 55 (24) ^e	-8-11 (2) ^e	ī		Enoksson & Samuelsson (1987)
Gullmar Fjord, W Sweden	Ĩ	Fine sand	ĺ	80	9	1	508	11.7	8.3	ſ		Anderson et al (1984)
Norsminde Fjord, Denmark	Annual	Silt	0.5-6.0	4084	1	2-20	583-3167	I	ł	l .	Oxygen and sul- phate fluxes cor- related with temperature	Jørgensen & Sørensen (1985)

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Location	Sampling Scheme	Sedimen Type	t characteristic % Org. C	ъsа Ф	Depth (m)	Temp. (°C)	0	Fluxes (µN N-NH4	1 m ⁻² h ⁻¹) N-NO ₃ + N-NO ₂	p-PO4	Comments	Source
ASKÖ transect, Baltic proper	Jul	Sand/silt	0.5	71	47	4.1	714 (8.7) ^f	13 (18) ^f	-2.2 (-1.0) ^f	-0.5 (12.6) ^f	Sediments well mixed	This study
partice proper	Jul	Silt	5.9	81	82	4.8	573 (≈0) ^f	4 (24) ^f	-8.7 (-9.0) ^f	0.1 (21.7) ^f	Sediments partially mixed	This study
	Jul	Silt	6.7	91	130	4.9	433 (33) ^f	$(29)^{f}$	(-14.3)	$(3.7)^{f}$	Clear striations present	This study
^a % Organic C au ^b Range in the su	nd ¢ represei uface 2 mm, 1	nt percent organ Entry in parenth	uic carbon and I eses is average	porosity. Ior sedi	respecti iments be	vely, in the dow 3 cm	upper 2 to	10 cm of the	sediment colu	иш	·	
^c Annual mean v ^d Flux recorded c	alue Juring anoxic	: period in summ	ler									
^e Average of rest	ults from 11 c	ores collected in	September									

^f Fluxes recorded during hypoxic/anoxic periods





and there is a usable pool of phosphorus available. The substantially lower phosphorus fluxes we observed at the deep station may reflect a depleted pool of available phosphorus because of the chronic exposure of these sediments to hypoxic or anoxic conditions.

Ammonium fluxes

Ammonium fluxes ranged from 1 to 35 μ mol N m⁻² h⁻¹ and agree with values recorded from elsewhere in the Baltic and the west coast of Sweden and Norway (Table 3). They are, however, at the low end of the range reported from other ecosystems (Seiki et al. 1989). Mean fluxes at Stns 1 and 3 were similar during the oxic phase (ca 15 μ mol N m⁻² h⁻¹) whereas Stn 2 had values about 6 to 7 times lower. During the anoxic phase fluxes increased by factors of 15 and 2 at Stns 2 and 3 respectively, while fluxes at Stn 1 were of equal magnitude throughout the 27 d measurement period.

Nitrite and nitrate

Nitrite fluxes were always extremely small and were directed both into and out of sediment (Table 2).

Nitrate fluxes ranged between 0.5 and $24 \ \mu mol \ N \ m^{-2}$ h^{-1} and were directed into sediments during the entire study period. During the anoxic phase any nitrate added after sampling was used up entirely during the following 3 d incubation period and hence fluxes measured during this period probably underestimate actual rates. Measured fluxes were within the range of values reported in other studies from northern Europe (see Table 3) although, as with oxygen and ammonium, they are low compared to results obtained in other, more productive, habitats. Nitrate entering sediments from the overlying water can be processed along 2 different pathways. First, it can be reduced to ammonium and enter the dissolved ammonium pool in the interstitial water and ultimately cross the sedimentwater interface to overlying waters. Alternatively, it can enter the denitrification pathway and be reduced to gaseous N_2 which then leaves the system. As we have described above, ammonium fluxes in our study were generally lower than we would predict from direct biodegradation of organic matter to N-NH₄. Because N-NH₄ fluxes were low it seems likely that a proportion of the nitrate entering sediments was denitrified.

During the measurement period nitrate fluxes were directly proportional to nitrate concentrations in overlying waters. When all nitrate fluxes were plotted against the N-NO₃ concentration at the start of each 3 d incubation period a linear relationship of the form $y = -1.78 \times -0.18$ (r² = 0.66; n = 62) was obtained (Fig. 3).



Fig. 3. Relationship between nitrate fluxes and nitrate concentrations in the overlying water in sediment cores from 3 depths in the northern Baltic proper. Points from both the oxic and anoxic phase of the experiment are plotted y = -1.78x - 0.18($r^2 = 0.66$, n = 62)

Nutrient flux ratios

During aerobic decomposition of organic matter derived from phytoplankton, approximately 13.25 oxygen atoms are consumed for every atom of nitrogen released as ammonium (Redfield 1934). It is useful to examine our measured sediment-water fluxes for departures from such stoichiometric relationships and to relate these to other possible transformations in sediments. Mean O:N flux ratios during the oxic phase of our experiment (using oxygen consumption and N-NH₄ fluxes) were 57, 529 and 13 for Stns 1, 2 and 3, respectively. Only at Stn 3 do calculated ratios agree with the expected Redfield ratio, indicating that organic material is being decomposed directly to N-NH₄. At Stn 1, and particularly at Stn 2, ammonium fluxes are much lower than expected, indicating that other pathways must be active. One clear possibility is that the ammonium produced in the sediments is oxidised to nitrate and then denitrified. Also, nitrate fluxes were always directed into the sediments and this nitrate may have also been denitrified. This pattern of anomalously high O:N ratios has been observed in other areas (e.g. Chesapeake Bay) under similar conditions (Boynton & Kemp 1985). The expected mean ammonium production based on our measured oxygen consumption values is about 50 µmol N m⁻² h⁻¹ for Stns 1 and 2 combined. This contrasts with an actual measured mean flux of about 7 μ mol N m⁻² h⁻¹, leaving 43 μ mol N m⁻² h⁻¹ unaccounted for. If we assume that this is denitrified and that the nitrate entering sediments from the water column (ca $6 \mu mol N m^{-2} h^{-1}$) is also denitrified we calculate a total denitrification rate of about 49 μ mol N m⁻² h⁻¹. This is in agreement with other calculated and measured denitrification rates from similar areas (Table 4).

Flux ratios of N:P (N-NH₄:P-PO₄) observed during the anoxic period of our experiment also depart markedly from expected Redfield ratios (1.3, 1.1 and 9.0 for Stns 1 to 3, respectively). These very low ratios resulted from extremely high P-PO4 fluxes observed under anoxic conditions which exhibited a relatively larger increase than fluxes of N-NHa under similar conditions. Since the organic source for these sediments is mainly phytoplankton detritus with an initial N:P composition ratio of about 12:1 (Leppänen 1988) it is probable that the extreme departures from stoichiometric ratios that we observed at Stns 1 and 2 were of a transient nature resulting from the rapid release of recently deposited P as well as P sequestered in oxidised surficial sediments via mechanisms such as those suggested by Klump & Martens (1981) and Carman & Wulff (1989). Interestingly, sediments at the deepest station had N:P flux ratios closer to those expected based on the composition of the source material. Given the general anoxic conditions in overlying waters at this station, P fluxes may be based solely on release from recently deposited material rather than a combination of both recently deposited and sequestered P compounds. It may be worth investigating how long very high P-PO₄ fluxes relative to N fluxes can prevail because such information is useful in developing nutrient management strategies.

Relationships between sediment characteristics and nutrient fluxes

The C:N:P ratio of organic material is often used to characterise particulates, both with regard to organic

Location	Sampling	Sedime	nt characteri	stics	Depth	Temp	Denitrification	Source
	scheme	Туре	% Org. C	Ф	(m)	(°C)	$(\mu M Nm^{-2} h^{-1})$	
Laholm Bay, W Sweden	Sept	Silt	3.4	80	20	10	19–24	Enoksson (1987) ^d
Aarhus Bight, Denmark	Annual	Silt/clay	2.6-11.4	41-75	15	4-13	4-21 (42) ^b	Jensen et al. (1988) ^e
Baltic Sea proper	Annual	-	~		0-450	4	16	Wulff & Stige- brandt (1989) ^f
Lendrup Vig, N Denmark	Spring Summer	Silt	1.5	-	<1	7–23	13-213	Andersen et al. (1984) ^e
Norsminde Fjord, Denmark	Annual	Silt (mud flat)	5–15	 ,	0.5	0-20	83–416 (205) ^c	Jørgensen & Sørensen (1988) ^e
Himmerfjärden, SE Sweden	Annual	Silt		-	30	1.5-10.0	1.5-24.0	Koop & John- stone (unpubl.)
Laholm Bay, W Sweden	-	Silt		.— '	17–20	10	10	Sahlsten & Sörensson (1986) ^e
Baltic proper deep water	Annual	-		-	>60	4	74	Shaffer & Rönner (1984) ^g
Baltic proper	Annual	-	_	-	55	4	14	Rönner (1985) ^e
^a % Org. C and Φ re	epresent perce	ent organic carb	on and poro	sity, respe	ctively, in	surface sed	iments	

Table 4. Environmental conditions and denitrification rates from the Baltic Sea and adjacent marine areas. -: no data available

^b Maximum observed rate

^c Mean annual rate

^d Estimated from ammonium fluxes

^e Acetylene inhibition technique

^f Estimated from nutrient budget calculations

matter quality and origin. Phytoplankton detritus characteristically has C:N ratios of around 7 and the ratio gets higher as the material becomes more refractory, with organic matter of terrestrial origin having values of 30 or more. The C:N ratios of surficial sediments along our transect were around 10 indicating that some of the nitrogen in the sedimenting phytoplankton material had been lost. Superimposed on this general pattern there was a depth gradient of C:N ratios with the shallowest station having a value of 8.2 (top 20 mm) increasing to 9.6 at the intermediate station and to 10.1 in the deepest sediments. This indicates a successive depletion of organic matter in N relative to C along the depth gradient. One explanation for this pattern is that material deposited in shallow water is likely to be resuspended frequently and, in the process, transported to deeper areas where it accumulates. During transport much of the labile nitrogen is likely to be rapidly recycled (Garber 1984) leaving only the more refractory material to be deposited in deeper areas. Such an explanation is consistent with the fact that organic matter content of sediments increases substantially with depth. An alternative explanation, which does not require invoking transport, is that the potential for denitrification of deposited

N increases with depth. This hypothesis could be tested by measuring denitrification along the depth gradient.

Particulate C:N:P ratios of surface sediment (top 20 mm) also exhibited a wide range along the depth gradient (Table 1). The measured value of 18:2:1 at the 47 m station indicates that this material was highly enriched in phosphorus relative to both C and N. This is in sharp contrast to sediments at the deeper Stns 2 and 3 which were both poorer in phosphorus with ratios of 103:11:1 and 134:13:1, respectively, quite close to the Redfield ratio of 106:16:1. High phosphate fluxes after the redox shift at Stn 1 and, particularly, Stn 2 suggest that there is a large pool of mobilisable phosphate in these sediments which is released when redox conditions permit. At Stn 3 fluxes during the anoxic phase of the experiment were much lower than at the other stations. Sediments at this station experience periodic anoxia in the bottom water during which the labile phosphorus fraction may be released into the water as during the anoxic phase of our experiment. Under such conditions most of the labile phosphorus is likely to have been removed from those sediments even though the total phosphorus content may still be rather high. Thus the P in apatite-phosphorus, for example, is very stable and is unaffected by ambient changes in oxygen

75

and pH. This appears to be the situation in the sediment at Stn 3 where fluxes during the anoxic phase of our experiment were only about 20 % of those recorded from Stn 1 and an order of magnitude less than those from Stn 2.

Influence of sediment processes on water column characteristics and nutrient budgets

The potential influence of sediment processes on nutrient budgets and water column characteristics can be evaluated by comparing the relative magnitude of appropriate terms. Thus we can compare the magnitude of release rates of N and P from sediments with phytoplankton demand for these nutrients as well as with the amounts of N and P lost through burial in deep sediments and that of N lost through denitrification. We are aware of the pitfalls of doing such comparisons using results from only one experiment. Obviously additional data are required before a more complete picture can be obtained. The following comparisons may, however, be useful as a first approximation.

In the Baltic proper annual rates of phytoplankton production have been estimated to be about 150 g C m^{-2} yr⁻¹ (Elmgren 1984, Leppänen & Kononen 1988). Since we did not have direct measurements of N and P uptake (e.g. Caperon et al. 1979) we used a Redfield composition ratio of 106:16:1 coupled with primary production rates to develop estimates of daily phytoplankton nutrient demand (Redfield 1934, Boynton & Kemp 1985). This approach yields demand estimates of about 216 and 14 µmol m⁻² h⁻¹ for N and P respectively. Comparing these values to the net sediment release rates of N-NH₄ and P-PO₄ during the oxic period (Table 2) we find that sediments supplied a small fraction of the required N (1 to 8 %) and essentially no phosphorus (0 to 2 %). If we consider the net flux of N-NH₄ plus N-NO₃ across the sediment-water interface, the amount of N potentially available to phytoplankton is even further reduced (Table 2) because nitrate was consistently consumed by the sediments.

In sharp contrast to these findings it appears that sediment regeneration of N-NH₄ and P-PO₄ during anoxic periods can supply significant proportions of the phytoplankton demand, particularly of P, at the 2 shallower stations. Specifically, ammonium fluxes could supply 9, 13 and 13 % of estimated N demands at Stns 1 to 3, respectively. During the same period phosphate fluxes could supply 119, 200, and 30 % of P demand, respectively. The large P fluxes observed at the 2 shallower stations during induced hypoxia are particularly significant in terms of current eutrophication trends in the Baltic. As shown by Carman & Wulff (1989) these sediments have the highest capacity for adsorbing phosphorus and also the highest concentrations of phosphorus calculated per unit area. From our present study it appears that they have a large capacity for releasing P under low oxygen conditions. Such conditions occur only occasionally at present (Larsson et al. 1985) but if they became more frequent, they would, all other factors being equal, have the effect of enriching bottom waters with P and further reducing already low N:P ratios of dissolved nutrients in deep waters (Table 1).

A second approach in evaluating the significance of our measured sediment release rates is to compare them with sediment related loss terms of burial and denitrification. We have assembled available estimates of burial and denitrification rates from the Baltic and adjacent areas (Tables 4 and 5). Burial rates of N and P ranged from 5 to 24 μ mol N m⁻² h⁻¹ and 0.5 to 0.8 μ mol

Location	Depth (m)	Estimated (µM m	burial rate ⁻² h ⁻¹)	Source			
		N	Р				
Kiel Bight	20	~5	~0.5	Balzer (1984) ^a			
Baltic proper	>60	~10	-	Shaffer (1987) ^b			
Baltic proper	0-450	22	0.5	Wulff & Stigebrandt (1989) ^b (note these values are estimates for burial + denitrification)			
N Baltic proper	150	~24	-	Leppänen (1988) ^a			
Fanafjorden, W Norway	60-90	8	0.8	Wassmann (1984) ^a			
ASKÖ transect	47	4.7	0.7	This study ^a (assuming vertical accretion rate of			
	82	29.4	0.9	$10 \text{ mm m}^{-2} \text{ yr}^{-1}$			
	130	15.1	0.3				

Table 5. Estimates of burial rates of N and P from the Baltic Sea and adjacent areas. -: data not available

^a Estimates based on vertical accretion rate of sediments, porosity and N and P content of sediments

^b Estimate based on nutrient budget calculations

P m⁻² h⁻¹, and were similar to those calculated from our study (Askö transect). These rates of burial were similar or slightly higher than net fluxes of N during the oxic period of our experiment and very similar during the anoxic period. There was an order of magnitude shift in the relative rates of burial and sediment release between oxic and anoxic periods with releases of P dominating during the latter period. A very considerable range in denitrification rates has been reported (1.5 to 416 µmol N m⁻² h⁻¹) as indicated in Table 4. However, if rates from very shallow, organically rich areas which also have an external supply of nitrate, are eliminated, the majority of values are in the range 2 to 24 µmol N m⁻² h⁻¹, again similar to rates of NH₄⁺ flux which we observed.

While there are some uncertainties associated with these types of comparisons, the relative uniformity of burial and denitrification rates leads us to believe that we are not making order of magnitude errors. The major conclusions we draw from these data is that sediment recycling of N and P is more than balanced by the loss terms of burial and denitrification, especially under oxic conditions. Should anoxic conditions become more prevalent in time and space, however, we would expect sediment recycling of nitrogen and phosphorus to assume increasing importance because of the loss of the denitrification pathway (which would result from lower levels of NO_3^- and the release of P previously bound in the sediments. Both of these changes would exacerbate eutrophication tendencies.

There are a number of measurements that need to be made before a more complete understanding of the role of sediment processes in the Baltic proper can be obtained. First, there are virtually no direct measurements of processes such as sediment recycling of nitrogen and phosphorus, burial or denitrification for zones of the Baltic deeper than 20 to 30 m. This may be an especially serious shortfall because our data suggest strong gradients related to depth and the mean depth of the Baltic is about 55 m. Secondly, there is the obvious need to obtain measurements of these processes over at least one annual cycle, particularly during and immediately after settling of the spring bloom when others have observed substantially increased rates of many sediment processes (e.g. Enoksson & Rüdén-Berg 1983, Wassmann 1984, Jensen et al. 1988). Finally, in the vast majority of sediment-water flux studies, measurements of N fluxes have included N-NH₄, N-NO₃ and N-NO₂ but not dissolved organic nitrogen (DON), partly because early attempts failed to find interpretable fluxes (e.g. Boynton et al. 1980). Enoksson (1987), however, has more recently measured significant DON fluxes in Laholm Bay, on the Swedish west coast. It seems that the old issue of sediment DON fluxes will have to be revisited if we are

to evaluate adequately total N processes at the sediment-water interface.

Acknowledgements. We thank Anders Sjösten and the chemistry laboratory of the Askö Lab. for doing some of the analyses and Lisa Matteson for preparing the figures. We appreciate the technical assistance received from Berndt Abrahamsson. Sven Blomqvist lent us some of his equipment, for which we are grateful. Discussions with Jon Tuttle and Mike Kemp were particularly useful. Financial assistance was provided through grants to K. K. and F. W. from the Swedish Environmental Protection Board. We thank B. O. Jansson for making available the facilities at the Askö field station.

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Manuscript first received: August 7, 1989 Revised version accepted: February 23, 1990