Effects of short-term anoxia on benthic denitrification, nutrient fluxes and phosphorus forms in coastal Baltic sediment

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ABSTRACT: Whether sediments act as sinks or sources of nutrients depends partly on the oxygen conditions at the seafloor. Laboratory experiments on coastal Gulf of Finland (Baltic Sea) sediment tested the sensitivity of denitrification to a 2 wk anoxia exposure and subsequent reoxidation of the bottom waters. At the same time we followed the rapidly (1 d) and more slowly (9 d) emerging changes in different forms of sediment P after oxic conditions were restored. The total denitrification rate (Dtôt) did not change during anoxic incubation, but shifted from coupled nitrification-denitrification (Dn) towards water column nitrate dependence (Dw). As the Dn rate did not decrease at the same rate as the Dw rate increased, the overall effect of 2 wk exposure to anoxia was an increase in Dtôt rate. Nitrification was enhanced in the manipulated sediment compared to natural conditions, despite anoxia. Anoxia quickly caused a release of dissolved P from its 2 most labile forms. The effect was readily reversible, but in nature the replenishment of oxygen stores is usually linked to an intense mixing of the water column, and it is possible that part of the P released during anoxia reaches the productive layer. In our experiments, anoxia affected P cycling more than N cycling.

KEY WORDS: Denitrification · Phosphorus fractions · Benthic nutrients · Anoxia

INTRODUCTION

Eutrophication is one of the main environmental problems in the Baltic Sea. In the most eutrophied part of the sea, the Gulf of Finland, nutrient loading exceeds average loading to the Baltic Sea by a factor of 2 to 3 (Pitkänen et al. 2001). In addition to loading, internal nutrient processing, such as nitrogen (N) removal (by denitrification and anammox) and phosphorus (P) binding to and release from the sediments, define the nutrient status of a water body. Whether sediments act as sinks or sources of nutrients depends partly on the oxygen (O₂) conditions at the seafloor. Anoxia stops nitrification, an aerobic process oxidizing ammonium to nitrite and nitrate that are reduced to N₂ in the anaerobic processes of anammox and denitrification. As a result, these N removing processes become limited by the lack of substrate. Consequently, denitrification rates in the Baltic Sea are very low in areas with frequent and prolonged anoxia (Tuominen et al. 1998). P bound to oxidized iron (Fe) compounds in the sediment can be released as a result of Fe reduction following decrease in O₂ concentrations and anoxia (Einsele 1936, Mortimer 1941, 1942). There is a strong correlation between the extent of anoxic sediments and the concentration of dissolved inorganic phosphorus (PO₄³⁻) in the Baltic Sea (Conley et al. 2002); the release of sediment P to the water column (internal P loading) plays a major role in eutrophication of the Gulf of Finland (Pitkänen et al. 2001). The Baltic Sea continually experiences anoxic episodes of variable frequency and duration (e.g. Laine et al. 1997, Sohle-
In the open Gulf of Finland, the replenishment of deep water O₂ is mainly regulated by salinity stratification, which is under the influence of irregular salt water intrusions from the western end of the Gulf and freshwater input from the River Neva at the eastern end. In coastal areas, seasonal mixing events restore O₂. The O₂ level at which harmful effects begin, the rates of reaction responses to changing O₂ conditions and the recovery times of system processes when oxic conditions are restored are as yet unclear.

We tested the sensitivity of denitrification to a short term (2 wk) anoxia exposure and a subsequent reoxidation of bottom waters, a common phenomenon, especially in the coastal areas of the Gulf of Finland in late summer. At the same time the effects of changing O₂ conditions on nutrient fluxes across the sediment-water interface were monitored. We also followed changes in different forms of sediment P after oxic conditions were restored.

**MATERIALS AND METHODS**

**Sediment and water.** Sediment used in the experiments was collected at the beginning of October 2004 from a coastal station (Gulf of Finland, Tvärminne Storfjärden, 59° 51’ 21” N, 23° 15’ 56” E) representing a characteristic, outer archipelago accumulation bottom consisting of soft mud. Water depth at the sampling station is 33 m. A box corer was used to collect the deeper, reduced sediment and an Ockelman sledge to collect oxidized surface sediment. At the time of the experiments, few macrofauna could be found at the sampling site, probably due to a low-oxygen period in August (Hietanen & Kuparinen in press). Therefore the mud used was gently sieved (1 mm for the reduced sediment and 0.5 mm for the surface slurry) to remove the scarce macrozoobenthos so that all subsamples were free of larger animals. Four aquaria (area: 600 cm², volume: 12 l) were packed with 8 cm of reduced mud covered with 2 cm of surface mud and filled with nutrient-free artificial seawater (Reef Crystals [Aquarium Systems] dissolved in distilled water) at a salinity matching that of the sampling location (6 psu). The experiment was conducted at 5°C in a cold room. This is the *in situ* bottom temperature in the study area. Aquaria were left to stabilize under oxic and anoxic conditions (see below) for 12 d before the first sampling. A similar approach of sieving and mixing sediment has been used in previous experiments with open Baltic Sea sediments (Tuominen et al. 1999). In those experiments (Tuominen et al. 1999), a NO₃⁻ peak similar to that occurring in undisturbed samples formed in just 2 d, indicating that the most disturbance-sensitive members of the microbial consortia, the nitrifiers, quickly recovered from the handling of the sediment.

**Anoxic-oxic regulation.** Aquaria were covered with 5 mm thick polyacrylic lids. Each was equipped with 2 magnetic stirrers constantly mixing the water phase at the highest possible rate without causing resuspension. Each aquarium lid had a hole for gas bubbling and, in 2 cases, for O₂ electrodes. Water phases of 2 of the aquaria were bubbled continually with air through the whole experiment (O₂-O₂ aquaria). The other 2 aquaria (N₂-O₂ aquaria) were connected to systems comprising an O₂ electrode (Pt-Ag, Strathkelvin Instruments Oxygen Electrode model 1302) immersed in the water phase of each aquarium and connected to an O₂ meter (Strathkelvin Instruments Oxygen Meter model 781), which was in turn connected to a regulator (PR Electronics) controlling a magnetic valve (Danfoss) connected to a nitrogen (N₂) gas bottle. Electrodes were calibrated in artificial seawater (matching the medium in experimental aquaria) subjected to vigorous air bubbling before calibration (100% O₂ saturation) and with N₂ gas (0% O₂ saturation). In order to control the level, the regulators were adjusted to 5% O₂ saturation (19 µM = 0.4 ml l⁻¹) with 2% tolerance, so that when O₂ concentration in the aquarium increased to 7% as measured by the electrodes, the regulator opened the magnetic valve to start N₂ gas bubbling, and kept it bubbling until 3% O₂ saturation was reached. As the gas supply tubing and the electrode were at the opposing ends of the aquaria, there was a short delay until the entire water phase was mixed and anoxia established at the electrode ends of the aquaria; the system caused N₂ bubbling always to proceed until aquaria were completely anoxic. Therefore, the actual O₂ saturation in the medium varied between 0 and 7% (up to 27 µM = 0.6 ml l⁻¹). Constant mixing with the magnetic stirrers prevented gas and nutrient gradient formation, and the lids reduced O₂ diffusion in from the atmosphere, as evidenced by the long pauses between N₂ bubbling periods.

Anoxia prevailed in the 2 aquaria for 17 d, with the exception of Day 12, when the N₂-O₂ vessels were oxic for 3 to 4 h due to technical problems while sampling. On Day 18, the N₂ bubbling was replaced with constant air bubbling in order to study process recovery after anoxia.

**Nutrients.** Samples for measurement of water phase concentrations of nutrients (ammonium NH₄⁺, nitrate + nitrite NO₃⁻ + NO₂⁻ = NOₓ, and phosphate PO₄³⁻) and sediment denitrification rate were taken on Days 12, 15 (anoxic and oxic), 19 and 27 (oxic). Samples for nutrient analyses (250 ml) were taken from the water phases of all aquaria and replaced with artificial seawater. Nutrients were analyzed using standard methods for seawater (Grasshoff et al. 1983).
**Denitrification.** Denitrification was measured using the isotope pairing technique (Nielsen 1992). Two replicate samples were collected from each aquarium in clear plastic cores (diameter 2.6 cm, height 9 cm) so that about half of the core was filled with the sediment and half with the water from above. Samples were enriched with potassium nitrate, $K^15NO_3$ (98% labelling, Cambridge Isotope Laboratories) to a final concentration of 100 $\mu M$ $^{15NO_3}$ in the water overlying the sediment, then incubated (with a magnetic stirrer on the lids of the cores) at 5°C in darkness for 3 to 4 h. Activity in the samples was terminated with $ZnCl_2$, and subsamples were stored in gas-tight 12 ml vials (Exetainer, Labco) to the National Environmental Research Institute, Silkeborg, Denmark for analysis of N2 isotope composition. Denitrification potential (D15: the denitrification of the added $^{15NO_3}$ was calculated from the formation of single-labelled ($^{14N}$-$^{15N}$) and double-labelled ($^{15N}$-$^{15N}$) dinitrogen assuming random isotope pairing of the uniformly mixed natural $^{14NO_3}$ and added $^{15NO_3}$ (Nielsen 1992):

$$D15 = (^{14N^{15N}}) + 2(^{15N^{15N}})$$

Naturally occurring denitrification (D14 or Dtot [total denitrification]) based on the unlabelled nitrate available for the denitrifiers was calculated as:

$$D14 = D15 \times [(^{14N^{15N}})/(2^{15N^{15N}})]$$

$Dtot$ (D14) can be divided into ‘direct’ denitrification (based on nitrate available in the water column: $Dw$) and coupled nitrification-denitrification ($Dn$: based on nitrate produced in the nitrification process) using the availability of the different nitrate sources as:

$$Dw = D15 \times (^{14NO_3})/[^{15NO_3}]$$

$$Dn = Dtot - Dw$$

where $[^{14NO_3}]$ and $[^{15NO_3}]$ represent the concentrations of the naturally occurring $^{14N}$ and the added $^{15N}$ in the NO$_3$ in the overlying water, respectively (Nielsen 1992).

**P fractionation.** Two replicate sediment samples for P fractionation studies were collected from each aquarium on Days 19 and 27. Samples were taken in clear plastic cores (see denitrification). The water above the sediment was removed (siphoned) immediately. Samples were capped and kept at 5°C in the dark until 2 sediment layers (0 to 2 cm and 2 to 4 cm) were separated from each core (O$_2$-O$_2$ treatment immediately; N$_2$-O$_2$ treatment the following day). N$_2$-O$_2$ cores were sliced in a N$_2$ atmosphere (O$_2$ content <5%) in a glove box. Subsamples were stored in small plastic bottles at 5°C. Chemical forms of sediment P were determined using a slightly modified P fractionation method by Jensen & Thamdrup (1993) (detailed description in Lukkari et al. in press). The method separates 6 P pools (Jensen et al. 1995): (1) loosely adsorbed and pore-water P (extracted with sodium chloride, NaCl, referred to hereafter as NaCl-iP), (2) redox-sensitive fraction of P bound to hydrated oxides of reducible metals (mainly those of Fe) (sodium dithionite, $Na_2S_2O_4$, in bicarbonate buffer NaHCO$_3$ at pH 7: NaBD-iP), (3) P bound to oxides of non-reducible Fe and aluminium (Al) (inorganic P in sodium hydroxide, NaOH: NaOH-iP), (4) apatite-P (hydrochloric acid, HCl: HCl-iP) and (5) residual, mainly organic, P (extracted with HCl after combustion: Res-P). The pool of (6) mobile organic P (non-reactive P: NRP) was determined as the difference between total P (TP) and dissolved inorganic P (iP), summarized from the first 3 steps. iP was determined from filtered (Nuclepore polycarbonate membranes, pore size 0.4 µm) extracts by UV-VIS spectrophotometry (Genesys 10uv Thermo Spectronic, with 50 mm flow injection cuvette) and TP with a spectrophotometer after acid persulphate digestion (Koroleff 1983). Total extractable P (TP$_{ex}$), calculated as the sum of all steps, is 90 to 95% of total P content in the sediment (TP$_{sed}$) determined after strong digestion (with aqua regia, fluoric acid and boric acid).

Sediment dry mass (DM) was first determined using a moisture analyzer (balance equipped with a halogen lamp dryer: Ohaus MB45) and the amount of fresh sediment extracted was determined as a fresh weight corresponding to 0.3 g DM. The volume of the extracts was always 30.0 ml, yielding a sediment DM-to-solution ratio of 1:100. Extraction was made at room temperature (on orbital shaker table) and during the first 2 steps (NaCl-iP and NaBD-iP) samples were handled in a N$_2$ atmosphere (O$_2$ < 5%) in a glove box. Supernatants were separated by centrifugation (3362 x g, 15 min) and extraction solutions and following rinsing solutions were combined into 1 sample in each step. Samples were preserved with sulphuric acid (pH 2, except those already containing acid) after filtration and stored at 5°C until analyses.

**Statistical analyses.** The results from the 2 aquaria per treatment were combined (independent variables: 2 treatments, 2 aquaria per treatment, 2 replicates per aquarium and different times). Differences in rates of $Dtot$, $Dn$, and $Dw$ were tested ($t$-tests, 2 sample assuming equal variances) between anoxic and oxic aquaria on each sampling occasion. Paired $t$-tests were used to evaluate the significance of the changes from one sampling occasion to the next. D15 based on the added $^{15NO_3}$ was analysed similarly. The significances of differences between concentrations of P fractions were tested on Days 19 and 27 within a treatment (paired $t$-test) and between treatments ($t$-test, 2 sample assuming unequal variances). For all tests, the significance level adopted was $p < 0.05$. 

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| Hietanen & Lukkari: Short-term anoxia and benthic processes | 295 |
RESULTS

Denitrification

Denitrification rates in the aquaria were high, averaging 650 µmol N m⁻² d⁻¹ (Fig. 1). On the first sampling on Day 12, no significant differences were found between the oxic and anoxic aquaria. From Day 12 to Day 15, Dtot (sum of Dn and Dw) and Dw (but not Dn) increased significantly in oxic and anoxic aquaria. There were significant differences in Dw and Dn between the treatments (Dw higher in anoxic, Dn higher in oxic), but not in Dtot rates. Oxic conditions were restored in N₂-O₂ aquaria on Day 17. From Day 15 to Day 19, the rates of Dw decreased and Dn increased significantly in the N₂-O₂ aquaria, and Dn decreased in the O₂-O₂ aquaria. Dtot and Dn (but not Dw) were significantly higher in the N₂-O₂ than in the O₂-O₂ aquaria on Day 19. From Day 19 to Day 27, only the Dw rate in N₂-O₂ aquaria decreased significantly, while other rates remained at the same level, and no differences between the treatments were found thereafter.

D15 doubled from Day 12 to Day 15 in the N₂-O₂ aquaria, but decreased back to the average level (300 µmol N m⁻² d⁻¹) by Day 19 when oxic conditions had been restored. There were no differences in D15 rates between Days 19 and 27 in the N₂-O₂ aquaria. In the O₂-O₂ aquaria, there were no differences in the D15 rates between days. The average level did not differ from the D15 rates in N₂-O₂ aquaria except on Day 15.

Nutrients

Nutrient dynamics clearly differed between the oxic and anoxic aquaria. As the water added to the aquaria was initially nutrient free (analysed at the beginning of the experiment and each time when added to the aquaria; average concentrations were NH₄⁺: 0.33 µM, NO₃⁻: 1.09 µM, PO₄³⁻: 0.03 µM), all nutrients in the water column originated from the sediment. Both NH₄⁺ and PO₄³⁻ occurred at high concentrations during anoxia, but after the shift from anoxic to oxic conditions they quickly decreased to the same low level measured in the oxic aquaria (Fig. 2). In contrast, while NO₃⁻ concentrations were higher in oxic than in anoxic aquaria (and were remarkably high in general), the difference was not especially marked, and there was no clear change in concentration after the shift from anoxic to oxic conditions (Fig. 2).

P fractionation

Concentration of the TPₐq was about 54 µmol (g DM⁻¹) at the surface (0 to 2 cm) and 35 µmol (g DM⁻¹) in the deeper (2 to 4 cm) layer. The NaBD-iP...
fraction formed the major part of TP in the surface layer, followed by Res-P in the deeper layer, HCl-iP and NRP were the most abundant and second most abundant fractions, respectively (Fig. 3).

NaCl-iP was the smallest P fraction, forming <1% of the TP$_{ext}$. NaCl-iP was slightly (but significantly) higher in O$_2$-O$_2$ than in N$_2$-O$_2$ aquaria on Day 19. Decrease in NaCl-iP with time in both layers of the O$_2$-O$_2$ aquaria was small, but statistically significant, whereas no change was detected in the N$_2$-O$_2$ aquaria. On Day 27, there were no significant differences in NaCl-iP between the 2 treatments.

NaBD-iP formed about 30 and 10% of the TP$_{ext}$ in the surface and deeper layers, respectively. On Day 19, NaBD-iP was significantly higher in the deeper layer of the N$_2$-O$_2$ aquaria (even though the difference was small), whereas it was higher in the surface layer of the O$_2$-O$_2$ aquaria small, but statistically significant, whereas no change was detected in the N$_2$-O$_2$ aquaria. On Day 27, there were no significant differences in NaBD-iP between the 2 treatments.

The share of the NRP fraction was slightly higher in the deeper layer (26%) than in the surface layer (19%), and it increased in both layers in all aquaria during the experiment. In N$_2$-O$_2$ aquaria, the increase was statistically significant in both depth layers. On both sampling days, N$_2$-O$_2$ aquaria had higher NRP concentrations than O$_2$-O$_2$ aquaria, but the differences were not significant.

Of the immobile fractions, NaOH-iP formed about 4 and 8% of the TP$_{ext}$ in surface and deeper layers, respectively, and did not vary significantly between sampling times or treatments. HCl-iP formed about 20 and 35% of the TP$_{ext}$ in the surface and deeper layers, respectively. Its concentration increased in the surface layer of all aquaria from Day 19 to Day 27, but the increase was not significant. Res-P fraction was slightly more abundant in the surface (about 27%) than in the deeper (about 21%) layer and its share decreased, but not significantly, during the experiment. There were no significant differences between the treatments despite the slightly higher concentration in the N$_2$-O$_2$ aquaria.

**DISCUSSION**

**Nitrification and denitrification**

The total denitrification rates were found to be less sensitive to O$_2$ deficiency than we expected. Denitrification based in the water column NO$_x$ (Dw) was
removing NO₃ directly from the water phase of the aquaria at a rate of about 12 µmol N aquarium⁻¹ d⁻¹, except on Day 15, when the removal rate doubled in anoxic aquaria. In addition, the coupled nitrification-denitrification used about 27 µmol N aquarium⁻¹ d⁻¹. Thus, denitrification was removing NO₃ at a rate of about 40 µmol N aquarium⁻¹ d⁻¹, and nitrification must have produced NO₃ at least at the same rate as there was no clear decrease in NO₃ concentrations (averaging 60 µM, 300 µmol aquarium⁻¹) between Days 12 and 27. The O₂ concentration during the ‘anoxic’ incubation period stayed below 7% (27 µM = 0.6 ml l⁻¹), with a short exposure to O₂ in the setting up of the experiments and during the first sampling. When filling the aquaria with artificial sea water at the onset of the experiments, NH₄⁺ probably leaked from the pore-water of the surface sediment to the originally nutrient-free water, despite all the care that was taken to prevent mixing of the sediment with the water (to the eye, there was virtually no mixing). The sampling further released NH₄⁺ from the sediment to the water. The short exposures (4 to 5 h at the onset and during the first sampling) to O₂ clearly provided the nitrifying bacteria with enough O₂ to oxidize NH₄⁺ released from the sediment, as shown by the high concentrations of NOₓ during the experiment, despite anoxia in the 2 aquaria. The small difference between the NO₃ concentrations in the oxic and anoxic aquaria was reflected in the higher concentration of NH₄⁺ in the anoxic aquaria. Nitrifiers have a high affinity for O₂ and the minimum concentration at which nitrification still proceeds can be as low as 2 µM O₂ (0.05 ml l⁻¹) (Gunderson 1966, Gunderson et al. 1966, Carlucci & McNally 1969, Henriksen & Kemp 1988). Indeed, nitrification is often fastest in sub-saturating oxygen conditions (Goreau et al. 1980, Prosser 1989, Voytek & Ward 1995a,b, Kester et al. 1997). However, in Chesapeake Bay, the nitrification rate is minimal at just below 125 µM O₂ concentration; sediment O₂ consumption in this system exceeds O₂ diffusion into the sediment and restricts nitrification to the sediment surface (Kemp et al. 1990). In our experiments, the dominant nitrate source for denitrification shifted from coupled nitrification-denitrification towards more water column NO₃ dependence during the anoxic incubations. This indicates that nitrification in the anoxic aquaria did not cease but concentrated in a shallower layer closer to the sediment surface leading to nitrate diffusion into the water column and uncoupling from denitrification. Clearly, nitrification dynamics cannot be predicted from the O₂ concentrations alone, but sediment O₂ consumption plays a role in defining the depth at which nitrification is still possible. In addition, an earlier exposure to low O₂ concentrations or even anoxia causes adaptations in nitrifying communities, so that bacteria repeatedly experiencing such conditions have a higher affinity for O₂ than bacteria from permanently oxic environments (Bodelier et al. 1996). During anoxic periods, bacteria in fluctuating O₂ environments conserve their capacity to nitrify, and are therefore able to react immediately to improved O₂ availability (Bodelier et al. 1996).

As the incubation cores used in the denitrification measurements were not gas-tight, O₂ was also diffusing into the cores during the 3 to 4 h incubation times. This diffusion is 50 µM O₂ h⁻¹ at 90 µM O₂ concentration in the water overlying the sediment (Hietanen & Kuparinen in press). According to Fick’s first law, diffusion was faster at the lower O₂ concentration used in these experiments. Therefore, it is also possible that some NH₄⁺ was oxidized to NO₃ in the cores during the incubation. However, the ambient NO₃ concentration in the aquaria (and consequently in the incubation cores) was so high that nitrification during incubation would have made no significant difference to the measured denitrification rates. The diffusion of O₂ into the cores and its subsequent consumption are also unlikely to have increased the O₂ concentration in the cores so quickly that it would have affected the depth of the denitrifying layer through increased O₂ penetration depth or changed the overall microbial activity in the cores.

D15 reflects Dw at unlimited NO₃ concentration. D15 and Dw doubled in the anoxic aquaria from Day 12 to Day 15. The increase may have been caused by a switch from oxic to anoxic metabolism in the facultatively anaerobic bacteria as the oxidized layer in the sediment became thinner. It may also be due to the possible uncoupling of nitrification and denitrification, as nitrification concentrated closer to the sediment surface and subsequently nitrate diffused into the water. However, as the Dn rate did not decrease at the same rate as the Dw rate increased, the overall effect of the 2 wk exposure to anoxia was an increase in the Dtot rate.

**Anammox**

In addition to denitrification, another N removing process, anaerobic oxidation of ammonium (anammox), has recently been found active in marine sediments (e.g. Thamdrup & Dalsgaard 2002, Dalsgaard & Thamdrup 2002, Trimmer et al. 2003). In this process, microbes oxidize NH₄⁺ with NO₃⁻ to form N₂. Accumulating knowledge about the process and the responsible microbes was reviewed by Dalsgaard et al. (2005). In marine sediments, both N removing processes can proceed at the same time, which complicates the calcu-
decreasing concentrations of PO$_4^{3-}$ in the water column to the sediment during a positive redox-shift in laboratory experiments has been described by Gächter et al. (1988), Hupfer & Uhmann (1991) Gunnars & Blomqvist (1997) and others. In addition, NaBD-iP did not increase in the previously anoxic aquaria between Days 19 and 27; 9 d after restoring the oxic conditions, there was no significant difference in sediment NaBD-iP fractions between the treatments. Time had no effect on the amount of Fe-bound P in the surface layers of the oxic aquaria, which indicates thatoxic incubation did not cause remobilization of NaBD-iP from the deeper to the surface layer. Instead, the increase of NaBD-iP in the deeper layer of the oxic aquaria towards Day 27 could have been caused by the prolonged exposure of the deeper layer to O$_2$ diffusion into the sediment (resulting from sediment disturbance during sampling). The slight decrease with time of NaCl-iP in the oxic treatment may have been caused by a slow diffusion of P from the pore water to the water column, where PO$_4^{3-}$ increased with time as well. Similar decrease of NaCl-iP with time did not occur in the previously anoxic aquaria, probably because the release of PO$_4^{3-}$ to the water column had already occurred at the beginning of the experiment. After the shift from anoxic to oxic conditions, the sediment started to oxidize and bind P more efficiently. By Day 27, the conditions had stabilized and there was no difference between treatments in NaCl-iP fractions.

The organic forms of sediment P (the more labile NRP and the more recalcitrant Res-P) also showed changes during the experiment, but these forms were not affected as clearly and quickly as the 2 most mobile P forms discussed above. Generally, NRP seemed to increase with time, while Res-P seemed to decrease and both fractions were higher in the anoxic treatment. However, the only statistically significant difference in these fractions was the increase of NRP with time in the anoxic treatment. The Res-P fraction is commonly included in the immobile forms of P (e.g. Jensen et al. 1995), but due to its organic nature, its composition and therefore its degradability vary depending on the deposited material and the prevailing conditions in the sediment. It is possible that part of the Res-P was degraded or transformed into more mobile P forms, but this cannot be confirmed with our results, and it is not clear whether the incubation time was long enough for such transformations. The NaOH-extractable NRP is not solely organic, but contains compounds formed in biological transformations, such as pyrophosphate (a degradation product of organic P) and polyphosphate, a storage compound of P found in many microorganisms (Hupfer et al. 1995a, Ahlgren et al. 2005). Microbial PO$_4^{3-}$ binding and polyphosphate synthesis in the sediment as well as release during changing O$_2$ conditions have been reported by Gächter et al. (1988) and the present study, but the effect of a 10 to 15% contribution of anammox on the results can be calculated according to Risgaard-Petersen et al. (2003, 2004; equations 15, 21 and 24). A share of 10% would overestimate Dtot by 2.5% in all but the anoxic samples, for which it would cause 4 and 6% overestimates of N$_2$ production on Days 12 and 15, respectively. A share of 15% would overestimate Dtot by 5% in all but the anoxic samples, for which it would cause 7 and 10% overestimates of N$_2$ production on Days 12 and 15, respectively. Interestingly, the higher error in the N$_2$-O$_2$ incubations during anoxic incubation decreased to the level in the oxic aquaria by Day 19, i.e. 1 d after the switch from anoxic to oxic conditions. The overestimations were not statistically significant in any case (t-test, p > 0.05).

**Phosphorus cycling**

The oxidation and reduction reactions of Fe and subsequent release and binding of P from and to the hydrated oxides of Fe occur relatively quickly (Davison & Seed 1983, Millero et al. 1987). Thus, we expected that fluctuating O$_2$ conditions would quickly affect the sediment P fractions, mainly those representing dissolved or loosely bound P (NaCl-iP) and P bound to reducible Fe-compounds (NaBD-iP). The first samples were taken when oxic conditions had prevailed for 1 d after the exposure to anoxia. The effect of anoxia could still be evident in the smaller NaCl-iP fraction in the previously anoxic aquaria, in which the reduced conditions had caused P release from sediment to the pore water and further diffusion to the overlying water (Krom & Berner 1980, Gächter et al. 1988, Hupfer & Uhmann 1991, Gunnars & Blomqvist 1997). The water column PO$_4^{3-}$ concentrations were also higher in the aquaria that had suffered from anoxia. On the other hand, no sign of the release of Fe-bound P (NaBD-iP) from the sediment was evident in the previously anoxic aquaria 1 d after the switch. Andersen & Ring (1999) reported that during long-term anaerobic incubation, most loss of sediment P occurred in the Fe-bound P fraction. In our study, however, given the fast oxidation of Fe$^{2+}$ to Fe$^{3+}$ (Davison & Seed 1983, Millero et al. 1987), one day of oxic conditions was probably sufficient for capture of the released P back to the sediment surface; this was also evident from the quickly decreasing concentrations of PO$_4^{3-}$ in the water columns of the aquaria. Similar sorption of PO$_4^{3-}$ from the water and further diffusion to the overlying water (Krom & Berner 1980, Gächter et al. 1988, Hupfer & Uhmann 1991, Gunnars & Blomqvist 1997). The water column PO$_4^{3-}$ concentrations were also higher in the aquaria that had suffered from anoxia. On the other hand, no sign of the release of Fe-bound P (NaBD-iP) from the sediment was evident in the previously anoxic aquaria 1 d after the switch. Andersen & Ring (1999) reported that during long-term anaerobic incubation, most loss of sediment P occurred in the Fe-bound P fraction. In our study, however, given the fast oxidation of Fe$^{2+}$ to Fe$^{3+}$ (Davison & Seed 1983, Millero et al. 1987), one day of oxic conditions was probably sufficient for capture of the released P back to the sediment surface; this was also evident from the quickly decreasing concentrations of PO$_4^{3-}$ in the water columns of the aquaria. Similar sorption of PO$_4^{3-}$ from the water column to the sediment during a positive redox-shift in laboratory experiments has been described by Gächter et al. (1988), Hupfer & Uhmann (1991) Gunnars & Blomqvist (1997) and others. In addition, NaBD-iP did not increase in the previously anoxic aquaria between Days 19 and 27; 9 d after restoring the oxic conditions, there was no significant difference in sediment NaBD-iP fractions between the treatments. Time had no effect on the amount of Fe-bound P in the surface layers of the oxic aquaria, which indicates thatoxic incubation did not cause remobilization of NaBD-iP from the deeper to the surface layer. Instead, the increase of NaBD-iP in the deeper layer of the oxic aquaria towards Day 27 could have been caused by the prolonged exposure of the deeper layer to O$_2$ diffusion into the sediment (resulting from sediment disturbance during sampling). The slight decrease with time of NaCl-iP in the oxic treatment may have been caused by a slow diffusion of P from the pore water to the water column, where PO$_4^{3-}$ increased with time as well. Similar decrease of NaCl-iP with time did not occur in the previously anoxic aquaria, probably because the release of PO$_4^{3-}$ to the water column had already occurred at the beginning of the experiment. After the shift from anoxic to oxic conditions, the sediment started to oxidize and bind P more efficiently. By Day 27, the conditions had stabilized and there was no difference between treatments in NaCl-iP fractions.

The organic forms of sediment P (the more labile NRP and the more recalcitrant Res-P) also showed changes during the experiment, but these forms were not affected as clearly and quickly as the 2 most mobile P forms discussed above. Generally, NRP seemed to increase with time, while Res-P seemed to decrease and both fractions were higher in the anoxic treatment. However, the only statistically significant difference in these fractions was the increase of NRP with time in the anoxic treatment. The Res-P fraction is commonly included in the immobile forms of P (e.g. Jensen et al. 1995), but due to its organic nature, its composition and therefore its degradability vary depending on the deposited material and the prevailing conditions in the sediment. It is possible that part of the Res-P was degraded or transformed into more mobile P forms, but this cannot be confirmed with our results, and it is not clear whether the incubation time was long enough for such transformations. The NaOH-extractable NRP is not solely organic, but contains compounds formed in biological transformations, such as pyrophosphate (a degradation product of organic P) and polyphosphate, a storage compound of P found in many microorganisms (Hupfer et al. 1995a, Ahlgren et al. 2005). Microbial PO$_4^{3-}$ binding and polyphosphate synthesis in the sediment as well as release during changing O$_2$ conditions have been reported by Gächter et al. (1988) and the present study, but the effect of a 10 to 15% contribution of anammox on the results can be calculated according to Risgaard-Petersen et al. (2003, 2004; equations 15, 21 and 24). A share of 10% would overestimate Dtot by 2.5% in all but the anoxic samples, for which it would cause 4 and 6% overestimates of N$_2$ production on Days 12 and 15, respectively. A share of 15% would overestimate Dtot by 5% in all but the anoxic samples, for which it would cause 7 and 10% overestimates of N$_2$ production on Days 12 and 15, respectively. Interestingly, the higher error in the N$_2$-O$_2$ incubations during anoxic incubation decreased to the level in the oxic aquaria by Day 19, i.e. 1 d after the switch from anoxic to oxic conditions. The overestimations were not statistically significant in any case (t-test, p > 0.05).
Effects of seasonal anoxia in the coastal Gulf of Finland

In our experiments, the shift from oxic conditions to anoxia in the water column was very sudden (only a matter of minutes), whereas in nature the O$_2$ concentration decreases more slowly in the course of days or even weeks, leaving the microbes time to adjust to changing conditions. In contrast, the reoxidizing of the bottom waters in the study area happens relatively quickly as a result of the autumn water column turnover, and the response in the processes is probably as fast in nature as it was in the experiments reported here. According to our results, the nitrification-denitrification system in the study area is highly flexible, and denitrification proceeds unaffected for at least 2 wk after the onset of anoxia. However, in the laboratory experiments, which were done with homogenised, sieved sediments, both the denitrification rates and the NO$_x$ concentrations were extremely high. In a year-round survey in the area where the sediment for the experiments was collected, the highest NO$_x$ concentration in the near-bottom water and the fastest denitrification rate measured were 3 µM and approximately 400 µmol N m$^{-1}$ d$^{-1}$ (Hietanen & Kuparinen in press), respectively, compared to 60 µM and 650 µmol N m$^{-1}$ d$^{-1}$, respectively, in the experiments reported here. Nevertheless, in our experiments, all of the NO$_x$ was produced by nitrification, which clearly was enhanced compared to natural conditions, despite the anoxia. A likely explanation for the enhanced nitrification is the increased availability of NH$_4^+$ (released during the setting up of the experiments and in sampling) combined with a thick (2 cm) oxidized layer, enabling nitrification to proceed, even during the anoxic period. If the oxidized layer had been thinner, nitrification might have been more sensitive to anoxia. Therefore, it is not only the length of an anoxic period, but also the frequencies of such periods and the capacity of the sediment to recover between exposures to anoxia that regulate the nitrogen dynamics in seasonally anoxic sediments.

The changes observed in the P fractions were also likely to have been more pronounced in the manipulated sediments than in nature as the pre-treatment of the sediment and the effective oxidation in a closed system may have enhanced oxidation and mineralisation processes in general. In our experiments, a short exposure to anoxia quickly caused a release of dissolved P from the 2 most labile forms of P (NaCl-iP and NaBD-iP). The effect was highly reversible, which was also evident in the quickly decreasing concentrations of the PO$_4^{3-}$ in the water phase afteroxic conditions were restored. As the replenishment of O$_2$ stores in nature is usually linked to an intense mixing of the water column (e.g. turnover or upwelling events), P released during anoxia may reach the productive layer instead of being readsorped into sediment, even when O$_2$ conditions are improved. However, in the oxic zone of the water column, dissolved P is rapidly bound to oxidized Fe-compounds (when Fe is present in sufficient concentration) and may settle with depositing particulates (e.g. Ellis-Evans & Lemon 1989, Gunnars & Blomqvist 1997, Gunnars et al. 2002). Furthermore, some of the organic P compounds participate in P cycling at the sediment-water interface over relatively short time scales and are affected by fluctuating O$_2$ conditions (e.g. Gächter et al. 1988, Hupfer et al. 1995b, Törnblom & Rydin 1998).

In a N limited ecosystem such as the Baltic Sea, a short exposure to anoxia may, according to our findings, further strengthen the N limitation by simultaneously enhancing N removal and increasing internal P loading. It has also been estimated that the eutrophied Baltic Sea has already entered a potentially self-sustaining vicious circle in which the increased nutrient availability enhances production and sedimentation of organic matter. Oxidation of this organic matter causes anoxia, leading to internal loading of P and a further lowering of the N:P ratio. The low N:P ratio, in turn, may lead to blooms of N fixing cyanobacteria, a common phenomenon in the eutrophied Gulf of Finland (Vahtera et al. 2007). Seasonal anoxia occurs in coastal areas when the increase in temperature both escalates the mineralisation rate of the organic matter and at the same time lowers O$_2$ solubility. In the open
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