Coupled anoxic nitrification/manganese reduction in marine sediments

STEFEAN HULTH,1,* ROBERT C. ALLER,2 and FRANCK GILBERT3

1Department of Analytical and Marine Chemistry, Göteborg University, S-412 96 Göteborg, Sweden
2Marine Sciences Research Center, State University of New York at Stony Brook, Stony Brook New York 11794-5000, USA
3Laboratoire d'Océanographie et de Biogéochimie (COM)-Campus de Luminy-Case 901, F-13228 Marseille Cedex 9, France

Abstract—Pore water and solid phase distributions of oxygen, manganese, and nitrogen from hemipelagic and shelf sediments sometimes indicate a close coupling between the manganese and nitrogen redox cycles. Reaction coupling must be sustained in part by biological reworking of Mn-oxide-rich surface sediments into underlying anoxic zones. Surface sediment from Long Island Sound (USA) was used in laboratory experiments to simulate such intermittent natural mixing processes and subsequent reaction evolution. Mixed sediment was incubated anoxically under either diffusively open (plugs) or closed conditions (jars). In closed anoxic incubations, pore water NO3− increased regularly to a maximum (up to 17 μM) after one to several days, and was subsequently depleted. Mn2+ was produced simultaneously with NO3−. N2O was also clearly produced and subsequently reduced, with a formation-depletion pattern consistent with coupled nitrification-denitrification in the anoxic sediment. Manipulative additions of Mn-oxides (5−10 μmol g−1 net) demonstrated that net anoxic NO3− production correlated directly with initial Mn-oxide content. During initial net NO3− production there was no evidence for SO4−2 reduction. A direct correlation was also observed between anoxic nitrification rates and estimated sulfate reduction rates; the larger the nitrification rates, the larger the eventual net sulfate reduction rates. Diffusively-open incubations using sediment plugs of four different thicknesses (2, 5, 10 and 20 mm) exposed to anoxic overlying water, also showed net production of pore water NO3− (<15−20 μM) despite the absence of NO3− in the overlying water for at least five days. In general, higher nitrate concentrations were maintained in the open relative to the closed incubations, due most likely to lower concentrations of dissolved reductants for NO3− in the open system. These experiments imply simultaneous coupling between the benthic nitrogen, manganese, and sulfur redox cycles, involving anoxic nitrification and sulfide oxidation to SO4−2. Anoxic nitrate production during Mn reduction indicates that nitrification and denitrification can occur simultaneously in subsurface sediments, without vertical stratification. The existence of anoxic nitrification implies new reaction pathways capable of increasing coupled sedimentary nitrification-denitrification, particularly in bioturbated or physically mixed deposits. Ltd

1. INTRODUCTION

Benthic nitrification-denitrification represents a significant component of the marine nitrogen cycle, particularly in organic rich shelf and hemipelagic sediments underlying oxygenated waters (Christensen, et al., 1987; Seitzinger, 1988; Reimers, et al., 1992; Seitzinger and Giblin, 1996). Nitrification and denitrification reactions in sediments are often conceptualized as occurring in a largely stratified reaction sequence characterized by an upper oxic layer, dominated by aerobic nitrification, and an underlying suboxic or anoxic region into which NO3− is transported and reduced during Corg remineralization (Froelich, et al., 1979; Jahnke, et al., 1982; Billen, 1982; Christensen and Rowe, 1984). Burrows, tubes, and fecal structures formed by benthic fauna superimpose various microenviromental patterns of radially or spherically distributed aerobic nitrification-anaerobic denitrification patterns onto this overall stratified reaction sequence (Jahnke, 1985; Kristensen, 1988; Aller, 1988; Brandes and Devol, 1995). In some sedimentary deposits, however, the average distributions of O2 and NO3− are apparently not consistent with nitrification exclusively in oxic regions, or with reduction of NO3− by Corg alone (Sørensen et al., 1987; Bender, et al., 1989; Aller, 1990; Murray et al., 1995; Luther, et al., 1997; Aller, et al., 1998). In particular, circumstantial evidence has mounted that within organic and Mn-rich regions of continental margin sediments, the N and Mn cycles may be closely coupled through reactions such as:

\[5 \text{Mn}^{2+} + 2 \text{NO}_3^- + 4 \text{H}_2\text{O} \rightarrow 5 \text{MnO}_2 + \text{N}_2 + 8 \text{H}^+ \]  

(1)

\[4 \text{MnO}_2 + \text{NH}_3^+ + 6 \text{H}^+ \rightarrow 4 \text{Mn}^{2+} + \text{NO}_3^- + 5 \text{H}_2\text{O} \]  

(2)

Evidence for the oxidation of Mn2+ by NO3− comes from the apparent oxidation of Mn2+ in surficial sediments where O2 is absent, that is O2 and Mn2+ distributions do not overlap, and where the only additional energetically favorable oxidants available in sufficient quantity, are NO3− and particulate MnO2. Depending on the assumed form of MnO2, reaction (1) is not thermodynamically favorable except in zones of relatively high NO3− and pH (Table 1). NO3− maxima well below the surficial oxic zone may sometimes result from biogenic irrigation and oxic microzones, but anoxic nitrification during MnO2 reduction, such as defined by reaction (2), is also clearly energetically possible under typical sedimentary conditions (Table 1). Because Mn-oxide enriched surficial sediment is rapidly mixed into subsurface anoxic zones during normal biogenic and physical reworking, an anaerobic, lithotrophic nitrification pathway could be a significant mode of biologically mediated N cycling.

In the present study, we experimentally simulated the natural

* Author to whom correspondence should be addressed.
Table 1. Change in standard free energy of the reaction R ($\Delta G_{R}$; kJ mol$^{-1}$) at conditions often found in marine sediments. Organic material is for simplicity represented as carbohydrate, CH$_2$O.

<table>
<thead>
<tr>
<th>Reaction Type</th>
<th>$\Delta G_{R}$ at pH=7</th>
<th>$\Delta G_{R}$ at pH=8</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>N OXIDATION</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nitrification</td>
<td>$2 \text{O}_2 + \text{NH}_4^+ \rightarrow 2 \text{H}^+ + \text{NO}_3^- + \text{H}_2\text{O}$</td>
<td>$-341$</td>
</tr>
<tr>
<td>Oxidation $\text{N}_2$ generation</td>
<td>$3/2 \text{MnO}_2 + \text{NH}_4^+ + 2 \text{H}^+ \rightarrow 4 \text{Mn}^{2+} + \text{NO}_3^- + 5 \text{H}_2\text{O}$</td>
<td>$-224$</td>
</tr>
<tr>
<td>Anoxic nitrification</td>
<td>$4 \text{MnO}_2 + \text{NH}_4^+ + 6 \text{H}^+ \rightarrow 5 \text{Mn}^{2+} + \text{NO}_3^- + 5 \text{H}_2\text{O}$</td>
<td>$-175$</td>
</tr>
<tr>
<td><strong>N REDUCTION</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Denitrification</td>
<td>$5/4 \text{CH}_2\text{O} + \text{NO}_3^- + \text{H}^+ \rightarrow 5/4 \text{CO}_2 + 1/2 \text{N}_2 + 7/4 \text{H}_2\text{O}$</td>
<td>$-596$</td>
</tr>
<tr>
<td>Anoxic sulfide oxidation I</td>
<td>$5/8 \text{FeS} + 4 \text{MnO}_2 + \text{NH}_4^+ + 7 \text{H}^+ \rightarrow 1/2 \text{N}_2 + 4 \text{Mn}^{2+} + 5/8 \text{SO}_4^{2-} + 5/8 \text{Fe}^{2+} + 11/2 \text{H}_2\text{O}$</td>
<td>$-590$</td>
</tr>
<tr>
<td>NH$_4^+$ oxidation</td>
<td>$5/3 \text{NH}_4^+ + \text{NO}_3^- \rightarrow 4/3 \text{N}_2 + 3 \text{H}_2\text{O} + 2/3 \text{H}^+$</td>
<td>$-421$</td>
</tr>
<tr>
<td>Anoxic sulfide oxidation II</td>
<td>$4 \text{MnO}_2 + \text{NO}_3^- + 5/8 \text{FeS} + 1/2 \text{N}_2 \rightarrow 1/2 \text{N}_2 + 5/8 \text{SO}_4^{2-} + 5/8 \text{Fe}^{2+} + 1/2 \text{H}_2\text{O}$</td>
<td>$-419$</td>
</tr>
<tr>
<td>Fe$^{2+}$ oxidation</td>
<td>$\text{NO}_3^- + 5 \text{Fe}^{2+} + 12 \text{H}_2\text{O} \rightarrow 5 \text{Fe(OH)}_3 + 1/2 \text{N}_2 + 9 \text{H}^+$</td>
<td>$-319$</td>
</tr>
<tr>
<td>Anoxic Mn$^{2+}$ oxidation</td>
<td>$5/2 \text{Mn}^{2+} + \text{NO}_3^- + 2 \text{H}_2\text{O} \rightarrow 5/2 \text{MnO}_2 + 1/2 \text{N}_2 + 4 \text{H}^+$</td>
<td>$-48.9$</td>
</tr>
<tr>
<td><strong>ADDITIONAL Mn REDUCTION</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mn reduction</td>
<td>$1/2 \text{CH}_2\text{O} + 3/2 \text{CO}_2 + 1/2 \text{H}_2\text{O} + \text{MnO}_2 \rightarrow 1/2 \text{Mn}^{2+} + 2 \text{HCO}_3^-$</td>
<td>$-190$</td>
</tr>
<tr>
<td>S$^0$ oxidation</td>
<td>$4/3 \text{H}^+ + \text{MnO}_2 + 1/3 \text{S}^0 \rightarrow 2 \text{Mn}^{2+} + 1/3 \text{SO}_4^{2-} + 2/3 \text{H}_2\text{O}$</td>
<td>$-157$</td>
</tr>
<tr>
<td>Anoxic sulfide oxidation III</td>
<td>$2 \text{H}^+ + \text{MnO}_2 + 1/4 \text{FeS} \rightarrow \text{Mn}^{2+} + 1/4 \text{SO}_4^{2-} + 1/4 \text{Fe}^{2+} + \text{H}_2\text{O}$</td>
<td>$-148$</td>
</tr>
<tr>
<td>Anoxic sulfide oxidation IV</td>
<td>$3 \text{H}^+ + \text{MnO}_2 + \text{HS}^- \rightarrow \text{Mn}^{2+} + \text{S}^0 + 2 \text{H}_2\text{O}$</td>
<td>$-136$</td>
</tr>
<tr>
<td><strong>C OXIDATION</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxic respiration</td>
<td>$\text{CH}_2\text{O} + \text{O}_2 \rightarrow \text{CO}_2 + \text{H}_2\text{O}$</td>
<td>$-521$</td>
</tr>
<tr>
<td>Sulfate reduction</td>
<td>$\text{CH}_2\text{O} + 1/2 \text{SO}_4^{2-} \rightarrow \text{HS}^- + \text{HCO}_3^-$</td>
<td>$-122$</td>
</tr>
<tr>
<td>Fe reduction</td>
<td>$\text{CH}_2\text{O} + 7 \text{CO}_2 + 4 \text{Fe(OH)}_3 \rightarrow 4 \text{Fe}^{2+} + 8 \text{HCO}_3^- + 3 \text{H}_2\text{O}$</td>
<td>$-107$</td>
</tr>
</tbody>
</table>

Conditions used to calculate $\Delta G_{R}$: [O$_2$] = 150 $\mu$M; [CO$_2$] = 50 $\mu$M; [HCO$_3^-$] = 2 000 $\mu$M; pN2 = 0.781 atm; [NO$_3^-$] = 5 $\mu$M; [NH$_4^+$] = 25 $\mu$M; [Mn$^{2+}$] = 100 $\mu$M; [Fe$^{2+}$] = 5 $\mu$M; [SO$_4^{2-}$] = 24 mM; [H$_2$S] = 25 $\mu$M; [HS$^-]$ = 5 $\mu$M. Activity coefficients were assumed to be 1. Values on the standard free energy of formation, $\Delta G_f$, were taken from Berner (1980) and Stumm and Morgan (1996).

2. MATERIAL AND METHODS

The general experimental approach was to physically mix the upper ~2 cm of sediment obtained from an organic-rich mud deposit in central Long Island Sound, and to either enclose the mixed material in sealed anoxic containers (closed incubation series), or expose defined thicknesses of sediment for a set period to an anoxic overlying water reservoir (open incubation series). Sediment and pore water were then sampled serially or at a discrete time subsequent to mixing in order to document net reaction patterns. Dissolved O$_2$ penetrates only a few millimeters into the sediment and the surface-most material (0–1 cm) is typically enriched in Mn-oxides by normal diagenetic cycling at the sampling site used (Aller, 1994). Mixing thus served to rapidly introduce Mn-oxides into underlying anoxic sediment. The naturally present Mn-oxides were augmented in one experimental series by addition of variable amounts of laboratory synthesized MnO$_2$.

2.1. Study Site and Sediment Sampling

Surface sediment was sampled at the PULSE study site in central Long Island Sound, USA (41 10.03N, 72 57.43W) during June, September, and November 1997. A number of studies have been carried out in the general location of the site, and biological, geochemical, and physical sedimentary properties are relatively well-characterized (Station NWC; Station P, PULSE site; e.g., Benninger, et al., 1979; Aller, 1994; Gerino, et al., 1998). Water depth averages ~15 m, sediments contain ~2% organic carbon, ~2% CaCO$_3$, and have an average porosity of ~0.8 (0–1 cm). Salinity is relatively constant between 24 and 25, and temperature varies sinusoidally between 2 and 22°C annually. The bottom was sampled with a Smith-MacIntire grab, obvious macrofauna and shell debris picked out, and the top ~2 cm of sediment removed. Sediment was sieved through a 0.45 mm Nylon mesh and homogenized. No water was added during sieving and homogenizing. Sediment collected in June was homogenized under N$_2$, initial handling of sediment in September and November was done under air.

2.2. Closed Incubations

A portion of the sieved and homogenized sediment was removed separately and placed in 20 ml (June and September) or 50 ml (November) polypropylene centrifuge tubes (jars). The completely filled jars were sealed, put in plastic bags, and incubated at room temperature in a bucket of anoxic mud obtained from the collection site (Fig. 1). Subsequent to the manual removal of air, the bags were placed with the upper edges at least 5 cm into the mud to ensure completely anoxic
conditions (surficial O\(_2\) penetration \(-1\text{--}2\) mm, Fe-monosulfides present). Temperatures were 20.7--21.6°C (June), 20--22.1°C (Sept.), and 21.2--22.1°C (Nov.). This type of incubation is referred to as “closed anoxic incubation” (e.g., Martens and Berner, 1974).

During the November incubation experiment, Mn-oxide was added to two separate sediment portions before enclosure. The MnO\(_2\) used was formed from the oxidation of Mn\(^{2+}\) by MnO\(_4^2-\) under basic conditions, equilibrated with sea water, and aged 12 years at 4°C (Balistrieri and Murray, 1982; Aller and Rude, 1988). The precipitate has been inferred as the mineral phase vernadite (\(\delta\)MnO\(_2\)) based on the average oxidation state of the solid (O:Mn = 1.94 ± 0.02), and its X-ray diffraction pattern (Burns and Burns, 1979; Balistrieri and Murray, 1982). In this case, the sieved and homogenized natural sediment was split into three portions, corresponding to three different treatments. No extra MnO\(_2\) was added to the first sediment portion (A), while an estimated amount of \(-13\text{--}33\) mmol g\(^{-1}\) dry sediment (assuming molecular weight of MnO\(_2\)) of the synthesized MnO\(_2\) was added to the second (B) and third (C) portion, respectively. All three portions were homogenized by hand using plastic spoons before packing the sediment into jars.

During each incubation series, 1--3 tubes were removed every 1--3 days, centrifuged (12 minutes, 3800 rpm), filtered (0.4 \(\mu\)m) and the obtained pore water analyzed for total carbonate (\(\text{CO}_2\) or C\(_T\)), alkalinity (A\(_T\)), DIN (NO\(_3^2\), NO\(_2^2\), and NH\(_4^+\)), Mn\(^{2+}\) and pH. The centrifuged sediment was frozen for later analyses of manganese.

2.3. Diffusively Open Incubations

Portions of the surface sediment from the June sampling were also used for anoxic incubations under diffusively open conditions. The sieved and homogenized sediment was placed in polycarbonate tube-sections sealed at the base (plugs), and the sediment surface was smoothed using a stainless steel spatula. Plugs had variable thickness of 2, 5, 10 and 20 mm. The sediment-filled plugs were placed in a polycarbonate reservoir containing filtered (0.2 \(\mu\)m), anoxic overlying water from the sampling location (salinity of 24). The anoxic overlying

---

**Fig. 1.** Schematic illustration of the experimental set-up used to study net reaction rates within sediments under diffusively closed conditions (closed anoxic incubation). To ensure completely anoxic conditions the jars were placed with their upper edges at least 5 cm into the mud. Reaction rates were inferred from the production rates of solutes to the pore water with time of incubation.
water reservoir was in a glove bag, and was constantly purged before
(over night) and during the experiment with a mixture of N\textsubscript{2} and CO\textsubscript{2}
gas scrubbed of O\textsubscript{2} using an Alltech O\textsubscript{2} trap and an O\textsubscript{2} indicating
assembly (Fig. 2). Previous experiments have demonstrated that O\textsubscript{2} is
maintained below detection in this set-up. The incubation was per-
formed in the dark at a temperature of 19.4–21.7°C. This type of
incubation is referred to as “diffusively open” because exchange of
solutes is possible between sediment and overlying water (e.g., Aller
and Mackin; 1989). The water reservoir was mixed by a centrally
located Teflon-coated magnetic stirring bar. Before use, the polycar-
bionate reservoirs, plug assemblies, and stirring bars were soaked in 4%
HCl for 10 days or more. After soaking they were thoroughly rinsed in
double distilled water (DDW) and soaked in DDW for at least 7 days.

Overlying water (~20 ml) was sampled periodically during the open
plug incubations for $\Sigma$CO\textsubscript{2}, DIN (NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, and NH\textsubscript{4}\textsuperscript{+}) and Mn\textsuperscript{2+}
using a polypropylene syringe and Tygon tubing extension. Prior to
use, the assembly was washed in 4% HCl, thoroughly rinsed with
DDW and then with sample. Overlying water was filtered through

Fig. 2. Typical experimental set-up illustrating the open plug incubation technique. Anoxic sediment was incubated in
plugs (52 mm diameter; 2, 5, 10 and 20 mm thickness) under diffusively open conditions, where exchange was possible
between the sediment and overlying water. The reservoir (15.4 L of overlying water) was kept in a glove bag and to ensure
anoxic conditions, the glove bag and the overlying water were constantly purged with a mixture of oxygen free N\textsubscript{2} and CO\textsubscript{2} gas.
in-line polycarbonate filters (0.40 μm pore size) directly into acid cleaned polypropylene vials.

After 12 days, plugs were sampled for pore water and solid phase composition. Plugs were individually removed from the water reservoir using stainless steel forceps, sediment quickly transferred to a polypropylene syringe (prewashed in 4% HCl and rinsed with DDW and overlying water from the container) using a stainless steel spatula. The front of the syringe contained a small amount (~2 cm²) of folded nylon screen to prevent sediment extrusion. The syringe was placed into a press which applied pressure by moving the plunger forward, and pore water passed through an in-line mounted 0.4 μm polycarbonate filter directly into an attached receiving polypropylene syringe without air contact (Aller, et al., 1998). The filter had been rinsed with overlying water from the incubation reservoir. Sediment from which the pore water was separated was frozen for later analyses of manganese in the solid phase.

Additional plugs were incubated for the determinations of pore water distributions of pH by direct insertion into sediment of a mini-combination electrode (Ag–AgCl) mounted on a micro-manipulator.

2.4. Analytical Methods

Sediment plug processing and pore water separation were complete within a few minutes (<5 min) after removal from the overlying water reservoir.

Samples for ΣCO₂ were analyzed immediately by FIA conductivity (Hall and Aller; 1992). DIN samples were frozen. Concentrations of NO₂⁻ and NO₃⁻ were determined using standard colorimetric methods (Strickland and Parsons, 1972) applied to a flow injection system (FIA). NH₄⁺ was determined using flow injection conductivity (Hall and Aller, 1992). Measurements of pH were standardized on the Hansson pH scale (Hansson, 1973). Alkalinity was determined by potentiometric titration and Gran evaluation according to the principles outlined by Haraldsson et al. (1997), with a precision (1 mL sample volume) better than 0.7% RSD. 50–100 μL of concentrated HCl was added to the Mn²⁺ samples, producing a pH of 1–2, before storage in a refrigerator. Dissolved manganese was determined using a Shimadzu flame AAS. Sulfate in the pore water from the closed anoxic incubations was determined gravimetrically with BaCl₂. Reactive solid phase Mn was estimated by leaching a small portion of the sediment for 15 hours using hydroxylamine/acetic acid (Chester and Hughes, 1967), and measuring manganese in the leachate.

Porosity (φ), was calculated from the weight of water loss after drying 5 mL of sediment at 70°C for at least 24 hours (until constant weight) at the beginning of each incubation series.

Net reaction rates in the closed incubations were calculated over specific time intervals from least squares linear fits to concentration versus time plots. Corrections for reversible adsorption were made for NH₄⁺, assuming a linear adsorption coefficient K = φK⁺ρₗ (1 − φ), where ρₗ = 2.6 g cm⁻³, and K⁺ = 0.77 (K⁺ previously measured, Aller and Aller, 1998).

Net production or consumption rates of pore water solutes (R) in plugs were calculated assuming the steady state relation (Aller and Mackin, 1989):

$$ R = 3^s D_s^* (C_{ss} - C_T) / L^2 $$  \hspace{1cm} (3)

where D_s is the whole molecular diffusion coefficient (D_s = ρ^s D₀ (Ullman and Aller, 1982); D₀ is the free solution diffusion coefficient at the appropriate temperature and salinity, estimated from Li and Gregory, 1974), Cₚₗ is the average steady state concentration in a plug of thickness L, and Cₚₗ is the overlying water solute concentration at the end of the experiment.

3. RESULTS AND DISCUSSION

3.1. Closed Jar Incubations—Unamended

The closed incubations demonstrated time-dependent patterns and rates of organic matter degradation typical of surface sediment in central Long Island Sound (Fig. 3; Table 2). Net remineralization rates, as evidenced by the initial linear pro-

Fig. 3. Organic matter mineralization during the June (filled circles), September (open squares), and November (filled triangles) closed jar incubations. A. Mobilization of total carbonate (ΣCO₂ or Cₜ) and alkalinity (Aₚ) to the pore water during the June, September, and November incubations. B. Production of ammonium during the un-amended and, C. amended November incubation. In the amended incubation, no MnO₂ was added to the control (A; •), while an amount of 13 and 33 μmol of MnO₂ g⁻¹ dry sediment was added to the (B; △) and (C; ▽) incubation series, respectively. The dashed lines represent the initial production used to calculate net rates of solute mobilization.
duction rates of $\Sigma$CO$_2$, alkalinity ($A_T$), and NH$_4^+$ were similar during the June and November incubations, but somewhat lower during the September incubation (Fig. 3; Table 2). These measured production rates would support, for example, a steady state flux of $\Sigma$CO$_2$ from the top 2 cm of ~21, 12, and 22 mmol m$^{-2}$ d$^{-1}$ (June, Sep., Nov.), values comparable to those observed in previous remineralization measurements in the sediment collection area (e.g., Aller, 1980; Westrich and

<table>
<thead>
<tr>
<th>Incubation</th>
<th>Mn$^{2+}$; pw</th>
<th>Mn$^{2+}$; sp</th>
<th>$K_{Mn}$</th>
<th>NH$_4^+$</th>
<th>$\Sigma$CO$_2$</th>
<th>$A_T$</th>
<th>Nitr.</th>
<th>Denitr.</th>
<th>SO$_4^{2-}$</th>
<th>ϕ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Closed</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>June</td>
<td>7.86</td>
<td>31.0</td>
<td>3.9</td>
<td>76</td>
<td>1390</td>
<td>–</td>
<td>11</td>
<td>–6.5</td>
<td>–175</td>
<td>0.77</td>
</tr>
<tr>
<td>Sep.</td>
<td>24.3</td>
<td>252</td>
<td>10</td>
<td>52</td>
<td>–</td>
<td>828</td>
<td>4.0</td>
<td>–0.61</td>
<td>–</td>
<td>0.71</td>
</tr>
<tr>
<td>Nov.; A</td>
<td>157</td>
<td>317</td>
<td>2.0</td>
<td>187</td>
<td>–</td>
<td>1450</td>
<td>0.44</td>
<td>–0.32</td>
<td>–147</td>
<td>0.76</td>
</tr>
<tr>
<td>Nov.; B</td>
<td>152</td>
<td>216</td>
<td>1.4</td>
<td>171</td>
<td>–</td>
<td>1440</td>
<td>1.0</td>
<td>–1.1</td>
<td>–211</td>
<td>0.76</td>
</tr>
<tr>
<td>Nov.; C</td>
<td>150</td>
<td>244</td>
<td>1.6</td>
<td>167</td>
<td>–</td>
<td>1400</td>
<td>2.3</td>
<td>–0.64</td>
<td>–243</td>
<td>0.76</td>
</tr>
<tr>
<td>Open</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L</td>
<td>Mn$^{2+}$; pw</td>
<td>Mn$^{2+}$; sp</td>
<td>$K_{Mn}$</td>
<td>NH$_4^+$</td>
<td>$\Sigma$CO$_2$</td>
<td>$A_T$</td>
<td>$\Sigma$NO$_3^-$</td>
<td>NO$_2^-$</td>
<td>SO$_4^{2-}$</td>
<td>ϕ</td>
</tr>
<tr>
<td>2</td>
<td>n.d.</td>
<td>384</td>
<td>–</td>
<td>623</td>
<td>n.d.</td>
<td>1090</td>
<td>68.2</td>
<td>–</td>
<td>0.77</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>42.0</td>
<td>363</td>
<td>–</td>
<td>108</td>
<td>1300</td>
<td>–</td>
<td>195</td>
<td>10.9</td>
<td>–</td>
<td>0.77</td>
</tr>
<tr>
<td>10</td>
<td>88.9</td>
<td>348</td>
<td>–</td>
<td>154</td>
<td>404</td>
<td>–</td>
<td>47.9</td>
<td>1.60</td>
<td>–</td>
<td>0.77</td>
</tr>
<tr>
<td>20</td>
<td>71.9</td>
<td>230</td>
<td>–</td>
<td>94.5</td>
<td>489</td>
<td>–</td>
<td>8.97</td>
<td>0.439</td>
<td>–</td>
<td>0.77</td>
</tr>
</tbody>
</table>

All rates are in $\mu$M d$^{-1}$. Net rates of nitrification (nitr.) were calculated from the initial mobilization of $\Sigma$NO$_3^-$ to the pore water. The phase of nitrate disappearance was used to estimate net denitrification rates (denitr.). Production rates of Mn$^{2+}$ were estimated both from net pore water concentration changes (pw), and from loss of solid phase reactive MnO$_2$ (sp). The absorption coefficient, $K_{Mn}$, was calculated as the (Mn$^{2+}$; sp/Mn$^{2+}$; pw) ratio. Also given are the actual plug thickness (L, mm) used during the open plug incubations, and the porosity (ϕ). A, B and C refer to the different additions of Mn$^{2+}$ oxides to the sediment prior to starting the November incubation (0, 13 and 33 $\mu$mol g$^{-1}$ dry sediment).

Fig. 4. The evolution of pH in the pore water during the September incubation is shown as a representative example on the behavior of pore water pH during the closed jar incubations. One to three replicate sediment samples were incubated. Average and standard deviation of these replicates are indicated. The samples were standardized using the Hansson pH scale (Hansson, 1973).
Berner, 1984; Sun, et al., 1991). $\Sigma$CO$_2$ and NH$_4^+$ net production rates were more or less constant during an initial period of four to six days, then decreased, most likely reflecting the depletion of labile organic matter, and/or in the case of CO$_2$, precipitation of solid phase carbonates (e.g., Ca, Mn-carbonates). During the build-up of decomposition products, pore water pH decreased exponentially from initial values near seawater, to about 7.3 (Fig. 4).

Elevated Mn$^{2+}$ concentrations, coupled with decreasing reactive solid phase Mn, are evidence of net Mn reduction during C$_{org}$ remineralization within the incubation series (Fig. 5, 6). Both the quantities of reactive Mn and the overall concentration of dissolved Mn$^{2+}$ differed in the various unamended incubations. Initial reactive Mn was highest in the June and November samples ($\sim$14 $\mu$mol g$^{-1}$) and lowest in September ($\sim$6 $\mu$mol g$^{-1}$). Likewise, maximum pore water Mn$^{2+}$ concentrations were 700–800 $\mu$M during the June and November incubations, but reached only $\sim$260 $\mu$M in the September samples (Fig. 6). The observations of the lowest net remineralization rates, as measured by $\Sigma$CO$_2$ and NH$_4^+$ release, and the lowest reactive Mn in the September incubation series, are consistent with partial loss of the surface most sediment layer or homogenization of a surface interval $>2$ cm during sample collection.

Nevertheless, the primary characteristics of all incubation series are relatively rapid remineralization and substantial Mn-oxide reduction. The differences in the estimates of Mn$^{2+}$ production rates based on pore water net concentration changes and those calculated from loss of solid phase reactive Mn (Table 2), imply reversible adsorption coefficients in the range of $K_{Mn}$ 2–10 (simple ratio estimates), similar to those reported from other sediments (Canfield, et al., 1993).

Although the closed incubations were anoxic, there was clear evidence of a pattern of progressive net NO$_3^-$ formation over a period of 2 to 4 days (Fig. 7A). Based on the measured remineralization rates of $\sim$1 mM $\Sigma$CO$_2$ d$^{-1}$ (Table 2), any O$_2$ occluded initially in the pore water could not have persisted for more than a few hours after enclosure (Table 2). The excess NO$_3^-$ produced during the first few days of incubation was subsequently depleted, but it was lost at a sufficiently slow rate given the intensity of $\Sigma$CO$_2$ remineralization, that continued production of NO$_3^-$ must have occurred during the entire period of incubation. Similarly, NO$_2^-$ was also obviously produced and depleted in a net sense, but again, concentrations were sustained at a readily detectable level throughout the incubations. Maximum NO$_3^-$ concentrations occurred after NO$_3^-$ maxima, indicating a coupling between the two constituents con-
sistent with denitrification. Most importantly, these time-dependent concentration patterns directly and categorically demonstrate the production of pore water NO$_3^-$ and NO$_2^-$ for extended periods in the absence of detectable O$_2$ or any conceivable source of dissolved O$_2$.

Although NO$_3^-$ and NO$_2^-$ were produced in all incubations, the net rates of production and the maximum NO$_3^-$ or NO$_2^-$ concentrations attained, differed substantially between the various incubation series (e.g., maxima: 0.7–17 μM NO$_3^-$, Fig. 7; rates: 0.4–11 μM NO$_3^-$ d$^{-1}$, Table 2). The anoxic nitrate production rate measured during the June incubation was the largest found in this investigation. The initial increase in this case corresponded to a net nitrification rate of 11 μM d$^{-1}$, or a steady state flux of 170 μmol m$^{-2}$ d$^{-1}$, assuming the rate applied over a 2 cm depth interval. Such a rate is similar to, or higher than, net nitrification rates reported from other coastal marine sediments open to O$_2$ diffusion (Szwerinski, 1981; Enoksson and Samuelsson, 1987). The only known sedimentary constituents present under the incubation conditions with sufficient oxidation potential and abundances capable of sustaining such nitrification rates are Mn-oxides. It is clear that Mn reduction occurred simultaneously with net NO$_3^-$ production and at rates generally sufficient to account for the observed NO$_3^-$ production rates assuming stoichiometric relations such as reaction (2) (Tables 1, 2).

3.2. Stimulation of Anoxic NO$_3^-$ Formation by Addition of MnO$_2$

Addition of Mn-oxide to sediment in the November closed incubation series stimulated the anoxic formation of NO$_3^-$ and increased the rate of NO$_3^-$ formation in direct proportion to the quantity of Mn-oxide added (Figs. 8, 9). The additions of ~13 and ~33 μmol of MnO$_2$ g$^{-1}$ dry sediment stimulated NO$_3^-$ production rates 2 to 5X compared to unamended controls. The calculated anoxic nitrification rates for the different treatments were 0.4, 1, and 2.3 μM d$^{-1}$, respectively (Table 2). In the present cases, the added Mn increased the extractable, reactive Mn by ~15–50% above the initial starting values in the unamended sediment, but the quantity extracted was well within the range found in diagenetically enriched surface sediments from Long Island Sound (e.g., Aller, 1994). Only ~20% of the total added MnO$_2$ was extracted by the hydroxylamine / acetic acid leach, presumably because the aged oxides were relatively well crystallized (Fig. 9B). The percent recovery, however, was constant.

Further support for substantial metal-oxide reduction came from the excess alkalinity production relative to net SO$_4^{2-}$ reduction observed in the incubations. Calculated rates correspond to C : S production / consumption ratios (A$_T$/ SO$_4^{2-}$) of ~ 9.9 for the incubation without additional MnO$_2$ (A), ~6.8 for the (B) incubation, and ~5.8 for the incubation with the largest
amount of MnO₂ added to the sediment (C) (Fig. 10; Table 2). These ratios are significantly larger than that expected from the reaction of net sulfate reduction alone (C : S ~2), and are consistent with alkalinity production during Mn reduction. When comparing net rates of ΔT or ΔCO₂ production with net rates of sulfate and manganese reduction, there appears to be additional alkalinity generated beyond that accountable by Mn²⁺ remaining in solution (Table 2). This indicates either Fe reduction (not measured) or ion exchange of Mn²⁺ with major elements such as Ca²⁺, Mg²⁺, Na⁺, or K⁺ (not measured).

Increased metal reduction in November incubations (B) and (C) were also indicated by pore water pH, where after 8–10 days of incubation, the pH was increased by approximately 0.05 pH units in the (B) and (C) incubations compared to the control (A) (Fig. 11). The increased pH with additions of MnO₂ is consistent with stimulated Mn-reduction (Emerson, et al., 1982). In each incubation series, there was an initial phase of obvious net Mn²⁺ production. The net Mn²⁺ build-up stabilized either as reactants were depleted or consumption processes, such as precipitation of Mn-carbonates, began to dominate (Fig. 12). The initial rates of net Mn²⁺-production, as estimated from pore water concentrations, were found to be more or less constant for the different treatments at 150–160 μM d⁻¹ (Table 2). However, there was a relatively lower net alkalinity attained with increasing additions of Mn-oxides, indicative of Mn-carbonate precipitation (Fig. 10A). After 8–10 days of incubation, alkalinity was found to be 6–7% higher in (A) compared to (B) and (C).

3.3. Effect of Diffusive Exchange

The diffusively open plug incubations also demonstrated clear evidence of NO₃⁻ production in sediment under anoxic conditions, but with some important contrasts compared to the closed incubations (Figs. 7, 8, 13). The initial overlying water NO₃⁻ concentrations were completely depleted after ~1 week,
while \(\text{NH}_4^+\) increased progressively with time, indicative of net denitrification and the absence of \(O_2\) in the overlying water as well as a net flux of \(\text{NH}_4^+\) from the sediment plugs (Fig. 14). Nitrate concentrations of 14–20 \(\mu\)M were found in the pore water of all the plugs incubated under anoxic conditions, despite the fact that no \(\text{NO}_3^-\) had been detected in the overlying water for more than five days prior to sediment sampling (Fig. 13), and there was no hint of oxidized sediment in the plugs (i.e., Fe-oxides). The calculated anoxic production rates of \(\text{NO}_3^-\) for the different plug thicknesses of 2, 5, 10 and 20 mm (based on equation 3) were 1090, 195, 48 and 9 \(\mu\)M d\(^{-1}\), respectively (Table 2). Net production of \(\text{NO}_2^-\) and \(\text{NH}_4^+\) were also evident in the plugs, with average formation rates also varying inversely with plug thickness as for \(\text{NO}_3^-\) (Fig. 15; Table 2).

Pore water and solid phase distributions of manganese suggested that Mn-oxide reduction was occurring in all plugs at the time the experiment was terminated and the pore water extracted (Fig. 16). Rates of Mn-reduction, as evidenced by the production rates of \(\text{Mn}^{2+}\) to the pore water (equation 3) at the time of experiment termination, were (2 mm not detectable), 42, 89, and 72 \(\mu\)M d\(^{-1}\) (Table 2). The lack of a calculatable production rate from the pore water data for the 2 mm plug reflects a relatively high overlying water Mn\(^{2+}\) concentration that resulted from the Mn\(^{2+}\) flux out of all plugs (i.e., small concentration difference in equation 3). The contents of reactive MnO\(_2\) in the sediment were clearly sufficient to support Mn reduction and anoxic nitrate formation in all plugs over the incubation period (Fig. 13, Table 2). Note the solid phase based estimates of reaction rates tend to integrate over a longer period of net reaction than do the pore water based estimates, the latter reflecting conditions close to the time of plug sampling.

The inverse relations of solute production rates with sediment plug thickness are typically found for a variety of diagenetically reactive constituents under diffusively open conditions, and reflect a combination of changes in both absolute and relative reaction rate (Aller and Aller, 1998). For example, the lowered build-up of reactive solutes in diffusively open sediment often results in less intense back-reactions such as authigenic mineral precipitation (e.g., MnCO\(_3\)), and thus a greater net release of produced solutes into the contacting water reservoir (e.g., Mn\(^{2+}\), HCO\(_3^-\)). In addition, there is evidence for an
absolute stimulation of anaerobic microbial activity under conditions of progressively greater solute exchange (Aller and Aller, 1998).

The primary conclusion from both types of incubation treatments is that NO₃⁻ and NO₂⁻ can be produced in the absence of O₂, and that N oxidation takes place simultaneously with Mn-oxide reduction. These results are consistent with net reactions such as reaction (2), but how is it that high nitrate concentrations could be maintained for a more extended period and at a greater concentration in the open anoxic, but not in the closed anoxic incubations (Fig. 7, 8 and 13)? Both treatments used exactly the same sediment, the only difference being the solute transport regime. We think the major differences in the incubations result primarily from changes in the rates of concentration-dependent reactions and multiple competitive reactions, rather than the occurrence of different reactions per se. The concentrations of reduced solutes in the closed incubations are higher than in the open plugs. Mn-oxides likely competitively react with reduced C, HS⁻, DOC, DON, and NH₄⁺. Higher concentrations of reduced reactants, for example DOC (e.g., Stone and Morgan, 1984), may either competitively suppress NH₄⁺ oxidation or use up oxidant (MnO₂) more rapidly than in a diffusively open case where a portion of reduced solutes escape into overlying water. In a similar way, NO₃⁻, once anoxically produced, is also exposed to higher concentrations of potential reductants (e.g., DOC, DON, NH₄⁺, HS⁻). Thus, local consumption rates for NO₃⁻ in the closed system are likely higher than in the open one. When sediment is otherwise equivalent, and the diffusively open system is not so thin that eventual concentration build-ups vanish, a higher concentration of NO₃⁻ should be sustained in the diffusively open case (higher production, lower consumption reactions). It is also possible that the open system results in diffusive loss of a dissolved
inhibitor of anoxic nitrification, thereby increasing production rates.

Because of pH gradients in the diffusively open plugs (Fig. 17), it is possible that oxidation of Mn$^{2+}$ by NO$_3^-$ (reaction 1) could occur more favorably in surficial regions of the sediment sections or in the overlying water reservoir (pH-dependent $\Delta G_R$, Table 2). Under such circumstances, or in the case of anoxic surface layer in the presence of oxygenated overlying water, an internal redox cycle could be set up in the sediment whereby production of a dissolved oxidized Mn (possibly chelated Mn$^{3+}$; Luther et al., 1994; Kostka et al., 1995) in a surface zone could diffusively supply oxidant to an underlying anoxic region.

3.4. Anaerobic, Lithotrophic Nitrification and Denitrification

We interpret the experimental results to indicate that anoxic nitrification occurs in sediments and, because of the favorable energetics, that the overall reaction is likely to be microbially mediated through a process of anaerobic, lithotrophic nitrification associated with Mn-oxide reduction (reaction 2). The depletion of NO$_3^-$ and NO$_2^-$ subsequent to their formation is additional evidence that denitrification is closely coupled to lithotrophic nitrification. The abundance of potential reductants of NO$_3^-$ in anoxic sediments implies that denitrification may not be strictly coupled to C oxidation as is often depicted. In particular, NH$_4^+$, Fe$^{2+}$, HS$^-$ (S$^0$, FeS, FeS$_2$), and C$_{org}$ are known reductants utilized by bacteria during denitrification (Bender, et al., 1989; Straub, et al., 1996; Luther, et al., 1997). The very rapid denitrification that would occur in subsurface sediments subsequent to anoxic nitrification is a likely reason why anoxic nitrification as a net reaction is not always evident in pore water profiles.

One consequence of a lithotrophic based nitrification cycle is that a greater proportion of benthic O$_2$ uptake could be directed to the N cycle indirectly through Mn-oxidation/reduction. Particle bioturbation and physical mixing processes would then play a dominant role in controlling solid phase oxidant transport into underlying sediments and promoting coupled lithotrophic nitrification-denitrification. The occurrence of a subsurface internal nitrification-denitrification cycle with small net production of NO$_3^-$ would complicate interpretation of N$_2$ production rates from NO$_3^-$ profiles (underestimates), and might cause discrepancies between various types of estimates

![Fig. 11. The evolution of pH in the pore water during the November closed jar incubation. After 8–10 days of incubation, the pH was increased by approximately 0.05 pH units in the (B) and (C) incubations compared to the control (A). This relative pH increase with increasing additions of Mn-oxides is consistent with increasing acid consumption following Mn-reduction. Indicated are average and standard deviation of one to three replicate sediment samples.](image-url)
Fig. 12. Manganese in the pore water during the November closed jar incubation. Following the initial phase of Mn$^{2+}$ production, net Mn$^{2+}$ build-up slowed down as reactants became limiting and consumption reactions were initiated (e.g. the formation of Mn-carbonates). Dashed lines represent initial mobilizations used to estimate net Mn$^{2+}$ production rates.

Fig. 13. Nitrate was found to be produced anoxically in all plugs, despite that no nitrate had been detected in the overlying water for more than five days. Calculated anoxic nitrification rates were 1090, 195, 47.9 and 8.97 μmol dm$^{-3}$ d$^{-1}$ for the 2, 5, 10 and 20 mm plugs, respectively.
of denitrification (e.g., Devol, 1991). Most likely, these types of reactions are of importance in nearshore sediments where anaerobic sediments are more or less constantly mixed with Mn-oxides from the oxidized surface layers, and in organic-rich, hemipelagic sediments close to hydrothermal vent sites. Luther et al. (1997) suggested that the direct formation of N₂ through the oxidation of organic-N and ammonium by Mn-oxides (Table 1) was probably the most important process for N₂ formation in Mn-rich sediments. However, it is clear from the present results that lithotrophic nitrification may be an intermediate step in eventual denitrification, and that the relative importance of the myriad possible pathways remains to be demonstrated in the field.

The documentation of an apparent lithotrophic nitrification pathway emphasizes the potential complex redox reaction web that can occur in physically and biologically mixed surface sediments. Previous experiments have shown that the reworking of Mn-oxides into anoxic sediments can be coupled to the biogenic oxidation of labile sulfides (HS⁻, FeS) to S⁰ and SO₄²⁻ (Burdige and Nealson, 1986; Aller and Rude, 1988; King, 1988). Overall reactions such as (4), (abiogenic; Burdige and Nealson, 1986) and (5), (biogenic, Aller and Rude, 1988; King, 1988) were proposed.

\[
\begin{align*}
3 \text{H}^+ + \text{MnO}_2 + \text{HS}^- & \rightarrow \text{Mn}^{2+} + \text{S}^0 + 2 \text{H}_2\text{O} \quad (4) \\
8 \text{H}^+ + 4 \text{MnO}_2 + \text{FeS} & \rightarrow 4 \text{Mn}^{2+} + \text{SO}_4^{2-} + \text{Fe}^{2+} + 4 \text{H}_2\text{O} \quad (5)
\end{align*}
\]

The apparent occurrence of lithotrophic nitrification suggests that additional coupled reactions, such as a combination of reactions (2) and (5), are likely to occur in parallel (6) or in series (7), resulting in overall sedimentary redox reactions such as:

\[
8 \text{NO}_3^- + 5 \text{FeS} + 8 \text{H}^+ \rightarrow 4 \text{N}_2 + 5 \text{SO}_4^{2-} + 5 \text{Fe}^{2+} + 4 \text{H}_2\text{O} \quad (6)
\]
By utilizing the nitrate formed during anoxic nitrification, bacteria such as *Thiobacillus denitrificans* could generate sulfate anoxically in a consortium with lithotrophic nitrifiers (e.g., Baalsrud and Baalsrud, 1954). These reactions are sufficiently energetic to support chemoaerotrophic C fixation ($\Delta G_{\text{r}}$'s in Table 1). We note that the lack of net SO$_4^{2-}$ depletion and the hint of net SO$_3^{2-}$ production during the initial stages of our incubation experiments (e.g., Nov.) suggest the occurrence of such sulfide oxidation reactions simultaneously with net NO$_3^-$ production (Fig. 10). In summary, a significant consequence of biogenic and physical reworking of Mn-oxides into underlying anoxic sediment is likely to be the serial redox reaction coupling of O$_2$, Mn, N, Fe, S, and C by opportunistic microbial consortia (Fig. 18).

Although we have emphasized redox interactions between the major oxidants, there may be additional effects on heterotrophic processes that result from nonsteady-steady introduction of Mn-oxide into anoxic sediments. Initially, the presence of Mn-oxide and produced NO$_3^-$ seemed to inhibit reduction of pore water sulfate, due either to competitive utilization of higher energy oxidants (Lovley and Phillips, 1986), or the coupled redox reactions outlined previously. Following the initial period of no net sulfate reduction, the rates of sulfate reduction that eventually occurred in the Mn-oxide amended incubations (Nov.) were modestly stimulated in direct proportion with the quantity of Mn-oxide added (Fig. 10B, Table 2). Assuming the proportional increases are real, such stimulation could result from 1) degradation of microbial biomass generated during the initial pulse introduction of the energetically favorable Mn-oxide; (2) degradation of partially oxidized degradation products formed during the period of enhanced Mn-reduction; and (3) removal and immobilization of inhibitory metabolites by adsorptive uptake onto the relatively fresh oxide surfaces.

\[ 5 \text{FeS} + 32 \text{MnO}_2 + 8 \text{NH}_4^+ + 56 \text{H}^+ \rightarrow 4 \text{N}_2 + 5 \text{SO}_4^{2-} + 5 \text{Fe}^{3+} + 32 \text{Mn}^{2+} + 44 \text{H}_2\text{O} \]  

Fig. 16. A. Pore water concentrations of manganese in the 2, 5, 10 and 20 mm plugs after terminating the open plug experiment. The concentration gradients between the overlying water and the pore water indicated a diffusive transport of manganese from the plugs towards the overlying water. This gradient increased with increasing plug thickness and the calculated manganese production rates for the plugs were n.d., 42, 89, and 72 \(\mu\text{mol dm}^{-3} \text{d}^{-1}\), respectively. B. Concentrations of manganese in the solid phase of the sediment. The pore water and solid phase distributions of manganese indicated that manganese reduction was occurring in the sediment. Clearly, the contents of reactive MnO$_2$ in the sediment were sufficient to support Mn reduction and anoxic nitrification in all plugs over the incubation period. Net production rates of Mn$^{2+}$, estimated from loss of reactive Mn-oxide, were 384, 363, 348 and 230 \(\mu\text{mol dm}^{-3} \text{d}^{-1}\) for the 2, 5, 10 and 20 mm plugs, respectively. Indicated are average together with maximum and minimum concentrations of duplicate sediment samples.

Fig. 17. One possible explanation why nitrate could be maintained for a more extended period and at greater concentrations in the open anoxic, but not in the closed anoxic incubations, may be due to elevated pH gradients close to the sediment-water interface in the diffusively open plugs. The pH dependent reaction of anoxic Mn$^{2+}$ oxidation (5 Mn$^{2+}$ + 2 NO$_3^-$ + 4 H$_2$O $\rightarrow$ 5 MnO$_2$ + N$_2$ + 8 H$^+$) may occur more favorable in surficial regions of the sediment or in the overlying water reservoir, where the pH is higher. An internal cycle could thus be sustained where a dissolved oxidant (chelated Mn$^{3+}$ ?) in the surface zone could be supplied to the underlying anoxic region with lower pH.
4. CONCLUSIONS

Introduction of Mn-oxides from oxidized surface sediment into underlying anoxic regions during biogenic or physical mixing results in anoxic nitrification and net production of NO$_3^-$ and NO$_2^-$.

Anoxic nitrification rates are proportional to the quantity of Mn-oxide available in the solid phase. Anoxic nitrification occurs simultaneously with Mn-oxide reduction, implying the likely microbial coupling of the two reactions through an anaerobic pathway of lithotrophic nitrification.

Because of the multiple possible reduction reactions for NO$_3^-$ under anoxic conditions ($\text{C}_x\text{H}_y\text{O}_z\text{N}_m\text{MeHPO}_4$, $\text{Fe}^{2+}$, $\text{H}_2\text{S}$), higher net NO$_3^-$ production is observed in diffusively open versus closed systems having greater metabolite concentrations.

The juxtaposition of pulsed inputs of Mn-oxides with reduced constituents in anoxic deposits makes likely the close coupling of the O$_2$, Mn, N, S, and C cycles in a complex redox web rather than a simple succession in surface sediments.

Such redox coupling are likely to be of greatest importance in nearshore sediments with active biological and physical reworking, and in regions near hydrothermal vents where oxidized metals frequently contact reduced solutes and organic-rich deposits.

**Acknowledgments**—S. Dunn, M. Green, A. Ingalls and M-Y Sun provided help during part of the field sampling. This study was supported
REFERENCES


by NSF grant OCE 9730933 to R. C. Aller, and by post doctoral fellowships from the Swedish Natural Science Research Council and the Swedish Institute to S. Hulth, and grants from the Societe de Secours des Amis de la Science and the GDR (1123) “HYCAR” to F. Gilbert.