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How soil characteristics and water quality influence the biogeochemical response to flooding in riverine wetlands

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Abstract Although phosphate concentrations have been reduced, the rivers Meuse and Rhine are still polluted with sulphate, which most probably affects vegetation development in newly created riverine wetlands. The influence of flooding with river water rich in sulphate was tested on three soil types from floodplains of the river Meuse using flow-through and batch experiments. Soils were selected for contrasting concentrations of iron and organic matter and originated from a floating fen (iron-poor, organic), an alder carr (iron-rich, organic) and a clay pit (iron-rich, low in organic matter). Flooding induced mobilisation of phosphate. Sulphate only enhanced this effect in the alder carr soil, where sulphide and phosphate competed for binding to iron. Only in the floating fen soil did the addition of sulphate result in the formation of free sulphide, which reduced the growth of Glyceria maxima, serving as a phytometer. In addition, the floating soil started to sink, due to falling methane concentrations. In the different soil types methane production was hampered by the presence of more favourable electron acceptors such as sulphate in the water and Fe(III) in the soil. It was concluded that the effects of inundation with sulphate-polluted water strongly depend on the soil type: under iron-poor circumstances, free sulphide may accumulate, leading to phytotoxicity, while in soils rich in iron, sulphide toxicity is prevented, but phosphate availability may be increased. In addition, shortage of easily degradable organic matter can limit the formation of potential toxicants such as ammonium, iron and sulphide. Results are discussed in terms of their implications for nature management.

Keywords Buoyancy • Eutrophication • Flooding • Floodplain • Methane • Sulphate • Water storage

Introduction

Although great efforts have been made since the 1970s to improve water quality in the rivers Meuse and Rhine, concentrations of many major ions are still elevated compared to their natural background values. Measures such as the introduction of wastewater treatment plants have effectively reduced the concentrations of ammonium and phosphorus in the river water, and decreased discharge from the industry and decreased atmospheric sulphur deposition have contributed to a decrease in concentrations of sodium, chloride and sulphate, respectively. However, today’s average sulphate concentrations in the river Meuse still lie around 600 µmol l⁻¹ (Waterstat 2006), whereas measured historical concentrations...
(before 1900) and calculated background concentrations for the Meuse and Rhine in the Netherlands are approximately 300 μmol l⁻¹. Concentrations of nitrate (around 50 μmol l⁻¹) are also still elevated compared to their background values (5 μmol l⁻¹) (Van der Weijden and Middelburg 1989; Van den Brink et al. 1991; Van den Brink 1994). This article focuses on the effects of the elevated sulphate concentrations.

It is known that sulphate pollution can play an important role in the deterioration of freshwater wetlands, especially in peatlands (Roelofs 1991; Smolders et al. 1996, 2003; Lamers et al. 1998, 1999, 2001, 2002a, b; Lucassen et al. 2004). Under anaerobic circumstances, for instance during flooding, sulphate is reduced to sulphide in the absence of more favourable electron acceptors such as oxygen or nitrate. Free sulphide can be very toxic to wetland plant species, as has for instance been shown for *Stratiotes aloides* (Smolders and Roelofs 1996) and *Caltha palustris* (Van der Welle et al. 2007). However, sulphide has a high affinity for binding to iron, which decreases the availability of free sulphide and hence its toxicity (Van der Welle et al. 2006). As the affinity to iron is higher for sulphide than for phosphate, the latter can be released during this process, causing eutrophication (Caraco et al. 1989; Roden and Edmonds 1997; Lamers et al. 1998). This mobilisation of phosphate is added to the release of phosphate that takes place upon reduction of iron after the flooding of semi-terrestrial wetlands, such as floodplains. After inundation and the depletion of oxygen from the pore water, iron can be used as an electron acceptor for the oxidation of organic matter. This causes a partial release of the phosphate that was bound to iron(hydro)oxides under aerobic circumstances (Patrick and Khalid 1974). It is, however, unclear if sulphate has similar detrimental effects on floodplain biogeochemistry.

Besides the phytotoxic and eutrophying effects of sulphate after reduction, sulphate is also known to inhibit bacterial methane production (Cappenberg 1974; Abram and Nedwell 1978; Lovley and Klug 1983; Westermann and Ahring 1987; Prieme 1994; Scholten and Stams 1995). It has been demonstrated that as a result of this, floating *Sphagnum* bog rafts, which depend on methane for floatation, sink under the influence of increased sulphate concentrations (Lamers et al. 1999; Smolders et al. 2002). As floating fen mats present in riverine systems also depend on methane for buoyancy (Hogg and Wein 1988a; Swarzenski et al. 1991; Sasser et al. 1996), a similar effect of elevated sulphate concentrations can be assumed to exist, although this has never been reported. Floating mats form an important phase in terrestrialisation and hence in vegetation succession of peatlands (Tallis 1973; Beltman et al. 1996). They can be found in river deltas across the world, such as in the rivers Mississippi, Danube, Nile and Amazon (Hogg and Wein 1988b; Sasser et al. 1996).

During the last decade, many new wetlands have been created in the floodplains of large rivers, including the rivers Rhine and Meuse, either for the purpose of ecological restoration, or for flood risk reduction, or for a combination of both objectives. Plans for riverine wetland creation often ignore river water quality and soil quality, although it may be hypothesised that the effects of sulphate pollution that have been found in peatlands may also play an important role in riverine floodplains. Since concentrations of iron and of easily degradable organic matter are expected to act as key factors differentiating the effects of sulphate pollution, we chose to examine three different floodplain soils from the river Meuse, differing in iron and organic matter contents.

### Materials and methods

#### Site description

Three different floodplain areas along the river Meuse in the Netherlands were selected for their soil characteristics. Figure 1 depicts the locations of the sites. The first soil (F) was taken from a floating fen at the bank of an old oxbow lake (the ‘Oude Maas’) anchored to the river dike (N 47°58′43.04″ E 3°18′56.38″). This lake discharges its water into the Meuse and only comes into direct contact with the river water during floods. The soil consists mostly of roots and plant litter. The vegetation type it supports, which is dominated by the fern *Thelypteris palustris*, has become very rare in Dutch floodplains. A vegetation relevé from 1954 shows that at that time, the oxbow lake was partly vegetated by *Stratiotes aloides* (Van Donselaar 1961). The second soil (C) was taken from the shores of a recently dug clay pit (N 47°58′41.42″ E 3°18′58.26″) in the
Romeinenweerd nature reserve. In this area, several small lakes have been excavated for the purpose of nature restoration and flood risk reduction. The excavated clay was used for the construction of flood protection systems in neighbouring towns. The lake from which we collected the soil, is a stagnant water body, which suffered severe eutrophication in the hot summer of 2003, when a bloom of cyanobacteria led to high mortality of fish and waterfowl (Pollux and Pollux 2004). No vegetation was present at the spot from which we took the soil. The third soil (A) was taken from a black alder carr (N 47 58'42.27" E 3 18'57.98"), called Heuloerbroek, within the winter bed of the river Meuse. The site receives discharge of iron-rich groundwater from higher situated areas. The top soil consists of litter, mainly from Alnus glutinosa.

Soil characteristics are summarised in Table 1. The clay soil was poor in organic matter (about 5%), while the other two soils were mostly organic. Both the alder carr and clay soils had a high concentration of iron, but the oxalate-extractable pool, which is regarded as the amorphous iron that is available for bacterial reduction, was much lower in the clay soil, both in absolute and in relative terms. The iron concentration of the floating fen soil was very low. Both total P and Olsen-extractable P were high in the alder carr soil.

Experimental set-up

Aquarium experiment

Twenty-four 9-l aquaria were filled with a 10-cm layer (1/3 of the aquarium) of carefully mixed soil (four replicates) (Fig. 2) and kept in a water basin at a temperature of 20°C in a climate chamber at a light intensity of 200 μmol m⁻² s⁻¹ (Hortilux Schreder HPS 400 W) for 12 h a day. Two 10-cm pore water samplers (Rhizon SMS -10 cm; Eijkelkamp Agrisearch Equipment, The Netherlands) were placed diagonally in the soil of each aquarium. A 5-10 cm layer of surface water collected in the field was added on top of the soil. After 1 week, the soils were further inundated with artificial river water containing either 1 mmol l⁻¹ SO₄ (as Na₂SO₄) or no SO₄ (Table 2), from 10 l stocks, each connected to their own aquarium. The medium was pumped through the aquaria at a speed of 10 l per week. Three shoots of Glyceria maxima and 0.25 g of Lemna minor, collected in the field, were added to each aquarium as phytometers. Other plant species than G. maxima, L. minor or algae were carefully removed from the aquaria. Pore water samples were taken by connecting 100-ml vacuum bottles to the rhizon samplers. Samples of both pore water samplers were pooled per aquarium to diminish the effects of heterogeneity. Surface water samples were filtered (Whatman GF/C). After 12 weeks, the aboveground biomass of G. maxima and the floating L. minor and algae biomass were harvested. In the aquaria with the alder carr soil, algae were harvested more often, to prevent die-off of G. maxima from light limitation.

Decomposition experiment

Fifty g portions of fresh soil were transferred to 350-ml glass bottles. The bottles were filled up to 170 ml with the same medium as used in the aquarium experiment and preflushed with N₂ to remove O₂ from the water. The headspace of the bottles was also flushed with N₂ before they were sealed with airtight
rubber stoppers. We will refer to this soil and water suspension as slurry. Analyses were done in quadruplicate. Initial concentrations in the slurry water were determined by filling smaller bottles with 25 g of fresh soil and filling them up to 85 ml with medium. These bottles were shaken overnight. Slurry water samples were taken by pore water samplers (Rhizon SMS-5 cm; Eijkelkamp Agrisearch Equipment, The Netherlands). Decomposition bottles were shaken in the dark at 20°C. Samples from the headspace were taken twice a week with 1-ml syringes and were analysed instantly for CH₄ and CO₂. After 42 days, slurry water samples were taken and the experiment was ended. Potential production rates were calculated by linear regression of the net production in the linear phase of the production rate of CH₄ or CO₂. Potential decomposition rates were calculated in the same way, expressed as the C production rates in the linear phase of the concentration build-up of the sum of CH₄ and CO₂. This does not equal the sum of the potential CH₄ and CO₂ production rates, because of the difference in lag phase. Potential net release rates were calculated from the difference between the pore water concentrations at the start and the end of the experiment, divided by the time elapsed.

**Chemical analyses**

Immediately after collection of the pore water, the concentration of free sulphide was determined in a 10 ml subsample, fixed with S²⁻ antioxidant buffer containing NaOH, NaEDTA and ascorbic acid (Van Gemerden 1984). S²⁻ was detected using an S²⁻ ion-specific Ag electrode and a double junction calomel reference electrode (Thermo Orion) (Van Gemerden 1984). pH was measured with a radiometer Copenhagen type PHM 82 standard pH meter. Alkalinity was determined by titrating 10 ml of sample with 0.01 mmol l⁻¹ HCl down to pH 4.2. For the ICP analyses, 1.7% of concentrated nitric acid was added to the samples to keep the oxidised iron in solution. Samples were stored in polypropylene tubes at 4°C. For the other analyses, 0.12 g citric acid/l was added to the samples. They were stored in iodated polyethylene bottles at −24°C until further analysis. Total concentrations of Ca, Mg, P, Fe and S were analysed by ICP-MS (X series, Thermo Electron Corporation.}

**Table 1** Soil characteristics

| Soil        | Organic matter content (%) | Oxalate-extractable Fe (µmol g⁻¹ dw) | Total Fe (µmol g⁻¹ dw) | P-Olsen (µmol g⁻¹ dw) | Total P (µmol g⁻¹ dw) |
|-------------|-----------------------------|--------------------------------------|------------------------|-----------------------|------------------------|
| Floating fen| 83                          | 50                                   | 100                    | 3                     | 30                     |
| Clay pit    | 5                           | 104                                  | 610                    | 1                     | 24                     |
| Alder carr  | 52                          | 950                                  | 1011                   | 5                     | 104                    |

**Table 2** Concentrations of macro-ions in the artificial river water treatments, with and without sulphate

|                | Medium −SO₄ (µmol l⁻¹) | Medium + SO₄ (µmol l⁻¹) |
|----------------|------------------------|------------------------|
| Na             | 2,000                  | 4,000                  |
| Ca             | 1,300                  | 1,300                  |
| Mg             | 250                    | 250                    |
| Cl             | 3,100                  | 3,100                  |
| HCO₃           | 2,000                  | 2,000                  |
| SO₄            | 0                      | 1,000                  |

Fig. 2 Experimental set-up. Picture: Lidwien van der Horst
and ICP-OES (IRIS Intrepid II XDL, Thermo Electron Corporation). Total Fe measured in the pore water was assumed to be Fe(II), since this is the dominant soluble species at the pH values we measured. Total S was assumed to be SO₄ (if no sulphide was present), as, at the concentrations occurring in our soils, only a very small part will be in organic form. This was confirmed by parallel SO₄ analysis of part of the samples with methylthymol blue and BaCl₂ (Bran + Luebbe, AutoAnalyser 3, Digital Colorimeter), o-PO₄ (Technicon AutoAnalyser II), NH₄⁺, NO₃⁻ (+NO₂⁻) and Cl⁻ (Bran + Luebbe AutoAnalyser 3) were analysed colorimetrically using ammonium molybdate, salicylate, hydrazine sulphate and ferriammonium sulphate, respectively. CO₂ and CH₄ were measured by infrared analysis (ABB Advance Optima Infrared Gas Analyser).

In order to determine the total concentrations of elements in plant and soil materials, 200 mg of homogenised and dried sample was digested for 17 min with 4 ml concentrated HNO₃ and 1 ml 30% H₂O₂ (Milestone microwave MLS 1200 Mega). C and N concentrations in the plant material (20 mg) were determined with a CNS analyser (Carbo Erla Instruments NA 1500).

Organic matter content of the soils was determined by loss-on-ignition (4 h, 550°C). Soil extractions were performed on fresh soils and corrected for moisture content afterwards, after drying (24 h, 105°C). The concentration of amorphous Fe, which is supposed to represent the fraction available for microbial Fe reduction, was determined by oxalate extraction (2.5 g of soil shaken for 2 h with 30 ml of a solution containing 16.2 g of (COONH₄)₂H₂O and 10.9 g of (COOH₂)₂H₂O per l) (Schwertmann 1964) and plant-available P by bicarbonate extraction (5 g of soil shaken with 100 ml of 0.5 M NaHCO₃) (Olsen et al. 1954).

Statistics

All concentrations were log (+1) transformed to improve their fit to a normal distribution, except for floatation measurements. All statistics were calculated with SPSS 13.0 (SPSS Inc., 2004). Effects of time series were tested with GLM repeated measures, using Greenhouse-Geisser corrections if terms of sphericity were not met in Mauchly's sphericity test. Effects of sulphate addition on the specific soil types were assessed with an independent samples T-test using data of the last measurement, unless stated otherwise. Effects of sulphate on concentrations and production rates of methane and on iron release were tested with 1-tailed tests, because the direction of the effect was already known. Differences between soil types were assessed by univariate ANOVA. Decomposition rates were calculated over the linear range of CO₂ or CH₄ production by linear regression. Significance was accepted at a confidence level of \( P \leq 0.05 \).

Results

Aquarium experiment

Pore water and surface water

Table 3 summarises the results of the repeated measures analyses of the concentrations of elements and compounds in the pore water and surface water. From this table it can be seen that concentrations changed over time for all elements we measured (time effect). These changes were different for the different soil types, as indicated by the significant time × soil interactions. Major changes in concentrations are discussed below. Initial concentrations of all measured elements and ions are given in Table 4.

The artificial river water replaced the original surface water in the aquarium within 2 weeks, as was indicated by the Cl⁻ concentration (results not shown), which can be used as an inert tracer. Within 1 month, the river water had penetrated all soil types, even the poorly permeable heavy clay soil, as can be seen from the Cl⁻ concentrations in the pore water (Fig. 3a).

The presence of high concentrations of soluble Fe in the pore water of the alder carr and clay soils right from the start of the experiment (Fig. 3b) showed that conditions were anoxic from the beginning. Concentrations of Fe in the pore water correlated with those of amorphous iron in the soil determined by oxalate extraction. In the floating fen soil, hardly any iron was present in the pore water. In the clay soil, the Fe concentration in the pore water stabilised at 400 μmol l⁻¹. In the alder carr soil, concentrations were extremely high, varying between 1,500 and 2,500 μmol l⁻¹. Concentrations of Fe in the surface water remained low, except for the alder carr soil
Table 3  P values of the time effects (time), time × soil interactions (time × soil), time × treatment interactions (time × treat) and time × soil×treatment interactions (time × soil × treat) of the pore water and surface water concentrations in the aquarium experiment, as tested by repeated measures (GLM)

|                     | Time | Time × Soil | Time × Treat | Time × Soil × Treat |
|---------------------|------|-------------|--------------|---------------------|
| P                   | 0.00 | 0.00        | 0.55         | 0.83                |
| Fe                  | 0.00 | 0.00        | 0.01         | 0.00                |
| Ca                  | 0.01 | 0.00        | 0.59         | 0.36                |
| NH₄                 | 0.00 | 0.00        | 0.35         | 0.08                |
| Cl                  | 0.00 | 0.00        | 0.69         | 0.31                |
| Na                  | 0.00 | 0.00        | 0.00         | 0.12                |
| K                   | 0.00 | 0.00        | 0.83         | 0.96                |
| SO₄                 | 0.00 | 0.00        | 0.00         | 0.00                |
| PO₄                 | 0.00 | 0.00        | 0.25         | 0.48                |
| Mn                  | 0.00 | 0.00        | 0.07         | 0.27                |
| Mg                  | 0.01 | 0.00        | 0.57         | 0.56                |
| pH                  | 0.00 | 0.00        | 0.04         | 0.27                |
| Alkalinity          | 0.00 | 0.00        | 0.03         | 0.42                |

|                     | Time | Time × Soil | Time × Treat | Time × Soil × Treat |
|---------------------|------|-------------|--------------|---------------------|
| P                   | 0.00 | 0.00        | 0.17         | 0.06                |
| Fe                  | 0.00 | 0.00        | 0.06         | 0.39                |
| Ca                  | 0.00 | 0.00        | 0.37         | 0.32                |
| NH₄                 | 0.00 | 0.00        | 0.00         | 0.02                |
| Cl                  | 0.00 | 0.03        | 0.40         | 0.65                |
| Na                  | 0.00 | 0.00        | 0.00         | 0.32                |
| K                   | 0.00 | 0.00        | 0.32         | 0.08                |
| SO₄                 | 0.00 | 0.00        | 0.00         | 0.24                |
| PO₄                 | 0.00 | 0.00        | 0.03         | 0.01                |
| Mn                  | 0.00 | 0.02        | 0.00         | 0.06                |
| Mg                  | 0.00 | 0.00        | 0.67         | 0.25                |
| pH                  | 0.00 | 0.00        | 0.06         | 0.12                |
| Alkalinity          | 0.00 | 0.00        | 0.01         | 0.47                |

Bold values indicate P ≤ 0.05

(Fig. 3c). Only in the iron-poor floating fen soil did sulphate addition lower the availability of iron significantly. At the end of the experiment, iron concentrations in the soil pore water in the floating fen soil were 4.5 μmol l⁻¹ on average in the treatment without sulphate and 0.7 μmol l⁻¹ in the treatment with sulphate.

Sulphate reduction took place even in the early stages of the experiment. The sulphate treatment caused sulphate concentrations (Fig. 3d) in the pore water in the iron-rich soils (C and A) to decrease and remain well below the added concentration of 1,000 μmol l⁻¹ after 1 month, even though Cl concentrations showed that the sulphate-rich medium had fully penetrated the soil. Initial sulphate concentrations were high for the clay soil, but fell considerably within the first month. The sulphide concentration in the pore water after 28 weeks was below the detection limit for all soils and treatments, except for the sulphate-treated floating fen soil (275 μmol l⁻¹; results not shown).

Starting from week 29, the distance between the floating sods and the bottom of the aquarium became significantly smaller in the sulphate-treated condition (Figs. 4 and 5). Sulphate addition had a significant influence on the CH₄ concentrations of the pore water in the floating fen soil. Average pore water concentration in the control was 1,700 μmol CH₄ l⁻¹ pore water, against 370 μmol l⁻¹ for the sunken sods in the sulphate treatment (results not shown).
Initial phosphate concentrations in the pore water were high for the fen soil (Fig. 3e), but decreased strongly within the first 2 weeks. Phosphate concentrations in the surface water of all soils (results not shown) also showed a steep drop at the beginning of the experiment, due to leaching. Concentrations of total P (results not shown) showed the same pattern as phosphate concentrations. Phosphate concentrations in the pore water became high especially in the alder carr soil after 10 weeks (13–20 μmol PO₄ P⁻¹ (Fig. 3e), 17–42 μmol total P⁻¹), compared to the other soil types. Sulphate addition clearly increased the phosphate availability in the pore water of the iron-rich alder carr soil, unlike the other soils. Phosphate concentrations in the pore water of this soil upon treatment with sulphate were significantly higher than in the control (weeks 6-20; about twice as high in week 20). In the surface water, however, this effect of sulphate on phosphate concentrations was only visible as a transient peak in week 20, when surface water concentrations (31 μmol P⁻¹) were 10 times higher in the sulphate-treated condition (results not shown).

NO₃⁻ was present in very low concentrations from the start of the experiment (results not shown). Within 2 weeks, it dropped to below 5 μmol L⁻¹ in both the pore water and the surface water, either by reduction under anoxic circumstances, or by leaching. During the experiment, concentrations of NH₄⁺ in the pore water (Fig. 3f) were high in the clay and the floating fen soil (10–100 μmol L⁻¹), and extremely high in the alder carr soil (up to 1,600 μmol L⁻¹). As the reduction of Fe and that of SO₄ are acid consuming processes, the pH in the pore water (Fig. 3g) rose in all soils during the experiment, the increase being greater for the sulphate treatment (time × treatment interaction). pH also increased in the surface water during the experiment (results not shown). During algal blooms, pH rose above 9 in the surface water of the alder carr soil, as a result of CO₂ depletion by photosynthesis.

**Biomass of the phytometers**

In the floating fen soil, the addition of sulphate had a negative effect on *G. maxima* biomass production. Here, the addition reduced the biomass by more than 60% compared to the control (results not shown). In the other soils, however, standard deviations of the
Fig. 3 Concentrations of ions in the pore water and the surface water in the aquarium experiments. Open symbols represent treatments without sulphate, closed symbols treatments with added sulphate. Squares: floating fen soil (F); triangles: clay soil (C); circles: alder carr soil (A). Error bars represent standard error of the mean. (a) Concentration of Cl in the pore water; (b) concentration of Fe in the pore water; (c) concentration of Fe in the surface water; (d) concentration of total S in the pore water; (e) concentration of PO$_4$ in the pore water; (f) concentration of NH$_4$ in the pore water; (g) pH of the pore water.
Decomposition experiment

Potential decomposition

Potential decomposition rates (Fig. 6, total C) were highest in the floating fen soil. No significant effect of sulphate addition on the potential decomposition rates was found. When corrected for the organic matter content of the soils, decomposition rates in the floating fen soil were four times higher than the rates in the alder carr soil.

Potential production of CH₄

The potential methane production differed considerably between the three soil types. Hardly any methane was produced in the clay soil. In contrast, the floating fen soil had a very high potential methane production rate (about 9 μmol g⁻¹ dry weight day⁻¹; Fig. 6), which was four times higher than the potential methane production rate in the alder carr slurry. Univariate ANOVA showed that the two organic soils also differed in the duration of the lag phase, i.e. the initial phase in which no methane is produced (4 h for the floating fen soil and 215 h for the alder carr soil). A significant effect of the addition of sulphate on the methane production rates was also demonstrated by univariate ANOVA (Table 5), showing the inhibiting effects of sulphate as a more favourable electron acceptor.
Table 5  \( P \) values of the soil and treatment effects of the CO\(_2\), CH\(_4\) and total C (CO\(_2\) + CH\(_4\)) production rates expressed per gram dry weight, as tested by univariate ANOVA (GLM)

|          | CO\(_2\) production rate | CH\(_4\) production rate | Total C production rate |
|----------|---------------------------|---------------------------|-------------------------|
| soil     | 0.000                      | 0.000                      | 0.000                   |
| treatment| 0.357                      | 0.021                      | 0.605                   |
| soil \( \times \) treatment | 0.537                      | 0.138                      | 0.043                   |

Bold values indicate \( P \leq 0.05 \)

**Potential production of CO\(_2\)**

Like that of CH\(_4\), the potential production rate of CO\(_2\) in the floating fen slurry was very high (28 \( \mu \)mol g\(^{-1}\) dry weight d\(^{-1}\); Fig. 6), four times higher than the potential production rate in the alder carr slurry. In the clay slurry, CO\(_2\) production was measured in the first 6 days of the experiment, but the production stopped after that time. Although the potential production rate expressed per gram dry weight was three times smaller in this clay slurry than in the alder carr slurry, when expressed as potential production rate per gram organic matter, the potential CO\(_2\) production rate in the clay slurry during these six days was even higher than in that of the alder carr.

**Potential release rates**

Figure 7a–d shows the net potential release rates for PO\(_4\), NH\(_4\), K and Fe. Both of the organic soils (F and A), had high net PO\(_4\), NH\(_4\) and K release rates. Net Fe release was highest in the alder carr slurry, analogous to the aquarium experiment. The ratio between the net release of C and NH\(_4\) in the slurry was three times higher in the floating fen slurry than in the alder carr soil. In the alder carr slurry treated with sulphate, phosphate release was three times higher \((P = 0.021)\) than in the control. This is comparable to the results of the aquarium experiment. In the iron-poor floating fen soil, the iron release rate was significantly reduced by the addition of sulphate.

**Discussion**

The three soil types showed large differences in potential anaerobic decomposition rates and release rates. In the clay soil, CO\(_2\) production stopped after 6 days. The potential decomposition rate per gram of organic matter in these first days was, however, even higher in this soil than in the alder carr soil. This means that during the first 6 days, easily degradable organic matter was present in the clay soil. Most likely, after a few days the concentration of degradable organic matter had become limiting for decomposition in the clay soil. The potential decomposition rate of the floating fen soil was very high (40 \( \mu \)mol C g\(^{-1}\) dw day\(^{-1}\)). Maximum values found in the literature vary between 10 \( \mu \)mol C g\(^{-1}\) dw day\(^{-1}\) for bog peat (Smolders et al. 2002) and 50 \( \mu \)mol C g\(^{-1}\) dw day\(^{-1}\), reported for the floating fen in Lake Emlagharan in Ireland (Bontes et al. 2005). This means that the organic matter in the floating fen soil was relatively easily degradable. Potential CH\(_4\) production rates also varied per soil type. Due to the lack of easily decomposable organic material, no CH\(_4\) was produced in the clay soil, i.e. there was a surplus of electron acceptors compared to the organic matter present. This was also the case in the alder carr soil during the lag phase. The high concentration of Fe(III) in this soil inhibited methanogenesis in this phase, because iron-reducing bacteria have a higher affinity to easily degradable organic compounds, such as acetate, than methane-producing bacteria (Roden and Wetzel 2003). The presence of a lag phase suggests that methanogenesis does not occur in the field, because of the continuous supply of sulphate.

Release rates were high for both of the organic soils. The alder carr soil released more iron, due to its higher amorphous iron concentration. The ratio between the net release of C and N in the alder carr slurry was, however, four times lower than in the floating fen soil, although NH\(_4\) release rates were equally high: this means that part of the ammonium released in the alder carr soil probably did not originate from the decomposition of organic matter, but from cation exchange. Cation exchange rates might be higher in the alder carr soil than in the other soils, because of the production of high concentrations of Fe\(^{2+}\) by the reduction of Fe\(^{3+}\).

In the aquarium experiment, iron concentrations in the pore water were extremely high in the alder carr soil. Lucassen et al. (2000) showed that field concentrations of 2,600 \( \mu \)mol Fe l\(^{-1}\) induce necrotic spots in leaves of Glyceria fluitans. Although no such
Fig. 7 Potential net release rates in the decomposition experiment. F = floating fen, C = clay soil, A = alder carr soil. − = control treatment without sulphate, + = treatment with sulphate. Error bars represent standard error of the mean. (a) Potential net release rate PO₄; (b) potential net release rate NH₄; (c) potential net release rate K; (d) potential net release rate Fe. Asterisks mark significant differences between treatments within a soil.

spots were found on Glyceria maxima in our experiment, concentrations found in our experiment (up to 2,500 μmol Fe l⁻¹) might still be toxic to sensitive plant species in the field. It is known that reduced iron from the soil pore water that diffuses towards the surface water is oxidised and precipitates on the soil-surface water interface due to the higher redox potential of the water column (Cornwell and Kipphut 1992; Gunnars and Blomqvist 1997; Moore et al. 1998). This explains the low iron concentrations in the surface water over the clay soil. In the alder carr soil, pore water concentrations were so high that exchange of Fe between the pore water and the surface water took place. Since the surface water was directly connected to the outflow of the aquarium, concentrations in the surface water can be regarded as outflow concentrations. The stability of the pore water iron concentrations and the high weekly iron outflow from the alder carr soil show that iron reduction was still taking place at the end of the experiment. Sulphate reduction rates were also high, indicating simultaneous activity of Fe— and SO₄— reducing micro-organisms (Sørensen 1982).

Iron reduction played a major role in the mobilisation of phosphate in this experiment. In the alder carr soil, which was rich in iron as well as in total and Olsen-extractable P, continuous iron reduction led to high concentrations of P in the pore water. In this P-rich soil, Fe(III) precipitations at the soil-water interface did not trap all P, and increasing P concentrations were also found in the surface water. In the iron-poor floating fen soil, on the other hand, most of the phosphate release might be attributed to decomposition of organic material. The decomposition experiment showed a high decomposition rate in this soil, as well as high concentrations of phosphate in the slurry at the end of the experiment. There was, however, a striking difference between the high phosphate release in the slurry of the floating fen in the incubation experiment and the low phosphate concentrations in the pore water of the aquarium experiment, whereas the P concentrations in the alder
carr were high in both experiments. One of the explanations for this apparent discrepancy might be the difference in phosphate leach-out between these soils. Pore water phosphate concentrations in the floating fen soil were high at the beginning of the aquarium experiment, but decreased very fast after the start of the experiment. Smolders et al. (2001) showed that the release of phosphate from the pore water of underwater soils to the water layer strongly depends on the Fe:PO₄ ratio in the pore water; below a value of 1 (mol mol⁻¹), they measured a strong release of PO₄ to the surface water. Fe:PO₄ ratios in our experiment were 0.3 (with sulphate) to 5 (without sulphate) for the floating fen soil and 82 (with sulphate) to 176 (without sulphate) for the alder carr soil. This may have caused a higher phosphate release to the water layer from the floating fen soil, and hence a larger leach-out.

Due to the anaerobic conditions, NH₄⁺ resulting from the decomposition of organic matter could not be nitrified, and accumulated in the pore water, particularly in the alder carr soil. Another possible source of the high NH₄⁺ concentrations in the pore water could be cation exchange of adsorbed NH₄⁺ by cations added with the artificial river water (Lucassen et al. 2006), and in the alder carr soil also by the mobilised Fe²⁺ originating from the reduction of Fe(III)oxides. In the alder carr soil, ammonium concentrations rose to 1,600 μmol ⁻¹. Lucassen et al. (2006) showed that the presence of target vegetation in alder carrs is correlated with low concentrations of ammonium (<200 μmol ⁻¹). Hence, the concentrations measured in this experiment, if present in the field, could seriously hamper the intended vegetation development.

The response of the different soils used in this study to inundation with sulphate-rich river water in the aquarium experiment varied substantially, reflecting their soil characteristics. Sulphide, produced by reduction of the added sulphate, was present in excess relative to iron in the floating fen soil. Hence, the binding of sulphide to iron reduced the availability of soluble iron in this soil, both in the aquarium and in the incubation experiment. As iron concentrations were low, phosphate release as a result of the binding of iron to sulphide was equally low. Part of the sulphide produced could not be bound to iron and was freely available in the pore water in toxic concentrations, hampering the growth of the planted Glyceria maxima. The sulphide concentrations we measured in the pore water have been shown to be toxic to several wetland plants (Smolders and Roelofs 1996; Van der Welle et al. 2006, 2007). Moreover, the addition of sulphate decreased the methane concentration in the floating fen soil in the aquarium experiment and the potential methane production rate in the incubation experiment. Methane concentrations determined the buoyancy of the floating fen, just as had been demonstrated earlier for other floating fens and peat bog rafts (Hogg and Wein 1988a; Swarzenki et al. 1991; Sasser et al. 1996; Lamers et al. 1999; Smolders et al. 2002). The floating fen soil sank under the influence of sulphate as a result of methanogenic bacteria being outcompeted by sulphate-reducing bacteria (Cappenberg 1974; Abram and Nedwell 1978; Lovley and Klug 1983; Westermann and Ahring 1987; Prieme 1994; Scholten and Stams 1995), in the same way as this is known to happen in floating peat bogs (Lamers et al. 1999; Smolders et al. 2002). In the other soils, sulphide formed by the reduction of sulphate was bound to the excess iron. In both the aquarium experiment and the incubation experiment, this led to an additional release of phosphate in the alder carr soil. Sulphate addition did not increase the decomposition rates, as had been found earlier in bog peat (Lamers et al. 1999; Smolders et al. 2002; Tomassen et al. 2003). In soils where sulphate increases decomposition, this seems to be an effect of the increased alkalinity produced by the reduction of sulphate (Lamers et al. 1999; Smolders et al. 2002; Tomassen et al. 2003). In our experiment, alkalinity of the artificial river water was already high (2,000 μmol HCO⁻1 1⁻¹), similar to concentrations in the water of the river Meuse). Since no effect of sulphate on the decomposition rate was found, it is likely that the additional release of phosphate in the sulphate treatment of the alder carr soil in the incubation experiment did not originate from decomposition of organic matter, but can be attributed to the competition between sulphide and phosphate for binding to iron.

Our findings show the importance of both soil type and water quality in the biogeochemical response to flooding. Inundation has the smallest impact on soils without organic matter, in terms of phosphate release by reduction of Fe (II) and sulphide (‘internal toxification’). Flooding of
soils containing organic matter, in contrast, leads to accumulation of ammonium and release of Fe-bound phosphate. This internal eutrophication could cause algal blooms in the water layer above such soils. Additionally, river water polluted with sulphate may cause sulphide phytotoxicity if the iron content is low, as was shown here for the floating fen soil. If the soil is richer in iron, sulphide will be bound to iron and may release phosphate that was bound to the iron, as took place in the alder carr soil. It should be noted that these two processes, sulphide toxicity and enhanced phosphate mobilisation, can also take place in the same soil, as was found for instance by Lamers et al. (1998) for a mesotrophic fen.

The presence of sulphate reduces methanogenesis, which is a very important process in ecosystems depending on methane for floating, such as floating fen mats and bog peat rafts. Floating fen mats form the first stage in the terrestrialisation process in fen systems. Our results show that inundation of these floating fens with sulphate-rich river water at levels occurring in the rivers Meuse and Rhine may thus destroy the floating fens and their rare vegetation as well as block the subsequent terrestrialisation process and the accompanying vegetation succession. Analogously, we expect the elevated nitrate concentrations in the river water to have a similar inhibiting effect on methanogenesis (Westermann and Ahring 1987; Scholten and Stams 1995) and on the buoyancy of floating fens.

The results presented in this article have important implications for the rehabilitation and creation of wetlands, and for the selection of locations for the creation of flood water storage areas. Completely mineral soils, such as those in clay and sand sedimentation areas where the topsoil has been removed, are not likely to be at great risk of eutrophication in the short term. However, wetland locations containing organic matter—and most soils do so—should be chosen with greater care. Here, other soil characteristics such as iron and phosphate concentrations, and water quality parameters such as sulphate concentrations, should be taken into account.

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