



# Wetland eco-engineering: measuring and modeling feedbacks of oxidation processes

# between plants and clay-rich material

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# Abstract

Interest is growing in using soft sediment as a building material in eco-engineering projects. Wetland construction in the Dutch lake Markermeer is an example: here the option of dredging some of the clay-rich lake-bed sediment and using it to construct wetland will soon go under construction. Natural processes will be utilized during and after construction to accelerate ecosystem development. Knowing that plants can eco-engineer their environment via positive





or negative biogeochemical plant-soil feedbacks, we conducted a six-month greenhouse experiment to identify the key biogeochemical processes in the mud when *Phragmites australis* is used as an eco-engineering species. We applied inverse biogeochemical modeling to link observed changes in pore water composition to biogeochemical processes. Two months after 25 transplantation we observed reduced plant growth and shriveling and yellowing of foliage. The N:P ratios of plant tissue were low and were affected not by hampered uptake of N but by enhanced uptake of P. Subsequent analyses revealed high Fe concentrations in the leaves and roots. Sulfate concentrations rose drastically in our experiment due to pyrite oxidation; as reduction of sulfate will decouple Fe-P in reducing conditions, we argue that plant-induced iron 30 toxicity hampered plant growth, forming a negative feedback loop, while simultaneously there was a positive feedback loop, as iron toxicity promotes P mobilization as a result of reduced conditions through root death, thereby stimulating plant growth and regeneration. Given these two feedback mechanisms, we propose that when building wetlands from these mud deposits Fe-tolerant species are used rather than species that thrive in N-limited conditions. The results 35 presented in this study demonstrate the importance of studying the biogeochemical properties of the building material and the feedback mechanisms between plant and soil prior to finalizing the design of the eco-engineering project.

40 **Keywords:** Drying; Fe-P; Iron toxicity; P mobilization; PHREEQC; Pyrite





### 1. Introduction

- Nowadays natural processes are being used across the world to achieve fast ecosystem 45 development while at the same time providing opportunities for developing hydraulic infrastructure, a concept called Building with Nature (BwN) (Temmerman et al., 2013). Though mostly focused on water safety and coastal protection (e.g. Borsje et al., 2011), BwN can also be applied for the management of fine sediments. A relevant application could be to use soft sediments as material for building freshwater wetlands. Here, vegetation can be used as an 50 eco-engineer (Jones et al., 1994), to modify the environment (Lambers et al., 2009). When fine sediments are used for the construction of wetlands, however, the use of eco-engineers is anticipated to pose challenges in relation to crest stability, consolidation and soil formation.
- In the Netherlands, a soft clay-rich lake-bed sediment in the Markermeer (an artificial lake of 691 km<sup>2</sup>) is causing serious turbidity problems: primary productivity is impeded and biodiversity 55 in the lake is declining (Vijverberg et al., 2011; Noordhuis et al., 2014). Because the lake is shallow, wind-induced waves frequently induce high bed shear stress, which causes sediment to be resuspended (Vijverberg et al., 2011). To improve the ecological conditions in the lake, it is planned to dredge some of the soft clay-rich sediment and use it to construct approximately 10,000 ha of wetland.
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Plants produce root exudates which influence soil formation by enhancing microbiological activity (Holtkamp et al., 2011), biological weathering and nutrient cycling (Taylor et al., 2009; Bradford et al., 2013). An example is the ability of plant roots to mobilize P by ligand exchange





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and dissolution of Fe-bound P (Fe-P) by citrate and oxalate excretion (Gerke et al., 2000). Plant
 roots may also enhance consolidation processes in substrate by increasing horizontal and
 vertical drainage (O'Kelly, 2006).

However, both negative and positive plant-soil feedbacks exist, in which the physical and chemical properties of the soil affect plant development and vice versa (Ehrenfeld et al., 2005). Therefore, when looking at soil formation it is important to study the sign and strength of these plant-soil feedback mechanisms. For example, nutrient conditions co-determine the type of plant community that develops (e.g. Olde Venterink, 2011), which in turn influences the nutrient conditions in the soil itself (Onipchenko et al., 2001). As feedback mechanisms differ between plant species (Ehrenfeld et al., 2005), in order to accelerate ecosystem development it is essential to decide which eco-engineer is best to introduce in these proto-soils.

De Lucas Pardo (2014) found that the Markermeer mud deposits had a high water content (20–60% of fresh weight) and were largely anoxic, with oxygen present only in the top 2 mm. Therefore, when such mud is taken from the lake and spread out in contact with the air, biogeochemical plant–soil processes related to oxidation and drying of the top soil are expected to play a significant role. It is intended to use two types of clay-rich deposits as building material for the wetland. They are the products of a combination of historical and present-day factors. Prior to 1932, the year in which the dam cutting off the Zuiderzee from the North Sea was completed, this was a marine environment into which several rivers discharged, including a branch of the river Rhine (the river IJssel). Hence, a near-shore marine deposit underlies the present-day soft clay-rich sediment. This soft clay-rich layer is produced by bioturbation and





physical weathering and continuously resuspends as a result of wave action (Van Kessel et al., 2008; De Lucas Pardo et al., 2013). This layer accumulated after 1976, when northward sediment transport was blocked by a second dam that separated Markermeer from IJsselmeer, thus allowing suspended matter to resettle on top of the marine deposit. We can therefore distinguish two layers: an upper disturbed mud layer prone to bioturbation and erosion, and a relatively undisturbed layer below.

With the aim of identifying the biogeochemical plant-soil feedback processes that occur when oxidation, drying and modification by plants alter the biogeochemical conditions of these two sediment types, thus in turn affecting vegetation development, we set up an experiment to monitor the chemical composition of pore water. Our study has two subsidiary aims: to ascertain how plants of *Phragmites australis* eco-engineer their environment by expediting biogeochemical processes in the deposits, and to simulate the geochemical differences between disturbed mud and undisturbed clay deposits and relate these to the processes identified from the pore water by using PHREEQC for inverse modeling. In addition, we altered the grain size of the disturbed mud deposit by adding inert sand to see how grain size distribution impacts pore water chemistry.





### 2. Material and Methods

2.1 Set-up

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A greenhouse experiment was conducted for six months at the test facility of Utrecht University. A basin of 4 m<sup>2</sup> (2 x 2 m) was filled with artificial rainwater and was refreshed every two weeks. At regular intervals, the chemistry of the water was checked to ensure that the water composition remained stable during the experiment. The artificial rainwater was made by adding 15 µmol NH<sub>4</sub>(SO<sub>4</sub>), 50 µmol NaNO<sub>3</sub> and 30 µmol NaCl to osmosis water. These values reflect the

average rainwater composition in the Netherlands for the period 2012–2013 (LMRe, 2014).

The sediments used include the soft clay-rich layer (Mud<sub>soft</sub>) and the underlying, consolidated, Zuiderzee deposit (Clay). In principle, both sediments have the same origin and were collected in the same area. We also included a third sediment type (Mud<sub>sand</sub>), as it is expected that Mud<sub>soft</sub> will be too soft for use as a building material: a 1:1 mixture was made by mixing mud with Dorsilit<sup>®</sup> crystal silica sand (c. 99% SiO<sub>2</sub>) which had been autoclaved for one hour at 120 °C prior to mixing. The Mud<sub>soft</sub> and Clay sediments were collected by mechanically dredging in the southern part of the lake and were stored in air-tight containers at 4 °C prior to the start of the experiment.

Plastic pots (diameter 10 cm, depth 18 cm) with a perforated base were filled to within 1 cm form the top with one of the three sediment types used (t = 0). In each pot, two soil moisture samplers (Rhizon Flex-5cm; Rhizosphere, Wageningen, the Netherlands) were installed horizontally at depths of 1 cm and 11 cm below the sediment surface (these depths are hereafter referred to as D<sub>1</sub> and D<sub>11</sub>). The pots were stood in rows in the basin. The water level was





maintained at 9 cm so that the sediment at D<sub>11</sub> remained saturated while the sediment at D<sub>1</sub> could oxidize and dry. Each sediment type had 13 replicates.

Reed seedlings (*Phragmites australis*) had been grown in nutrient-poor peat and when 35– 40 days old (experimental time t = 22 days), a single reed seedling was planted per pot in eight of the replicates, leaving five replicates unplanted. Any other seedlings that germinated spontaneously in the pots were removed immediately.

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#### 2.2 Chemical analysis

Soil moisture at D<sub>1</sub> and D<sub>11</sub> was collected from the moisture samplers on days 0, 3, 10, 22, 36, 64, 92, 134 and 174 from five of the pots per condition. The samples from the five replicates were pooled and chemically analyzed. Chloride, NH<sub>4</sub>, NO<sub>2</sub>, NO<sub>3</sub> and SO<sub>4</sub> were determined using ion chromatography (IC); Ca, Fe, K, Mn, Na, P, Si and Sr were determined with Inductively Coupled Plasma Optical Emission Spectrometry (ICP-OES), pH by an ion-specific electrode, and alkalinity was measured by a classic titration method.

Sediment samples were collected for each sediment type at t = 0 and were freeze-dried and stored anoxically prior to geochemical analysis. The major elements were determined using
ICP-OES following an aqua regia destruction. Total S content was measured on an elemental CS analyzer and the mineralogical composition was determined with X-ray diffraction (XRD). A sequential extraction method based on Ruttenberg (1992) was applied to characterize solid P speciation. The method involves five steps (Table 1), the first four of which were carried out anoxically. Loss on ignition (LOI) was determined by slowly heating to 1000 °C. LOI was also





- used as a proxy for organic matter content and total carbonates by calculating the weight loss between 105–550 °C for organic matter and the weight loss between 550–1000 °C for total carbonates (Howard, 1965). Cation exchange capacity (CEC) of the sediments was calculated from the organic matter content and the amounts and types of clay minerals present (Bauer and Velde, 2014).
- Fifty seedlings of *P. australis* randomly chosen from the seedlings grown for the experiment were used to determine the initial tissue contents of Fe, K, P, and N. Their roots, shoots, and leaves were separated and air dried. The air–dried material was then ground and analyzed with total reflection X-ray fluorescence (TXRF) to determine tissue contents of Fe, K, and P. Nitrogen was determined on an elemental CN analyzer. At the end of the experiment (t = 174), the plants in the pots were harvested and subjected to the same procedure, to determine the tissue contents of Fe, K, P, and N.

#### 2.3 Modeling of biogeochemical processes

To identify important biogeochemical processes during the incubation experiments, we modeled with PHREEQC (Parkhurst and Apello, 2013). PHREEQC modeling is frequently used in geochemical research focusing on issues of water quality: examples include investigating mineral weathering in a mountain river (Lecomte et al., 2005), deducing geochemical processes in groundwater (Belkhiri et al., 2010) and investigating the interaction between two aquifers (Carucci et al., 2012). Here, we applied it to identify biogeochemical plant–soil processes during the oxidation and natural drying out of the soil.





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The model approach is based on mass-balance equations of preselected mineral phases (reactants). The mineral phases can either precipitate (leave the solution) or dissolve (enter the solution) and these are expressed in mole transfers. As we only know the dynamics in concentrations of the pore water, we applied inverse modeling in which all possible combinations of the mass-balance equations are accepted within a range of measured pore water concentrations  $\pm 4\%$ . We can simulate infiltration or evaporation rates from the pore water. Since in freshwater mud deposits, the dissolution or precipitation of salts (e.g. NaCl) is negligible and can be ignored, the change in pore water CI concentration was used to calculate the amount of water evaporated or infiltrated.

- To enable the model to attribute some of the chemical changes to cation-exchange processes we included an assemblage of exchangers (X):  $CaX_2$ ,  $FeX_2$ , KX,  $MgX_2$ , NaX and  $NH_4X$ . The sum of this assemblage was defined as CEC calculated from the sediment composition. CEC is important, since it can buffer some of the biogeochemical processes in sediments by adsorption or desorption of cations.
- We used three time frames in our models: 1) oxidation and natural drying out of the soil before the seedlings were transplanted into the pots (t = 0-22 days); 2) initial stage of plant growth (t = 22-64 days); and 3) the stage in which roots started to influence pore water chemistry (t = 64-176 days).

Inverse modeling was applied for all combinations (sediment type, plant/no plant, and depth) for each time frame. For every combination, several valid simulations were found, due to small

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differences in the amount of mole transfers attributed to the mineral phases. Here we present the plausible simulation with the least amount of mole transfers for each combination.

### 190 **3. Results and Discussion**

Below, first the three sediment types will be compared in terms of certain geochemical and mineralogical elements. Next, the composition of the pore water will be introduced and will be linked to biogeochemical processes by presenting and discussing the PHREEQC model simulations. Then, the plant response is presented and discussed in terms of biomass and plant

tissue chemistry. Lastly, the implications for eco-engineering will be discussed.

# 3.1 A brief comparison between sediment types

Table 2 shows the geochemical composition of the disturbed Mud<sub>soft</sub> and Mud<sub>sand</sub> and undisturbed Clay sediments used in this study. The differences between Mud<sub>soft</sub> and Mud<sub>sand</sub> are solely attributable to the presence of inert Dorsilit<sup>®</sup>.

The total sediment concentrations of Al, Fe, Mg, Mn, Na, P, and Zn were significantly higher in Clay then in Mud<sub>soft</sub>. The quartz content was also higher in Clay, which suggests that there were more reactive minerals in this type of sediment.

Sequential P extraction revealed that the significant difference in total P consists of a significantly lower content of Fe-P in Mud<sub>soft</sub> than in Clay (279 mg/kg versus 772 mg/kg; p < 0.01); the other P pools did not differ significantly. The presence of Fe-P in the anoxic Clay





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sediment was unexpected, as in anoxic conditions Fe prefers to bind with S to form FeS<sub>2</sub>. However, after exhaustion of S, precipitation of Fe(II) phosphates may occur (Jilbert and Slomp, 2013). Another possibility is that the reduction of crystalline Fe(III) is not complete in the anoxic sediment because kinetic processes are slow (Canavan et al., 2007). This is likely the case in Markermeer, given our strict anoxic procedures for storage and analysis of the samples. The exchangeable (or loosely sorbed) P was low in Mud<sub>soft</sub> and Clay, indicating that only a small part of the total P found in the sediments was readily available for uptake. The other three P-pools were fairly similar and did not differ significantly between the two types of sediment.

The mineralogical analysis (XRD) showed not only that the quartz content was lower in Mud<sub>soft</sub> than in Clay (37% versus 48%) but that the amounts of calcite and pyrite did not differ between the two types of sediment (9% calcite and 0.6% pyrite). The amount of phyllosilicates (sum of illite, smectite, kaolinite, and chlorite) was higher in Mud<sub>soft</sub> than in Clay: 43% versus 30%. This must also have caused the CEC to be higher in Mud<sub>soft</sub>, as the organic matter content did not differ much between the two (7.2% in Mud<sub>soft</sub> and 6.8% in Clay).

#### 3.2 Pore water composition

Figure 1 presents time series for the pore water concentrations of the three macronutrients N, P, and K. The initial decrease in NH<sub>4</sub> and increase in NO<sub>x</sub> at a depth D<sub>1</sub> for the planted conditions was most likely caused by nitrification as a result of oxidation (Figure 1a–f). At the end of the experiment, almost all dissolved inorganic nitrogen had been removed from the pore water in the pots with plants, whereas in the pots without plants the NH<sub>4</sub> concentrations remained





substantial. Furthermore, a high peak of NO<sub>x</sub> was observed in Clay sediments at day 10 of the experiment. At a depth  $D_{11}$ , no large changes were found in general for NH<sub>4</sub> and NO<sub>x</sub>.

A sharp decline in soluble P was visible at D<sub>1</sub> for all three sediments, probably because P precipitated with Fe(III) when oxygen penetrated the top layer (Figure 1g–i). However, in Clay this decline was preceded by an increase in P. After several weeks, a thin moss layer started to develop on top of the Mud<sub>soft</sub> sediment, which probably prevented oxygen from penetrating and thereby increased the P concentrations (Figure 2g). Similar developments were observed for Mud<sub>sand</sub> although here the moss layer developed much later. In Clay, no moss grew throughout the experiment.

Concentrations of K were higher than concentrations of N and P and increased in the first few weeks (Figure1j–I). No difference was found between pots at  $D_{11}$  with or without plants. However, K was significantly higher at  $D_1$  in the planted pots with Mud<sub>sand</sub> (p < 0.05).

Although it may be important to study measured concentrations of nutrients in pore water in order to understand plant functioning, deriving biogeochemical processes from measured data is problematic because it must be taken into account that changes in pore water can be caused by multiple processes such as drying, dilution, dissolution, and precipitation. Figure 2 reveals that the drying of soils at D<sub>1</sub> was probably an important factor, because we observed an initial increase in Cl that indicated that Cl could not dissolve in the three sediments used (e.g. halite dissolution). Drying will have influenced other variables as well, such as sulfate (Figure 2d–f). Comparing the patterns of Cl and SO<sub>4</sub> suggests that the change in SO<sub>4</sub> concentrations at D<sub>1</sub> should be partly attributed to drying out of soils and partly either to dissolution (e.g. pyrite





oxidation) or to precipitation (e.g. gypsum formation). This highlights the need to use geochemical reaction models like PHREEQC to inversely derive biogeochemical processes from measured data.

### 3.3 Pore water processes (PHREEQC model simulations)

The main pore water processes modeled by PHREEQC are presented in Table 3. For clarity,

255 only major reactants are included in this Table. Supplementary Tables A1 and A2 present mole transfers for all reactants used, as well as the number of valid simulations per combination found.

# 3.3.1 Phase 1: Oxidation and drying (t = 0-22 days)

As discussed in section 3.2, initial drying of soils occurred at D<sub>1</sub> immediately after exposure to air. In the model, this is illustrated by high evaporation rates expressed as H<sub>2</sub>O loss (2300–3400 mmol l<sup>-1</sup> day<sup>-1</sup>; Table 3). The model accounts for this loss by adjusting the solution fractions before calculating other mole transfers.

Exposure to air also leads to oxidation, more so at D<sub>1</sub> than at D<sub>11</sub> (Table 3). The increase in measured sulfate is partly explained as pyrite oxidation (109–270 µmol l<sup>-1</sup> day<sup>-1</sup> for D<sub>1</sub> and 20.1– 36.2 µmol l<sup>-1</sup> day<sup>-1</sup> for D<sub>11</sub>, respectively). Oxidation of pyrite also produces iron oxyhydroxides and protons which in turn promotes dissolution of calcite. The overall reactions are  $FeS_2 + 3.75O_2 + 3.5H_2O \rightarrow Fe(OH)_3 + 2SO_4^{2-} + 4H^+$  (1)



(2)



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followed by calcite dissolution

 $CaCO_3 + H^+ \rightarrow Ca^{2+} + HCO_3^-$ 

The mole transfers for pyrite and calcite presented in Table 3 indicate that not enough calcite is dissolved to buffer all H<sup>+</sup> produced by dissolution of pyrite. Indeed, a drop in pH was observed at the beginning of the experiment (not shown). However, the mineralogical composition presented in Table 2 shows that the amount of calcite (9%; 900 mmol) far exceeds that of pyrite (0.6%; 50 mmol). These numbers suggest that even if all pyrite were to be oxidized, enough calcite is present to buffer all H<sup>+</sup> produced (200 mmol). Note that for Mud<sub>sand</sub> these values are lower due to mixing with Dorsilit<sup>®</sup>.

Some aeration occurred at D<sub>11</sub>. The O<sub>2</sub> fluxes ranged between 61 and 119 µmol l<sup>-1</sup> day<sup>-1</sup>, which resulted in small amounts of pyrite being oxidized (20–36 µmol l<sup>-1</sup> day<sup>-1</sup>). However, sulfate concentrations did not rise, as a result of subsequent precipitation with Ca to form gypsum (53– 73 µmol l<sup>-1</sup> day<sup>-1</sup>).

Furthermore, the cation-exchange-capacity (CEC) of the sediments buffered some processes in pore water chemistry by net adsorption of cations at  $D_1$  and net desorption at  $D_{11}$ .

The processes described above occurred in all three sediments, although some differences were noted. Oxidation was higher in Mud<sub>soft</sub> than in Mud<sub>sand</sub> and Clay, probably because higher evaporation rates in Mud<sub>soft</sub> enhanced oxidation and affected other reactants related to oxidation.

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### 3.3.2 Phase 2: Initial stage of plant growth (t = 22-64 days)

While the pore water compositions did not show clear differences between unplanted and planted conditions, the inverse modeling provided clear evidence for differences at D<sub>1</sub>. However, chemical differences between unplanted and planted conditions for Mud<sub>sand</sub> might simply be attributed to concentration/dilution due to H<sub>2</sub>O loss/gain (-996 to 380 mmol l<sup>-1</sup> day<sup>-1</sup>).

Overall, more pyrite was oxidized in the planted conditions, though the rates are much lower than in the first phase (0–64.3 µmol l<sup>-1</sup> day<sup>-1</sup>). This observation provides evidence that plants may enhance pyrite oxidation by radial oxygen loss (i.e. root aeration). Ferric oxide production on pyrite surfaces probably impeded further oxidation of pyrite, which is a common phenomenon in carbonate-buffered conditions (Nicholson et al., 1990). Indeed, the total pyrite that had

in carbonate-buffered conditions (Nicholson et al., 1990). Indeed, the total pyrite that had oxidized after 64 days (6.3 mmol for Mud<sub>soft</sub>, 2.5 mmol for Mud<sub>sand</sub> and 6.2 mmol for Clay, calculated from the rates presented in Table 3) corresponds to a small fraction of total pyrite present (50 mmol).

Saturation with gypsum led to precipitation of SO<sub>4</sub> and Ca at D<sub>1</sub>. Table 3 shows that with the exception of Mud<sub>sand</sub>, mole transfers were lower for planted conditions; the probable reason is that citric acid production by root tips retarded gypsum precipitation (Prisciandaro et al., 2005). This process was not relevant at D<sub>11</sub>, as here aeration (and subsequent sulfate production) by plant roots was minor (in the case of Clay) or absent (in the case of Mud<sub>soft</sub> and Mud<sub>sand</sub>).

The thin moss layer that started to develop after several weeks in the unplanted condition on top of the Mud<sub>soft</sub> sediment slowed down the aeration rate to 2.62 µmol l<sup>-1</sup> day<sup>-1</sup> and might be





the reason for the moderate increase in P, which probably resulted from  $Fe(OH)_3$  dissolution (0.95 µmol l<sup>-1</sup> day<sup>-1</sup>) (Figure 1g, Table 3).

3.3.3 Phase 3: Root influence (t = 64-176 days)

Phase 3 took place in the autumn, when temperatures were lower and therefore the soils did not dry out; hence there was a net gain in H<sub>2</sub>O. The gain was less in planted conditions, due to uptake of water by roots.

The fully grown plants continued to influence pore water chemistry at D<sub>1</sub>, but in the unplanted conditions the chemical changes were minor (Table 3). Radial oxygen loss continued the oxidation processes described in the previous sections. It should be noted that *P. australis* is known to have higher radial oxygen loss than other wetland species (Brix et al., 1996; Dickopp et al., 2011; Smith and Luna, 2013), so the aeration effect found in this study cannot be assumed to hold for other species.

In contrast to the previous phase, in phase 3 the influence of roots was clearly visible at D<sub>11</sub> for all three sediments. All planted sediments showed increased aeration and subsequent oxidation of pyrite due to radial oxygen loss, with a notable difference between Mud<sub>soft</sub> (lower) and Mud<sub>sand</sub> (higher). This is somewhat surprising, as the belowground biomass was significantly higher in Mud<sub>soft</sub> (section 3.4). It indicates that increasing the average grain size by adding sand enhanced aeration, even when root biomass production was low.

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#### 3.4 Plant response

Above- and belowground biomass were significantly higher in Mud<sub>soft</sub> and Clay than in Mud<sub>sand</sub> (Figure 3; p < 0.02). The difference between the two Mud sediments cannot be explained by nutrient concentrations in pore water or light conditions in the greenhouse, as these were the same for the two sediments. As biomass production in Mud<sub>sand</sub> was not limited by chemical or biological properties relative to Mud<sub>soft</sub>, it seems likely that the reason for the lower biomass production in Mud<sub>sand</sub> is a difference in physical properties. Voorhees et al. (1975) and Bengough and Mullins (1990) showed that so-called mechanical impedance (i.e. the resistance to penetration by the root tip) was higher in loamy sand than in clay, which was attributed to the higher bulk density of the loamy sand. Therefore, increasing the bulk density of Mud<sub>soft</sub> by mixing with sand increased the mechanical impedance and this might explain the lower biomass production we observed in Mud<sub>sand</sub>.

*P. australis* invested more in its root system than in its shoots and leaves for all sediments (Figure 3). More investment in roots implies a limitation of N, P, and/or S (Ericsson, 1995; Shipley and Meziane, 2002). Figures 1a–i and 2d–f show that the N and P concentrations were indeed low in the planted conditions but that SO<sub>4</sub> was high, which rules out S limitation. During the experiment we had observed reduced plant growth and shriveling and yellowing of foliage 2 months after transplantation, which might have been caused by nutrient limitation.

Figure 4 shows the N, P, and K contents as well as the N:P ratio for the roots of *P. australis* at the beginning and end of the experiment for the three sediment types. The N, P, and K contents in the roots increased in time, while the N:P ratio clearly decreased. The reduction in





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N:P ratio from 11 to 2–3 suggests N was the limiting nutrient as an N:P ratio of < 14 in plant tissue is indicative of N limitation (Koerselman and Meuleman, 1996). However, root N and P concentrations of *P. australis* should typically range between 0.64–1.04% for N and 0.06–0.13% for P (Wang et al., 2015). Figure 4 shows that the root N and P concentrations were above these values, and that P was particularly high: by a factor of 5 to 10 (N: 1.14–1.63% and P: 0.52–0.62%). Hence the concentrations of these nutrients in the roots do not indicate that nutrient limitation is a likely cause of the reduced plant growth and shriveling and yellowing of foliage.

We hypothesize that co-precipitation of P with Fe on roots enhanced the concentrations of P in the plant roots (Snowden and Wheeler, 1995; Jørgenson et al., 2012). Snowden and Wheeler (1995) showed that this so-called iron plaque formation enhances uptake of Fe and P. This may cause iron toxicity and is probably responsible for the elevated P concentrations in tissue, and for the stunted growth and leaf decay we observed in the experiment. Note that the plant roots of *P. australis* initiate this process by oxidizing their environment and thereby enabling ferrous iron to oxidize into P-bearing ferric iron, which precipitates on roots.

The Fe concentration in the leaves and in the roots supports the "Fe-P co-precipitation hypothesis": we measured an approximately 20-fold increase by comparison with the initial concentration in the seedlings (Figure 5). Furthermore, ferric oxide, a product of pyrite oxidation, precipitates on root surfaces (Jørgenson et al., 2012), and hence pyrite oxidation in sediments is directly linked to iron toxicity in plants.

Further evidence to support our hypothesis is provided by the results of the sequential phosphorus extraction conducted on the sediments: it revealed that the dominant P pool in the



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sediments is the Fe-P fraction (Table 2). P co-precipitates with Fe on roots if it is bound to ferric oxides.

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### 3.5. Implications for eco-engineering

Our results strongly point in the direction of iron toxicity as a major bottleneck prohibiting healthy development of *P. australis*. Since the candidate material for the construction of the Markermeer wetland has high contents of Fe and Fe-P, we recommend using Fe-tolerant plant species as test species in the new wetland, rather than species optimized for growing in N-limited conditions.

Concomitantly with iron toxicity, a high Fe-P content in soil will trigger P mobilization if that soil is rewetted after having dried out and contains high amounts of SO<sub>4</sub> (Smolders and Roelofs, 1993; Lucassen et al., 2005). In some cases, this can result in elevated levels of sulfide, thereby promoting S toxicity in plants (Lamers et al., 1998; Van der Welle et al., 2007).

Figure 6 summarizes the important feedbacks and processes we expect play an important role in the clay-rich sediments. Following the feedback loops between plant and soil, we see a negative feedback loop that arises because plant roots induce aeration, which promotes iron toxicity that decreases plant growth and results in plant death. Also, we see a positive feedback loop, as iron toxicity induces reduction processes as a result of root death, which leads to P mobilization and hence enhances plant growth and regeneration. Negative feedback loops diminish or buffer changes, whereas a positive feedback loop amplifies changes. So, a negative feedback loop normally stabilizes the system, in our case via the toxic effect of iron oxides on





plants, but plant growth may increase due to the positive feedback loop via P mobilization. The relative strengths of these two feedback loops and the sensitivity of species to Fe toxicity determine the ultimate effect on vegetation development in wetlands built from these sediments. As drying–rewetting cycles are likely to occur in these future wetlands and since the Fe-P concentrations in the building material are high, these feedbacks might be an important factor influencing soil formation and ecosystem development. We therefore recommend studying the ultimate effects of the use of this material on ecosystem development by testing with various plant species and drying–rewetting cycles.

#### 4. Conclusions

The results of this study show that plants expedite biogeochemical processes by oxidizing and modifying their environment, which in turn affects the growth conditions of the plants. In the mud deposits from Markermeer, the key processes influencing pore water chemistry are pyrite oxidation and associated calcite dissolution. The former is especially likely to be important as it is linked to iron toxicity and P mobilization and thus has the potential to initiate two feedback mechanisms between plant and soil. We found strong indications for a negative feedback loop, where plant-induced iron toxicity is hampering plant growth and a positive feedback loop, where iron toxicity promotes P mobilization and thereby enhances plant growth. The strength of these feedbacks and the balance between them will play an important role in regulating ecoengineering conditions for plants.





415 We found conclusive evidence that the low N:P ratio found in plant tissue was not caused by N limitation, as the ratio suggests, but probably results from enhanced P uptake, as a result of co-precipitation with Fe on roots.

The magnitude of the feedback mechanisms is expected to differ between the building materials used. The soft clay-rich layer has less Fe-P than the underlying clay layer and therefore P mobilization is expected to be less in mud. However, when the mud is mixed with sand, the enhanced aeration brought about by the change in grain-size composition results in higher oxidation rates, increasing the impact on P mobilization and iron toxicity.

To study the effects of iron toxicity and P mobilization in greater detail we recommend further testing with different plant species and drying-rewetting cycles. This is important because we

425 expect these mechanisms to influence soil formation and ecosystem development in the future wetland.

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**Table 1**. List of steps used in the extraction procedure of phosphorus (based on Ruttenberg, 1992).

Step	)	Extractant	Separated P fraction
Ι		1M MgCl <sub>2</sub> , 30 min	Exchangeable or loosely sorbed P
Ш	А	Citrate-dithionite-bicarbonate (CDB), 8 h	Easily reducible or reactive ferric Fe- P
	В	1M MgCl <sub>2</sub> , 30 min	
Ш	А	Na acetate buffer (pH 4), 6 h	Amorphous apatite and carbonate P
	В	1M MgCl <sub>2</sub> , 30 min	
IV		1M HCl, 24 h	Crystalline apatite and other inorganic P
V		Ash at 550 °C, 2h; 1M HCl, 24 h	Organic P





**Table 2.** Geochemical and mineralogical composition of the sediment types used in this study. Significant differences between Mud<sub>soft</sub> and Clay are indicated by \* (p < 0.05).

		n	Mud <sub>soft</sub>		Mudsand		Clay			
	Unit per		Mean SD		Mean	SD	Mean	SD		
Aqua regia / CS		type								
AI*	mg/kg	15	16593	3130	6394	2439	21989	4512		
Са	mg/kg	15	45635	6020	18877	3572	48031	3032		
Fe*	mg/kg	15	20745	2987	7804	2281	27766	3764		
К	mg/kg	15	4102	641	1723	742	5371	1262		
Mg*	mg/kg	15	6636	906	2531	558	8041	1017		
Mn*	mg/kg	15	577	160	238	62	710	166		
Na*	mg/kg	15	526	158	219	64	992	379		
P*	mg/kg	15	649	169	259	56	1186	217		
S	mg/kg	15	5586	698	3001	846	5727	710		
Sr	mg/kg	15	135	26	62	14	148	21		
Ті	mg/kg	15	312	77	125	44	312	74		
Zn*	mg/kg	15	110	29	43	18	159	58		
Seq. P extraction										
Exchangeable P	mg/kg	15	11.9	3.50	5.9	1.79	14.3	6.81		
Fe- bound P*	mg/kg	15	279	61.7	94.5	29.0	772	263		
Ca-bound P	mg/kg	15	121	30.9	36.8	13.1	146	43.3		
Detrital P	mg/kg	15	169	14.1	51.5	10.9	147	16.5		
Organic P	mg/kg	15	117	25.1	47.7	8.38	99.6	20.0		
XRD										
Quartz	%	1	37		n.a.		48			
Calcite	%	1	9		n.a.		9			
Pyrite	%	1	0.6		n.a.		0.6			
Illite	%	1	21		n.a.		15			
Smectite	%	1	14		n.a.		11			
Kaolinite	%	1	5		n.a.		3			
Chlorite	%	1	3		n.a.		2			
Other										
Organic matter	%	5	7.2	0.6	2.8	0.4	6.7	0.6		
CEC (calculated)	meq/100g	meq/100g 37.2			12.4		30.0			





Table 3. Main pore water processes expressed in mole transfers (μmol l<sup>-1</sup> day<sup>-1</sup>) as modeled by PHREEQC with pore water data

retrieved at 1 cm and 11 cm below sediment surface (D1 and D11 respectively). Positive values indicate dissolution, negative values

indicate precipitation. Cation exchange capacity (CEC) is the sum of Ca, Fe, K, Mg, Na, and NH<sub>4</sub>.

			Ca	cite	Gyp	sum	Fe(	OH)₃	Py	rite	ΣC	EC	H₂O (	x 10³)	c	<b>)</b> 2
Phase	Condition		D1	<b>D</b> 11	D1	D <sub>1</sub> D <sub>11</sub>	D1	<b>D</b> <sub>11</sub>	D1	<b>D</b> 11	D1	<b>D</b> 11	D <sub>1</sub>	<b>D</b> 11	D <sub>1</sub>	<b>D</b> 11
1. Oxidation	Mud <sub>soft</sub>	No plant	267	111	0.00	-72.5	-277	0.00	270	36.2	-31.3	20.2	-3364	0.00	1009	119
t=0-22 days	Mudsand	No plant	0.00	59.6	0.00	-40.7	-116	0.00	109	21.7	-4.99	7.92	-2591	0.00	432	69.5
	Clay	No plant	120	55.2	0.00	-53.4	-160	0.00	159	20.1	-91.4	14.0	-2364	0.00	659	61.9
2. Initial root	Mud <sub>soft</sub>	No plant	27.1	0.00	-236	0.00	0.95	-0.24	0.00	0.00	-23.1	1.43	0.00	0.00	2.62	0.00
development		Plant	48.8	19.8	-208	-3.81	-10.0	-6.19	9.76	0.00	-7.63	1.43	0.00	0.00	45.5	0.00
t=22-64 days	Mudsand	No plant	39.3	71.7	0.00	0.00	0.00	-41.2	0.21	0.00	1.90	1.46	380	0.00	0.00	0.00
		Plant	7.10	83.8	-83.4	0.00	0.00	-51.2	3.58	0.00	5.40	3.40	-996	0.00	0.00	0.00
	Clay	No plant	0.00	27.1	-32.1	0.00	-21.4	-25.0	21.2	0.00	0.01	-0.23	-286	0.00	41.9	0.00
	-	Plant	36.9	16.2	0.00	0.00	-14.3	0.00	64.3	11.9	28.4	4.53	-6.67	0.00	186	40.5
3. Root influence	Mud <sub>soft</sub>	No plant	0.00	-3.21	-19.2	0.00	-1.34	-0.80	0.00	0.00	-1.07	-1.43	56.3	0.00	0.00	0.00
t=64-176 days		Plant	25.8	0.00	0.00	0.00	-4.20	0.00	23.8	4.11	7.88	-4.65	49.1	0.00	83.6	13.6
	Mudsand	No plant	8.13	0.00	-7.59	0.00	-10.6	-1.34	0.00	0.00	-1.78	1.42	74.1	0.00	0.00	0.00
		Plant	0.00	0.00	-14.8	0.00	-13.3	-23.2	13.8	7.95	0.12	-10.6	-357	-652	44.7	32.6
	Clay	No plant	0.00	11.5	0.00	0.00	0.00	-13.8	33.3	0.00	23.9	0.36	134	0.00	113	0.00
	,	Plant	115	18.7	0.00	0.00	-58.5	-8.48	58.3	8.57	45.4	-5.73	0.00	-98.2	215	28.4

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Figure 1. Time series of NH<sub>4</sub> (a–c), NO<sub>x</sub> (d–f), P (g–i) and K (j–l) concentrations. Each
column represents one sediment type: Mud<sub>soft</sub> (a, d, g, j), Mud<sub>sand</sub> (b, e, h, k), and Clay
(c, f, i, l). The variable and the scale of the x-axis are the same for each row, except
for the scale in f.















- Figure 3. Above- and belowground biomass in grams dry weight, with error bars (n =
- 5). Significant differences between sediment types are indicated by different letters,
- and non-significant differences are indicated by a similar letter.







Figure 4. N, P, and K concentration in root tissue (t = 176) in % of dry weight (a) as
well as the N:P ratio (b) with error bars when n = 5. Significant differences between
sediment types are indicated by different letters, and non-significant differences are
indicated by a similar letter.







- **Figure 5.** Fe concentration (% of dry weight) in leaf and root tissue with error bars
- when n = 5. Significant differences between sediment types are indicated by different
- 620 letters, and non-significant differences are indicated by a similar letter.







Figure 6. Most important biogeochemical processes and feedbacks identified in this
study. + indicates positive feedback, - indicates negative feedback.





#### 623 Appendix

#### 624

Table A1. Pore water processes expressed in mole transfers (μmol l<sup>-1</sup> day<sup>-1</sup>) as modeled by PHREEQC with pore water data

retrieved at 1 cm below sediment surface. Positive values indicate dissolution, negative values indicate precipitation.

		Phase 1.	Oxidation	(t=0-22)	Phase 2. Initial root development (t=22-64)						Phase 3. Root influence (t=64-176)						
		No plant	No plant	No plant	No plant	Plant	No plant	Plant	No plant	Plant	No plant	Plant	No plant	Plant	No plant	Plant	
Reactant	Composition	Mud <sub>soft</sub>	Mudsand	Clay	Mud <sub>soft</sub>		Mudsand		Clay		Mud <sub>soft</sub>		Mudsand		Clay		
Calcite	CaCO₃	267	0.00	120	27.1	48.8	39.3	7.1	0.00	36.9	0.00	25.8	8.13	0.00	0.00	115	
Gypsum	CaSO <sub>4</sub> :2H <sub>2</sub> O	0.00	0.00	0.00	-236	-208	0.00	-83.4	-32.1	0.00	-19.2	0.00	-7.59	-14.8	0.00	0.00	
Hydroxyapatite	e Ca₅(PO₄)₃(OH)	-5.00	-3.64	0.00	0.24	0.00	-0.02	-0.04	0.00	0.00	0.18	0.00	0.09	0.00	0.00	0.00	
Chalcedony	SiO <sub>2</sub>	-19.1	-15.5	-18.2	0.95	0.71	1.91	-3.37	-1.67	-2.14	0.71	0.00	0.54	1.43	0.00	-0.36	
Fe(OH)₃ (a)	Fe(OH)₃	-277	-116	-160	0.95	-10.0	0.00	0.00	-21.4	-14.3	-1.34	-4.20	-10.6	-13.3	0.00	-58.5	
Pyrite	FeS₂	270	109	159	0.00	9.76	0.21	3.58	21.2	64.3	0.00	23.8	0.00	13.8	33.3	58.3	
Rhodochrosite	MnCO <sub>3</sub>	-11.8	-11.4	-2.27	2.86	1.19	1.23	0.34	-0.24	-0.24	-0.63	-0.89	0.09	0.18	0.00	0.00	
CEC	CaX <sub>2</sub>	0.00	20.9	55.5	63.1	41.9	-9.11	0.00	0.00	0.00	2.50	0.00	-9.73	0.00	-9.64	-85.4	
	FeX <sub>2</sub>	0.00	0.00	0.00	0.00	0.00	-0.19	-4.11	0.00	-50.2	1.61	-19.8	11.7	0.00	-33.3	0.00	
	кх	-8.64	-5.00	-17.7	-4.76	0.00	3.78	-8.30	-6.19	0.00	-2.14	-2.14	-2.77	-7.68	0.00	0.00	
	MgX <sub>2</sub>	31.4	-16.8	36.8	-39.8	-30.5	7.42	-1.35	0.00	21.7	-3.04	12.0	0.00	0.00	19.1	39.8	
	NaX	-20.9	0.00	-166	-46.4	-25.7	0.00	25.1	25.2	77.6	0.00	19.7	0.00	12.0	49.4	92.9	
	NH₄X	-33.2	-4.09	0.00	4.76	6.67	0.00	-5.94	-19.0	-20.7	0.00	-1.88	-0.98	-4.20	-1.70	-1.88	
H₂O (g)	H₂O x 10 <sup>3</sup>	-3364	-2591	-2364	0.00	0.00	380	-996	-286	-6.67	56.3	49.1	74.1	-357	134	0.00	
O <sub>2</sub> (g)	<b>O</b> <sub>2</sub>	1009	432	659	2.62	45.5	0.00	0.00	41.9	186	0.00	83.6	0.00	44.7	113	215	
CO <sub>2</sub> (g)	CO2	-827	-532	-650	35.2	0.00	39.7	0.00	-55.5	-84.8	0.00	-33.1	0.00	44.6	-31.7	-115	
1	lo. models found	2	2	2	3	4	2	2	5	2	6	2	1	2	2	1	





**Table A2.** Pore water processes expressed in mole transfers (μmol l<sup>-1</sup> day<sup>-1</sup>) as modeled by PHREEQC with pore water data

628	retrieved at 11	cm below sediment s	surface. Positive	values indicate	dissolution,	negative va	alues indicate	precipitation.
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		Phase 1. Oxidation (t=0-22)			Phase 2	root deve	Phase 3. Root influence (t=64-176)									
		No plant	No plant	No plant	No plant	Plant	No plant	Plant	No plant	Plant	No plant	Plant	No plant	Plant	No plant	Plant
Reactant	Composition	Mud <sub>soft</sub>	Mudsand	Clay	Mud <sub>soft</sub>		Mudsand		Clay		Mud <sub>soft</sub>		Mudsand		Clay	
Calcite	CaCO <sub>3</sub>	111	59.6	55.2	0.00	19.8	71.7	83.8	27.1	16.2	-3.21	0.00	0.00	0.00	11.5	18.7
Gypsum	CaSO <sub>4</sub> :2H <sub>2</sub> O	-72.5	-40.7	-53.4	0.00	-3.81	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Hydroxyapatite	Ca₅(PO₄)₃(OH)	0.00	0.51	1.45	-0.24	0.00	0.00	0.00	0.00	0.00	0.00	-0.09	0.00	-0.45	0.00	-0.18
Chalcedony	SiO <sub>2</sub>	4.44	5.32	6.74	1.90	3.33	3.10	3.81	1.67	0.95	-0.18	-1.07	-0.27	-3.48	0.00	-1.07
Fe(OH)₃ (a)	Fe(OH)₃	0.00	0.00	0.00	-0.24	-6.19	-41.2	-51.2	-25.0	0.00	-0.80	0.00	-1.34	-23.2	-13.8	-8.48
Pyrite	FeS₂	36.2	21.7	20.1	0.00	0.00	0.00	0.00	0.00	11.9	0.00	4.11	0.00	7.95	0.00	8.57
Rhodochrosite	MnCO₃	0.00	1.18	0.31	0.00	0.48	1.19	0.95	0.00	0.24	0.00	0.00	0.00	-0.71	0.18	0.09
CEC	CaX <sub>2</sub>	0.00	0.00	0.00	-1.43	-5.95	-50.7	-63.3	-7.86	0.00	1.70	8.39	0.00	0.00	-3.75	0.00
	FeX₂	-35.5	-20.9	-19.0	0.00	0.00	42.4	51.7	0.00	-11.9	1.07	-3.66	-0.54	15.2	4.29	0.00
	кх	7.00	5.87	3.76	0.00	0.00	2.62	2.86	-5.95	1.67	-0.89	-1.79	0.00	-3.84	0.00	-1.70
	MgX <sub>2</sub>	15.4	13.0	4.87	0.00	4.76	7.14	8.57	8.10	7.38	-1.25	0.00	0.00	-4.11	2.59	5.71
	NaX	25.2	9.95	24.4	0.00	0.00	0.00	5.24	0.00	6.43	-4.29	-4.20	1.96	-12.4	0.00	-5.54
	NH₄X	8.12	0.00	0.00	2.86	2.62	0.00	-1.67	5.48	0.95	2.23	-3.39	0.00	-5.80	-2.77	-4.20
H₂O (g)	H₂O x 10 <sup>3</sup>	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	-652	0.00	-98.2
O <sub>2</sub> (g)	<b>O</b> <sub>2</sub>	119	69.5	61.9	0.00	0.00	0.00	0.00	0.00	40.5	0.00	13.6	0.00	32.6	0.00	28.4
CO <sub>2</sub> (g)	CO <sub>2</sub>	156	0.00	43.0	0.00	0.00	0.00	0.00	0.00	14.5	0.00	0.00	0.00	-67.3	0.00	-13.7
No	o. models found	2	2	1	4	4	2	2	3	2	1	4	2	4	2	1