

# Sediment phosphorus speciation and mobility under dynamic redox conditions

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Abstract. Anthropogenic nutrient enrichment has caused phosphorus (P) accumulation in many freshwater sediments, raising concerns that internal loading from legacy P may delay the recovery of aquatic ecosystems suffering from eutrophication. Benthic recycling of P strongly depends on the redox regime within surficial sediment. In many shallow environments, redox conditions tend to be highly dynamic as a result of, among others, bioturbation by macrofauna, root activity, sediment resuspension and seasonal variations in bottom-water oxygen (O2) concentrations. To gain insight into the mobility and biogeochemistry of P under fluctuating redox conditions, a suspension of sediment from a hypereutrophic freshwater marsh was exposed to alternating 7-day periods of purging with air and nitrogen gas  $(N_2)$ , for a total duration of 74 days, in a bioreactor system. We present comprehensive data time series of bulk aqueous- and solidphase chemistry, solid-phase phosphorus speciation and hydrolytic enzyme activities demonstrating the mass balanced redistribution of P in sediment during redox cycling. Aqueous phosphate concentrations remained low ( $\sim 2.5 \,\mu\text{M}$ ) under oxic conditions due to sorption to iron(III) oxyhydroxides. During anoxic periods, once nitrate was depleted, the reductive dissolution of iron(III) oxyhydroxides released P. However, only 4.5 % of the released P accumulated in solution while the rest was redistributed between the MgCl<sub>2</sub> and NaHCO3 extractable fractions of the solid phase. Thus, under the short redox fluctuations imposed in the experiments, P remobilization to the aqueous phase remained relatively limited. Orthophosphate predominated at all times during the experiment in both the solid and aqueous phase. Combined P monoesters and diesters accounted for between 9 and 16% of sediment particulate P. Phosphatase activities up to 2.4 mmol  $h^{-1}$  kg<sup>-1</sup> indicated the potential for rapid mineralization of organic P (P<sub>o</sub>), in particular during periods of aeration when the activity of phosphomonoesterases was 37% higher than under N<sub>2</sub> sparging. The results emphasize that the magnitude and timing of internal P loading during periods of anoxia are dependent on both P redistribution within sediments and bottom-water nitrate concentrations.

#### 1 Introduction

It is widely recognized that accelerated eutrophication of freshwater aquatic environments is caused primarily by anthropogenic increases to dissolved phosphorus (P) concentrations in surface water (Smith and Schindler, 2009). Rapid cultural eutrophication of oligo or mesotrophic lacustrine and palustrine systems is often attributed to increased external P loadings originating in agricultural run-off or waste water treatment plant (WWTP) effluent. The resultant excessive algal growth negatively impacts aquatic ecosystems and economic activity (Pretty et al., 2003) and increases the risk of infectious diseases (Chun et al., 2013). Strategies to mitigate eutrophication have aimed to reduce point source and diffuse external phosphorus loadings by instituting agricultural best-management practices in the surrounding watershed (McLaughlin and Pike, 2014; Sharpley et al., 1994), limiting P inputs to domestic waste water (Corazza and Tironi, 2011) and upgrading WWTPs (Mallin et al., 2005). However, internal loading of P, from sediments to surface water, remains poorly quantified in many systems and is often the largest source of error in hydrodynamic and ecological phosphorus models (Kim et al., 2013). Early diagenesis and mineralogical removal of labile autochthonous organic phosphorus ( $P_o$ ) from solution is a complex process and is poorly understood in highly dynamic systems despite exerting a strong influence on the magnitude and timing of internal P loading. This is particularly true in shallow lakes and wetlands due to the high ratio of sediment surface area to water column depth (Søndergaard et al., 2003).

As policy and infrastructure improvements continue in order to mitigate external P inputs to aquatic systems, the relative importance of internal P loads from legacy P in sediments to overall P budgets in eutrophic systems is likely to increase (Reddy et al., 2011).

It has been widely demonstrated through laboratory and field investigations, particularly in seasonally anoxic lakes, that sustained anoxic conditions induced by water column stratification typically result in greater P mobility and correspondingly higher water column P concentrations (Krom and Berner, 1981; Einsele, 1936; Hongve, 1997; Katsev et al., 2006; Mortimer, 1941, 1971; Penn et al., 2000). The microbially mediated reductive dissolution of iron(III) oxyhydroxides or iron(III) phosphate during sustained periods of anoxia at the sediment–water interface (SWI) has long been considered the main mechanism responsible for P release under anoxic conditions (Bonneville et al., 2004; Hyacinthe and Van Cappellen, 2004).

More recently, considerable microbial polyphosphate accumulation and release in response to alternating oxic– anoxic conditions at the SWI in lacustrine environments has also been shown to occur (Gächter et al., 1988; Gächter and Meyer, 1993; Hupfer et al., 2007; Sannigrahi and Ingall, 2005). In some cases this accumulation of polyphosphate by the microbial community may account for 10 % of total phosphorus (Hupfer et al., 2007).

However, redox conditions in shallow, heavily bioturbated sediments are more spatially and temporally variable than in seasonally anoxic lakes (Aller, 1994; Gorham and Boyce, 1989), resulting in short-term redox oscillations even with continuous oxia at the SWI.

Additionally, the coupled biogeochemical cycles of other redox sensitive elements such as sulfur and carbon have been shown to play important and complex roles in phosphorus mobility (Gächter and Müller, 2003; Joshi et al., 2015; O'Connell et al., 2015). For example, high bottom-water sulfate concentrations have been shown to increase aqueous P in sediments by decreasing the permanent mineralogical removal of P within vivianite  $[Fe_3(PO_4)_{2(s)}]$  and by decreasing the abundance of iron(III) oxyhydroxides near to the SWI (Caraco et al., 1989). This is due to the scavenging of iron during the formation of iron sulfide minerals within sediment during diagenesis (Gächter and Müller, 2003). Carbon cycling also exerts considerable control over phosphorus mobility within sediment. The stoichiometry of freshly deposited organic matter (OM) in eutrophic water bodies approaches that of primary production, i.e.  $\sim C_{106} : N_{16} : P$ (Berner, 1977). Appreciable P may therefore be released to the aqueous phase when organic carbon is mineralized during microbial respiration of oxygen, nitrate, ferric iron or sulfate. In addition to driving N, Fe and S cycling, mineralization of organic carbon and concomitant P release has, in some places, been shown to be the primary mechanism controlling phosphorus mobility at the SWI (Joshi et al., 2015).

Core incubations and in situ flux chambers frequently examine the effects of anoxia on P mobility from sediments but the effects of repetitive redox oscillations are rarely investigated in a controlled setting (Frohne et al., 2011; Matisoff et al., 2016; Nürnberg, 1988; Thompson et al., 2006). Consequently, the cumulative and reversible effects of oxic–anoxic cycling on P distribution, speciation and mobility within sediments are poorly understood.

The aim of this study is to elucidate the microbial and geochemical mechanisms of in-sediment phosphorus cycling and release associated with commonly occurring short redox fluctuations (days) in surficial sediments in shallow eutrophic environments. Particularly, we aimed to (1) quantify the redistribution of P between aqueous and mineral sediment pools during fluctuating redox conditions; (2) determine whether the activity of hydrolytic phosphatase enzymes acting on  $P_0$  were influenced by redox conditions; (3) assess whether the proportions of orthophosphate,  $P_0$ , and polyphosphate varied systematically with redox conditions; and (4) ascertain whether P mobilization/immobilization mechanisms were reversible or cumulative.

We conducted controlled bioreactor experiments using sediment suspensions, designed to reproduce cyclic redox conditions analogous to those occurring in nature (Aller, 1994, 2004). A combination of aqueous chemistry, sediment sequential extractions (Ruttenberg, 1992), <sup>31</sup>P NMR spectroscopy (Cade-Menun, 2005) and extra-cellular enzyme assays (Deng et al., 2013) were used to produce a comprehensive dataset describing bulk chemistry, microbial and mineralogical controls on P mobility and speciation during redox oscillations.

#### 2 Methods

#### 2.1 Field site and sampling

Surface sediment (0–12 cm), sediment cores (34 cm long, 10 cm diameter), overlying water and green filamentous algae (GFA) were collected on 5 September 2013 from West Pond in Cootes Paradise Marsh (43.26979° N, 79.92899° W) following established guidelines (US EPA, September 2009). Cootes Paradise is a hypereutrophic, coastal freshwater marsh, which drains into Lake Ontario via Hamilton Harbour (see Fig. 1a–c). The marsh system suffered severe degrada-



**Figure 1. (a)** Photograph of the sampling location taken on the day of sampling illustrating the abundance of green filamentous algae. **(b)** Map of Cootes Paradise and West Pond showing the sampling location, local hydrological network and the King Street Waste Water Treatment plant in Dundas, Hamilton. Colour represents area covered by surface water at different water levels (m.s.l. is metres above sea level). **(c)** Overview map showing the hydrological connection between Cootes Paradise, Hamilton Harbour and Lake Ontario.

tion due to rapid urbanization, population growth and nutrient loadings in the 20th century (Chow-Fraser et al., 1998). West Pond in particular received extremely high external P loads from the King Street (Dundas) WWTP for several decades prior to the installation of sand filters in 1987 (Painter et al., 1991). The addition of sand filters, and other improvements, decreased P loadings from the WWTP from  $45 \text{ kg P day}^{-1}$  in the early 1970s (Semkin et al., 1976) to  $4.5 \text{ kg P day}^{-1}$  in the 1980s (Chow-Fraser et al., 1998) and  $2.59 \text{ kg P day}^{-1}$  in 2011 (Routledge, 2012). However, high external P loads resulted in accumulation of legacy P in West Pond sediments with total phosphorus concentrations reaching 200  $\mu$ mol g<sup>-1</sup> by the 1980s (Theysmeyer et al., 1999). Consequently, dredging was conducted in 1999 in an attempt to remediate the areas most affected by growth of green filamentous algae (Bowman and Theysmeyer, 2014).

Despite these restoration efforts and decreases to the external P load, pervasive growth of GFA during the summer persists in parts of Cootes Paradise (Fig. 1a). Cyanobacteria are not commonly observed at this location, potentially due to the high N : P ratios often associated with WWTP which utilize tertiary P removal treatment (Conley et al., 2009; Stumm and Morgan, 1996).

#### 2.2 Sediment characterization

Sediment cores were sliced every 3 cm, homogenized and characterized with bulk sediment samples prior to bioreactor experiments. Organic carbon and carbonate depth profiles were determined by thermogravimetric analysis (TGA-Q500, TA Instruments Q500) (Pallasser et al., 2013). Water

content and bulk density ( $\rho_b$ ) of the sliced sediment core were determined gravimetrically after oven drying (Gardner, 1986). Identification and quantification of crystalline mineralogy was determined by powder X-ray diffraction (XRD) (Empyrean Diffractometer and Highscore Plus software Ver. 3.0e PANalytical). The density of benthic macroinvertebrates was also quantified after sieving two additional cores (7.5 cm diameter, 18 cm deep) through 500 µm mesh.

# 2.3 Bioreactor experiment and redox oscillation procedure

An initial concentrated sediment suspension of approximately 500 g  $L^{-1}$  (dry weight equivalent) was prepared from freshly sampled sediment (0-12 cm) and filtered overlying water ( $< 0.45 \,\mu\text{m}$ ). Surface water was used, rather than distilled water, to provide background ionic strength and avoid osmotic shock to the microbial community. The concentrated suspension was stirred vigorously for 5 min then passed through a  $< 500 \,\mu\text{m}$  stainless steel sieve to remove larger solid organic material and macro-invertebrates. This procedure was repeated until a homogeneous suspension was achieved. The dry weight was then re-determined, and the sieved solution was diluted with filtered surface water to a final concentration of  $247 \pm 2 \text{ g L}^{-1}$ . The resulting suspension was transferred to a bioreactor system (Applikon Biotechnology) after Thompson et al. (2006) and Parsons et al. (2013). In addition to affording precise temperature control and continuous logging of temperature, redox potential  $(E_h)$  and pH, the system offers significant advancements over previous designs (Thompson 2006; Guo, 2007; Parsons, 2013).

The  $E_h$ , pH and dissolved oxygen (DO) were measured using a combined autoclavable Mettler Toledo InPro 3253i/SG open-junction electrode and an AppliSens low drift polarographic sensor. The InPro electrode system, using a common reference electrode, was chosen to help avoid potential interference between two electrodes in close proximity. A multiparameter transmitter was used to display current pH,  $E_h$  and temperature to automatically temperature-correct pH values and to adjust measured  $E_h$  to the standard hydrogen electrode (SHE). DO was calibrated using 100 % saturation in air (approximately 0.2905 atm) and 0 % saturation in N<sub>2</sub> at constant sparging of 30 mL min<sup>-1</sup>

The suspension was stirred continuously and sparged with  $30 \text{ mL min}^{-1}$  air for 11 days to equilibrate prior to the redox oscillation procedure. During the 11-day oxic equilibration period, CO<sub>2</sub> emissions were monitored in the reactor exhaust gas using an IR sensor (Applikon Biotechnology).

Subsequently, redox potential  $(E_h)$  variation was induced by the modulation of sparging gases  $(30 \text{ mL min}^{-1})$  between  $N_2: CO_2$  and  $O_2: N_2: CO_2$ . The suspension was subjected to five cycles of anoxia (7 days) and oxia (7 days) at constant temperature (25 °C) in the dark, while recording  $E_h$ , pH, DO and temperature data. The suspension was sampled on days 1, 3, 5 and 7 of each half-cycle. To separate solid and aqueous components from the sediment suspension, syringe-extracted samples (15 mL) were centrifuged at 5000 rpm for 20 min and the supernatant filtered through 0.45 µm polypropylene membrane filters prior to all aqueous analysis. For samples taken during anoxic half-cycles, centrifugation, filtering and subsampling were performed in an anoxic glove box (N<sub>2</sub>: H<sub>2</sub> 97:3 %, O<sub>2</sub> < 1 ppmv). Time periods were chosen to be representative of short temporal fluctuations to redox conditions experienced by surficial sediments (Aller, 1994; Nikolausz et al., 2008; Parsons et al., 2013). Similarly, the temperature and dark conditions were chosen to reflect those measured in surficial sediment during summer months at the field site. Summer conditions were chosen as this is when maximum bioturbation, microbial activity and OM input are expected within the sediment. Similar, long-running batch reactor experiments using soil or sediment have previously experienced a slowdown of metabolic processes due to depletion of labile organic carbon (Parsons et al., 2013). Therefore, gaseous carbon and nitrogen losses from the reactor were balanced by the addition of 3 g of freeze-dried and ground GFA to the suspension at the onset of each anoxic cycle. The amount of algae added was determined based on  $CO_2$  production from the reactor during the initial 11-day oxic period.

#### 2.4 Aqueous-phase methods

All reagents used were of analytical grade from Fluka, Sigma-Aldrich or Merck unless stated otherwise and were prepared with 18.2 M $\Omega$  cm<sup>-1</sup> water (Millipore). Total dissolved Na (70), K(100), Ca(20), Mg(0.5), Mn(1), Fe(3), Al(100), P (2), Si (15) and S (15) concentrations (MDL in  $\mu$ g L<sup>-1</sup>, in parentheses) were determined by ICP-OES (Thermo Scientific iCAP 6300) after filtration (< 0.45 µm) and acidification with HNO<sub>3</sub> to < pH 2. Matrix-matched standards were used for all calibrations and NIST validated multi-elemental solutions were used as controls. SRP concentrations were determined by the molybdenum blue/ascorbic acid method on a LaChat QuickChem 8500 flow injection analyzer system (4500-P E: Phosphorus by Ascorbic Acid, National Water Quality Monitoring Council, 1992; Murphy and Riley, 1962) (MDL 1.2 µg PL<sup>-1</sup>). DOC was determined using a Shimadzu TOC-LCPH/CPN analyzer (Shimadzu) following HCl addition (< pH 2) to degas dissolved inorganic carbon (MDL 71 µg CL<sup>-1</sup>).

Chloride, nitrate, nitrite and sulfate concentrations were measured by ion chromatography using a Dionex ICS 5000 equipped with a capillary IonPac<sup>®</sup> AS18 column. Aqueous sulfide was stabilized with 20  $\mu$ L 1% zinc acetate per mL (Pomeroy, 1954) after filtering and determined by the Cline method (Cline, 1969) (MDL 0.5  $\mu$ M). Fe<sup>2+</sup><sub>(aq)</sub> was determined by the ferrozine method immediately after filtering (Stookey, 1970; Viollier et al., 2000) (MDL 3.8  $\mu$ M). All aqueous analyses were conducted in triplicate. The precision and accuracy for all techniques was < 5 RSD % and ±10 % with respect to certified reference materials (where commercially available).

# 2.5 Solid-phase methods: phosphorus and iron speciation

Phosphorus partitioning within the solid phase in the reactor experiment was evaluated over a time series by both sequential extractions, using a modification (Baldwin, 1996) of the SEDEX extraction scheme (Ruttenberg, 1992) and solution <sup>31</sup>P NMR spectroscopy (Cade-Menun, 2005). The two approaches are complementary; <sup>31</sup>P NMR spectroscopy provides information on the molecular speciation of phosphorus, while sequential extraction provides information on the association of the P species with operationally defined solid-phase fractions. Therefore the combination of these two methods reveals redistribution of P within the solid phase over time during oxic–anoxic transitions.

The original SEDEX extraction scheme quantifies five different P reservoirs within sediment by consecutively solubilizing progressively more recalcitrant phases by using extracts of increasing severity. The reaction mechanisms associated with each extraction step are discussed in detail within Ruttenberg (1992). A modification of the SEDEX extraction scheme proposed by Baldwin (1996), used here, incorporates an additional 16 h, 1 M NaHCO<sub>3</sub> step (P<sub>Hum</sub>) after the P<sub>Ex</sub> step, to differentiate OM-associated P which may otherwise be co-extracted during the P<sub>Fe</sub> step. The pH of the NaHCO<sub>3</sub> extraction step was adjusted to 7.6 to minimize dissolution of carbonates prior to the P<sub>CFA</sub> extraction step. A total of 15 samples between days 11 and 74 of the reactor experiment were analysed in duplicate by sequential extraction. A sum-

Step	Extractant	Conditions	Target phase
1a	1 M MgCl <sub>2</sub>	pH 8 for 2 h at 25 °C	Exchangeable or loosely
1b	$1 \text{ M MgCl}_2$	pH 8 for 2 h at 25 °C	sorbed $P(P_{Ex})$
1c	$18.2 {\rm M}\Omega {\rm cm}^{-1} {\rm H}_2{\rm O}$	2 h at 25 °C	
2a	1 M NaHCO <sub>3</sub>	pH 7.6 for 16 h at 25 °C	Organic associated P
2b	1 M NaHCO <sub>3</sub>	pH 7.6 for 2 h at 25 °C	(P <sub>Hum</sub> )
2c	1 M NaHCO <sub>3</sub>	pH 7.6 for 2 h at 25 °C	
2d	1 M NaHCO <sub>3</sub>	pH 7.6 for 2 h at 25 °C	
2e	1 M MgCl <sub>2</sub>	pH 8 for 2 h at 25 °C	
3a	0.3 M Na <sub>3</sub> citrate with 1 M NaHCO <sub>3</sub> with 1.125 g of Na dithionite (CDB)	pH 7.6 for 8 h at 25 °C	Fe-bound P (P <sub>Fe</sub> )
3b	CDB	pH 7.6 for 2 h at 25 °C	
3c	1 M MgCl <sub>2</sub>	pH 8 for 2 h at 25 °C	
4a	1 M Na acetate with acetic acid	pH 4 for 6 h at 25 °C	Authigenic carbonate fluorapatite
4b	1 M Na acetate with acetic acid	pH 4 for 2 h at 25 °C	plus biogenic apatite plus
4c	1 M MgCl <sub>2</sub>	pH 8 for 2 h at 25 °C	CaCO <sub>3</sub> -bound P (P <sub>CFA</sub> )
5	1 M HCl	16 h at 25 °C	Detrital apatite plus other inorganic P (P <sub>Detri</sub> )
6	1M HCl	16 h after ashing at 550 °C at 25 °C	Residual/organic P (P <sub>Resi</sub> )

**Table 1.** Summary of the modified SEDEX sequential extraction protocol used on solid samples taken over a time series during the reactor experiment. Results of this extraction are shown in Fig. 4.

mary of the full sequential extraction method used, including target phases, reactants, pH, temperature and reaction times is provided in Table 1.

Changes to iron speciation were also evaluated through a time series during the experiment. To account for surfacesorbed or freshly precipitated Fe, total Fe<sup>2+</sup> production during anoxic half-cycles was estimated by a partial extraction (1 h, 0.5 N HCl) on sampled suspensions. Fe<sup>2+</sup> / Fe<sup>3+</sup> ratios were determined in extracts using a modification of the ferrozine method (Stookey, 1970; Viollier et al., 2000). Additionally, a thermodynamic model was implemented in PHREEQC (Parkhurst et al., 1999) to assess the saturation index (SI) of various minerals over time during the reactor experiment using measured pH, temperature,  $E_h$  and concentration data.

# 2.6 NaOH-EDTA extraction and solution <sup>31</sup>P NMR spectroscopy

Molecular changes to P speciation were evaluated over a time series by solution <sup>31</sup>P NMR. Phosphorus was extracted directly from suspension samples (~2g dry weight equivalent) prior to <sup>31</sup>P NMR analysis. The method used has been shown to allow quantitative analyses of P<sub>o</sub> (monoesters and diesters), polyphosphates and orthophosphates (Amirbahman et al., 2013; Cade-Menun et al., 2006; Cade-Menun and Preston, 1996; Reitzel et al., 2007; Turner et al., 2003b). Briefly, samples were extracted in 25 mL of 0.25 M (NaOH) and Na<sub>2</sub>EDTA (0.05 M) at ambient laboratory temperature

(~ 22 °C) for 4 h. Subsequently, the tubes were centrifuged (2300 g for 20 min), and the supernatant extracted via syringe and then neutralized with 2 M HCl to a pH of 7 to avoid the breakdown of polyphosphates during freeze-drying (Cade-Menun et al., 2006). This solution was then filtered to < 0.45  $\mu$ m. Prior to freeze-drying 1 mL aliquots of each sample were diluted and analysed by ICP-OES spectroscopy for Al, Ca, Fe, Mg, Mn and P. The remaining extracts were frozen at -80 °C and freeze-dried for 48 h. The freeze-dried extracts were re-dissolved in 1.0 mL D<sub>2</sub>O, 0.6 mL 10 M NaOH and 0.6 mL of the NaOH–EDTA extractant solution and were allowed to stand for 10 min with occasional vortexing. Samples were centrifuged for 20 min at 2300 g, transferred to 10 mm NMR tubes and stored at 4 °C before analysis within 12 h.

Solution  ${}^{31}$ P NMR spectra were obtained using a 600 MHz spectrometer equipped with a 10 mm broadband probe. The NMR parameters were 90° pulse, 0.68 s acquisition time, 4.32 s pulse delay, 12 Hz spinning, 20 °C and 2200 to 2900 scans (3–4 h) (Cade-Menun et al., 2010). Phosphorus compounds were identified by their chemical shifts relative to an external orthophosphoric acid standard, with the orthophosphate peak in all spectra standardized to 6 ppm. Peak areas were calculated by integration on spectra processed with 10 and 7 Hz line broadening, using NUTS software (Acorn NMR, Livermore CA, 2000 edition). Peak assignments were grouped into compounds or groups of specific compound classes when direct identifications could not be made (Cade-Menun, 2005).



Figure 2. (a) Proportions of bioturbating macro-invertebrates identified in the top 18 cm. (b) Depth profiles of sampled sediment, water content weight % (inverted black triangles), bulk density (white diamonds) OM % (white squares) and carbonate % (black circles). (c) Mineralogical composition of sediments from the zone of bioturbation determined by XRD (top 12 cm).

#### 2.7 Extracellular enzyme assays

Rates of enzymatic hydrolysis of Po were estimated through extracellular enzyme activities. Degradation rates for phosphomonoesters, phosphodiesters and pyrophosphate were determined fluorometrically through use of the MUF-tagged substrates: MUF phosphate (MUP), bis(MUF)phosphate (DiMUP, Chem-Impex International) and MUF pyrophosphate (PYRO-P, Chem-Impex International) respectively. Additionally MUF  $\beta$ -D-glucopyranoside (MUGb) was used in order to compare phosphatase enzyme activity to the activity of  $\beta$ -glucosidase (cellulase) (Dunn et al., 2013). Enzyme activities were determined using a microplate reader (Flexstation3, Molecular Devices) using a modification of Deng et al. (2013). Briefly, 1 g dry weight equivalent of suspension from the reactor was stirred with 100 mL of 100 mM HEPES buffer at pH 7.5 in a pyrex dish for 10 min at 280 rpm to allow for complete homogenization. Subsamples (100 µL) of the buffered soil suspension were removed during continuous mixing using a multi-channel pipette and placed into microplate wells, which were loaded into the microplate reader. Four replicate wells were filled per substrate. Plates were left to equilibrate at 30 °C for 5 min inside the reader before the automatic addition of 100 µL of substrate, resulting in a final substrate concentration of 667 µM. Each well was triturated thoroughly during addition of the substrate. Excitation fluorescence was set at 365 nm. Emission intensity at 450 nm was recorded at 5 min intervals over a 6 h period. The effect of fluorescence quenching was accounted for in each sample by preparing MUF calibration curves in the same soil suspension as used for the analysis. The limits of detection and quantification were determined to be 1.1 and 3.3 µM MUF respectively, equivalent to  $1.1 \,\mu\text{M}$  of phosphate for the determination of phosphomonoesterase activities.

#### 3 Results and discussion

# 3.1 Sediment characterization and evidence of bioturbation

Characterization of sediment cores revealed physical and chemical solid-phase homogeneity within the top 10 cm, with a bulk density of  $\sim 1.3$  g cm<sup>-3</sup>, water content of  $\sim 50$  % (by weight), OM of  $\sim 3$  % and a carbonate of  $\sim 25$  % (Fig. 2b). Between 10 and 15 cm depth increases in bulk density and decreases to the sediment water content, OM % and carbonate fraction occurred as soft sediment transitioned to clay.

A benthic macro-invertebrate density of approximately 49 500 individuals per m<sup>2</sup> was determined, consistent with previously reported values (Pelegri and Blackburn, 1995). The community (Fig. 2a) was dominated by aquatic earthworms (*Tubificidae* 60% and *Branchiura sowerbyi* 8%), which typically feed and mix sediment within the top 5–10 cm (Fisher et al., 1980; McCall and Fisher, 1980). Other groups identified included *Ceratopogonidae* (no-see-ums or biting midges, 22%) including *Sphaeromias, Probezzia, Bezzia* and *Chironomidae* (midges, 6%) including *Cryptochironomus, Tanypus* and *Nemotoda* (round worms, 4%), and a single *Hyalella azteca* (scud < 1%).

Bioturbating organisms, such as those identified, have previously been shown to alter biogeochemical cycling within surface sediments (Hölker et al., 2015). Reported influences include increased solute fluxes (Furukawa et al., 2001; Ma-

tisoff and Wang, 1998), mixing of solid sediment (Fisher et al., 1980) and bioconveying of sediment particles (Lagauzère et al., 2009). These processes have been shown to enhance sediment oxygen demand (McCall and Fisher, 1980; Pelegri and Blackburn, 1995), degradation of OM (Aller, 1994), rates of denitrification, transport of contaminants to surface water (Lagauzère et al., 2009) and temporal fluctuation of redox conditions (Aller, 1994). Efficient sediment mixing allows frequent re-oxidation of reduced sediments and therefore regeneration of terminal electron acceptors (TEAs) such as nitrate, ferric iron and sulfate, which often limit mineralization of OM in sediments underlying hypereutrophic water bodies (Reddy and DeLaune, 2008). Electron donors in the form of fresh autochthonous necromass are also rapidly redistributed vertically within the zone of bioturbation. This environment should therefore support a metabolically diverse, abundant and highly active microbial community (DeAngelis et al., 2010).

Quantitative XRD analysis of the top 12 cm of sediment (Fig. 2c) showed close agreement with the carbonate fraction determined by TGA (~ 25 % by TGA vs. 27 % by XRD) indicating a calcite-dominated, carbonate-buffered system. The remaining mineral assemblage was dominated by quartz and clay minerals (illite 30 % and chamosite 2 %). No pyrite or vivianite was detected by XRD, suggesting either their absence or presence in low abundance (< 1 %) with poorly crystalline structures.

#### 3.2 Experimental redox oscillation: aqueous chemistry

 $E_h$  within the bioreactor oscillated between +470 and -250 mV (Fig. 3) consistent with  $E_h$  ranges of wetland sediments (Nikolausz et al., 2008). A slight pH oscillation was also present between  $\sim$  7.4 during anoxic halfcycles (N\_2 sparging) and  $\sim 7.7$  during oxic half-cycles  $(N_2:O_2:CO_2 \text{ sparging})$  this variation, shown in Fig. 3, is consistent with calcite/dolomite-buffered sediment equilibrating with changing  $p_{CO_2}$  caused by sparging gas composition and microbial respiration. After the 11-day equilibration, ionic strength of the aqueous phase in the reactor suspension remained at  $\sim 0.025 \pm 0.004$  M for the duration of the experiment. This range of  $E_h$  / pH conditions and ionic strength is consistent with the range measured within surficial sediment at the field site and transitions across the thermodynamically predicted stability boundaries for multiple redox couples, e.g.  $MnO_2 / Mn^{2+}$ ,  $NO_3^- / NO_2^- / NH_4^+$ ,  $Fe(OH)_3 / Fe^{2+}$ ,  $SO_4^{2-} / HS^-$ , during each 14-day redox cycle. The upper  $E_h$  values recorded during oxic cycles are significantly lower than predicted by the  $O_2 / H_2O$  couple (820 mV at pH 7) but are consistent with the  $O_2 / H_2O_2$  couple (300 mV at pH 7), which is considered to control electrode measured  $E_h$  under oxic conditions (Stumm and Morgan, 1996).

Aqueous chemistry data, shown in Fig. 3, demonstrate the consumption of TEAs in order of decreasing nominal en-



**Figure 3.** Aqueous chemistry and iron extraction data with time during reactor experiments: the solid red line is  $E_h$ , solid black line is DO, solid blue line is pH, full black circles are NO<sub>3</sub><sup>-</sup>, white squares are NO<sub>2</sub><sup>-</sup>, inverted red triangles are Fe<sub>(aq)</sub>, inverted white triangles are Mn<sub>(aq)</sub>, red triangles are Fe<sup>2+</sup> 0.5 M HCl extractable, white triangles are Fe<sup>2+</sup> / Fe<sup>3+</sup> ratio in 0.5 M HCl extract, red circles are S<sub>2</sub><sup>-</sup> and black diamonds are DOC. Sampling points for <sup>31</sup>P NMR and extracellular enzyme assays (EEA) are shown on the  $E_h$  curve (<sup>31</sup>P NMR is indicated by open blue squares and EEA by open green circles).

ergetic yield, coupled to the oxidation of labile OM. Upon physical removal and consumption of residual oxygen by aerobic respiration, nitrate concentration decreased in the solution. Decreases to nitrate concentration coincided with peaks of nitrite concentration within the first hour of oxygen removal, indicative of microbial denitrification. Subsequent increases to  $Mn_{(aq)},\,Fe_{(aq)}^{2+}$  and  $HS_{(aq)}^{-}$  imply sequential reduction of MnO<sub>2</sub>, Fe(OH)<sub>3</sub> and SO<sub>4</sub><sup>2-</sup> as more energetically efficient electron acceptors were depleted. Mn (predicted as  $Mn^{2+}$  by the thermodynamic model) and  $Fe^{2+}$  were detected in solution earlier within each subsequent anoxic cycle, but the apparent order of reduction remained consistent across all five redox cycles  $(O_2, NO_3^-, NO_2^-, MnO_2, Fe(OH)_3, SO_4^{2-})$ . The consistent order and relative magnitude of reduction imply that the main biogeochemical functioning of the sediment suspension did not change dramatically between cycles during the experiment.

Although relatively low concentrations of  $Fe^{2+}$  (up to 71 µM) were measured in solution, 0.5 M HCl extractions revealed that significantly greater Fe<sup>2+</sup> was produced during each anoxic cycle than was measured in the aqueous phase. Fe<sup>2+</sup> generated by dissimilatory iron reduction has been shown to sorb to mineral surfaces in sediment (Gehin et al., 2007; Klein et al., 2010; Liger et al., 1999) or precipitate as ferrous carbonate (Jensen et al., 2002), ferrous sulfide or other mixed ferrous/ferric phases (Rickard and Morse, 2005). The 0.5 M HCl extractions targeted this sorbed or poorly crystalline freshly precipitated Fe<sup>2+</sup>. During each anoxic cycle HCl extractable Fe<sup>2+</sup> concentration increased by 50-70  $\mu$ mol g<sup>-1</sup>, equivalent to 12.31 to 17.29 mM of iron reduction within the reactor as a whole. Thus, only  $0.63\pm0.4$  % of microbially reduced Fe<sup>2+</sup> was measurable in solution. The aqueous phase of the reactor was shown to be supersaturated with respect to mackinawite (FeS), pyrite (FeS<sub>2</sub>), vivianite (Fe<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>: 8H<sub>2</sub>O) and siderite (FeCO<sub>3</sub>) during anoxic halfcycles, indicating thermodynamic favorability for precipitation of these minerals. The kinetic constraints on precipitation were not, however, considered. No significant cumulative change to extractable  $Fe^{2+}$  /  $Fe^{3+}$  occurred after five full reduction-oxidation cycles, indicating that solid-phase Fe redox cycling was reversible, potentially due to rapid oxidation of solid  $Fe^{2+}$  in the presence of  $O_2$  and carbonate (Caldeira et al., 2010).

DOC concentration also fluctuated systematically during oxic-anoxic cycles (Fig. 3). Higher DOC concentrations were measured during anoxic conditions than oxic conditions. DOC may be replenished by both enzymatic hydrolysis of particulate organic matter (Vetter et al., 1998) and desorption of mineral-associated OM (Grybos et al., 2009). The addition of algal matter at the beginning of anoxic cycles resulted in observable sharp peaks of DOC, which was rapidly removed from solution, probably due to a combination of mineralization of labile DOC to HCO<sub>3</sub><sup>-</sup> and sorption processes (Chorover and Amistadi, 2001; Grybos et al., 2009). The peak of DOC supplied by addition of algal matter represented labile DOC, which was readily mineralized in comparison to the residual DOC, which persisted in the system throughout the experiment. The differences in residual OM mobility between oxic and anoxic cycles were unlikely due to oxide dissolution as differences to DOC concentration are observed prior to increases in Mn and Fe concentration in solution. We therefore postulate that solubility changes to humified DOC were driven by pH changes (Fig. 3) between oxic and anoxic conditions caused by changes in  $pCO_2$  between oxic and anoxic conditions as previously shown in wetland sediments (Grybos et al., 2009).

Lowest aqueous phosphorus concentrations ( $\sim 2.5-3 \,\mu$ M), shown in Fig. 4, occurred during oxic half-cycles and increased dramatically during anoxic half-cycles to a maximum concentration of 50–60  $\mu$ M per cycle, 88 % of which occurred as SRP. The range of TDP concentrations within the aqueous phase of the reactor suspension is similar to that re-

Figure 4. Aqueous- and solid-phase phosphorus speciation from sequential chemical extractions with time during the reactor experiment. White panels correspond to time periods with air sparging, and grey panels correspond to time periods with  $N_2 : CO_2$  sparging. Black symbols are total P concentration, and white symbols are SRP concentration.

ported in situ at the field site by Mayer et al. (2006). The timing of phosphorus release to the aqueous phase corresponded well with increasing  $Fe_{(aq)}^{2+}$  concentration. This is reflected in a strong positive correlation between TDP and Fe concentrations (n = 37, R<sup>2</sup> = 0.93, p < 0.0001). Increases to aqueous P concentration occurred only after depletion of residual O<sub>2</sub>, NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup>, after increases to Mn<sub>(aq)</sub> and before increases to HS<sup>-</sup><sub>(aq)</sub>. The timing of P release is suggestive of an iron(III)-oxyhydroxide or ferric phosphate control on phosphorus mobility (Bonneville et al., 2004; Hyacinthe and Van Cappellen, 2004) and indicates that complete nitrate depletion was required prior to phosphorus release to the aqueous phase.

# 3.3 Sequential chemical extractions and solid-phase P partitioning

The sum of P concentrations from all sequential extraction reservoirs  $(61 \pm 5 \,\mu\text{mol}\,g^{-1})$  was consistently within 10%



of a total P extraction  $(57 \pm 4 \,\mu\text{mol g}^{-1})$ , indicating acceptable analytical precision from the sequential extraction procedure. Highest P concentrations were associated with the P<sub>Hum</sub> (~26 % of TP) and P<sub>Fe</sub> (~24 %) fractions with lower P concentrations in the P<sub>Ex</sub> ~16 %, P<sub>CFA</sub> ~15 %, P<sub>Detr</sub> ~12 % and P<sub>Res</sub> ~7 % fractions (Fig. 4). The largest variations in P concentration were observed for P<sub>Fe</sub> and decreased in the order P<sub>Fe</sub> (2.6) > P<sub>Hum</sub> (1.5) > P<sub>Ex</sub> (0.4) > P<sub>Detr</sub> (0.4) > P<sub>Resi</sub> (0.35) > P<sub>CFA</sub> (0.25), where the numbers in parentheses are standard deviations (µmol g<sup>-1</sup>). High variability over time within the P<sub>Fe</sub> and P<sub>Hum</sub> fractions suggests that P was exchanged to and from these fractions during redox oscillation, whereas changes to the P<sub>Ex</sub>, P<sub>Resi</sub> and P<sub>CFA</sub> fractions were comparatively insignificant (Fig. 4).

The P<sub>Fe</sub> fraction was the only P pool in which concentration consistently decreased during anoxic conditions and increased during oxic conditions (Fig. 4). When a P mass balance (Fig. 5) was attempted to account for increases to aqueous phosphorus (P<sub>Aq</sub>) from the iron-bound (P<sub>Fe</sub>) pool during anoxic periods, it became evident that only approximately 4.5% of variability observed in the P<sub>Fe</sub> pool (total P<sub>Fe</sub> variation of up to 4.5 µmol g<sup>-1</sup> during anoxic periods) was necessary to account for the changes to inter-cycle P<sub>Aq</sub> concentrations (50 µM). The remainder of P<sub>Fe</sub> lost during anoxic cycles appears to be reversibly redistributed to the P<sub>Ex</sub> (~30%) and P<sub>Hum</sub> (~65%) pools, which generally increased during anoxic conditions and decreased during oxic conditions (Fig. 4).

According to Ruttenberg (1992) the  $P_{Ex}$  fraction corresponds to P mobilized via the formation of  $MgPO_4^-$  complexes and(or) mass action displacement by Cl<sup>-</sup>. It is therefore considered that MgCl<sub>2</sub> effectively extracts P loosely associated with mineral surfaces. However, Ruttenberg (1992) also demonstrated that plankton were efficiently extracted by MgCl<sub>2</sub> as well as  $\sim 25 \%$  of P associated with biogenic CaCO<sub>3</sub>. Consequently, it is likely that P associated with microbial biomass, CaCO<sub>3</sub>, and other loosely sorbed P contributed to the PEx fraction. Slight increases to PEx concentration during anoxic periods and corresponding decreases during oxic periods likely reflect the combination of two processes: (1) equilibration between P surface complexes and aqueous P species due to fluctuating aqueous concentrations which were consistently higher during anoxic periods (Olsen and Watanabe, 1957); and (2) pH-driven sorption/desorption as pH was consistently slightly higher during aerobic periods (Fig. 3), favouring desorption from mineral surfaces including illite, which comprised 30 % of the crystalline mineralogical fraction (Fig. 2c) (Manning, 1996).

The NaHCO<sub>3</sub> extraction step was originally added to the SEDEX method to target OM-associated P, which would otherwise be liberated during the  $P_{Fe}$  extraction step (Baldwin, 1996). Baldwin noted a brown coloration in  $P_{Hum}$  extracts and that absorbance at 250 nm was positively correlated with SRP. Absorbance at 254 nm has been shown to be indicative of aromatic OM, commonly associated with humic sub-



**Figure 5.** Change in P distribution between the start and end of each oxic and anoxic period (7-day change). Iron-bound P (P<sub>Fe</sub>) appears to be reversibly redistributed to the loosely sorbed (P<sub>Ex</sub>) and humic-bound (P<sub>Hum</sub>) and aqueous fractions (P<sub>Aq</sub>). The dashed line is 1 : 1. Linear regression of the data results in an  $R^2$  of 0.95, a slope of -1.1 and p < 0.0001).

stances (Weishaar et al., 2003). A light brown colour was also present in the NaHCO<sub>3</sub> extracts recovered during this experiment despite comparatively low sediment OM content (Fig. 2b). Absorbance spectra for these extracts were not determined.

Li et al. (2015) have recently demonstrated that the SEDEX P<sub>Fe</sub> extraction step co-extracted P associated with fine iron oxide-OM complexes when a prior NaHCO3 step was not incorporated. However, Li et al. (2015) also suggested that these complexes may be more recalcitrant than pure minerals. Iron was present within the P<sub>Hum</sub> extract at concentrations between 18 and  $25 \,\mu mol g^{-1}$ , but the original speciation of this iron is prior to extraction is unknown. Chemically similar extractions used in soil sciences such as Hedley's extraction (0.5 M NaHCO<sub>3</sub>, pH 8.5, 16 h) and the Olsen P test (0.5 M NaHCO<sub>3</sub>, pH 8.5, 30 min) have been shown to extract Mg and Ca phosphates as well as some organic P (Hedley et al., 1982; Olsen et al., 1954). Approximately two-thirds of P extracted within P<sub>Hum</sub> was present as SRP, suggesting that approximately one-third of this fraction may be P<sub>o</sub> species. The pH of the NaHCO<sub>3</sub> extract used here was adjusted to 7.6 to minimize the dissolution of Mg and Ca phosphates prior to the P<sub>CFA</sub> extraction. Despite this, 30-44 µmol g<sup>-1</sup> of Ca was extracted within the P<sub>Hum</sub> fraction. The origin of the extracted Ca is, however, unknown and may have been complexed with OM or part of labile Ca phosphates. It is still expected that the majority of Ca-phosphate minerals were quantified as part of the PCFA or PDetri extractions, which included  $\sim$  500–700 and  $\sim$  50–70 µmol Ca g<sup>-1</sup> respectively.

Humic acids are known to compete with orthophosphate for surface-binding sites on various minerals, including goethite (Sibanda and Young, 1986) and poorly ordered Feoxides in the short term (Gerke, 1993). However, sorption of natural OM to freshly precipitated Fe oxides may increase the long-term sorption capacity of ferric oxides towards P by decreasing recrystallization over time (Gerke, 1993) and through the formation of OM-Fe-P complexes (Gerke, 1993). Although previous studies have provided evidence for ternary complexes between ferric iron, OM and phosphate (Kizewski et al., 2010b) there is currently no direct spectroscopic evidence for the existence of mixed OM-Fe(III)-phosphate complexes in natural waters. Identification of such complexes in natural environments is inherently challenging due to the complexity of natural geochemical matrices (Kizewski et al., 2010a). Recent studies have however successfully investigated the structure of synthetic OM-Fe(III)-phosphate complexes (Kizewski et al., 2010a) and similar OM-Fe(III)-arsenate complexes spectroscopically (Mikutta and Kretzschmar, 2011; Sharma et al., 2010). These studies suggest that similar and perhaps more complex heterogeneous ternary complexes are also likely to be present in natural freshwater environments (Kizewski et al., 2010a). This suggestion is also supported by the observation that more than 80% of soluble P in some natural waters is associated with high molecular weight OM (Gerke, 2010). As spectroscopic characterization of the P associated with the P<sub>Hum</sub> fraction was not performed in this study, the P<sub>Hum</sub> pool is considered to represent a variety of OM-associated P in addition to small amounts of P from labile Ca-phosphate minerals. OM-associated P extracted within this fraction is likely coordinated with ferric iron  $(18-25 \,\mu\text{mol Fe g}^{-1} \text{ co-}$ extracted), Ca (30–44  $\mu$ mol Ca g<sup>-1</sup> co-extracted) or Al (0.8– 1.5  $\mu$ mol Al g<sup>-1</sup> co-extracted). These metals may, in turn, be associated with various mineral surfaces within the sediment.

Sequential extraction data demonstrate that the  $P_{Hum}$  fraction is the dominant P fraction in all samples analysed, which highlights the significance of this fraction. P mass balance also suggests that reversible re-partitioning between this and the  $P_{Fe}$  fraction occurs during redox condition changes. As the exact chemical nature of the  $P_{Hum}$  fraction is not known, interpretation of concentration changes over time are challenging. Speculatively, increases to the  $P_{Hum}$  fraction under anoxic conditions may be due to the release of occluded OM–metal–P complexes within iron(III) oxyhydroxides during reductive dissolution or simply re-equilibration of solid-phase OM–metal–P complexes with increased aqueous P.

We consider that the majority of P extracted within the  $P_{Fe}$  fraction was co-precipitated with iron(III) oxyhydroxides which were reductively dissolved by dithionite during the extraction (Ruttenberg, 1992). This interpretation is supported by the relatively high concentrations of iron extracted within the  $P_{Fe}$  fraction (72–91 µmol Fe g<sup>-1</sup>). Aqueous Fe<sup>2+</sup> produced during this extraction is subsequently chelated by citrate and therefore solubility of Fe and P is maintained. The bicarbonate component functions as a pH buffer to ensure maximum preservation of apatite and CaCO<sub>3</sub>-bound P during this reaction step (Ruttenberg, 1992). Neither the  $P_{CFA}$ ,  $P_{Detri}$  nor  $P_{Resi}$  fractions varied systematically between oxic and anoxic conditions or changed consistently during the course of the experiment, suggesting their comparative stability during short periods of redox fluctuation. This is supported by calculated SI for hydroxyapatite, which remained between +0.86 and +6.24 for the duration of the experiment.

The P contribution from individual algal additions  $(\sim 1.5 \,\mu\text{mol} \,P \,g^{-1})$  was relatively small compared to total P in the reactor (61  $\mu$ mol P g<sup>-1</sup>) and within the margin of analytical error associated with solid extractions. Additionally, no single fraction shows a clear increase over the course of the experiment, and therefore quantification of the redistribution of P added with algal additions is not possible.

#### 3.4 Fe: P ratios

Sequential extraction data (shown in Fig. 4), aqueous chemistry data (shown in Fig. 3) and the correlation between aqueous Fe and P (n = 37,  $R^2 = 0.93$ , p < 0.0001) all suggest that P released to solution under anoxic conditions originated in the PFe pool. Although the maximum molar ratio for phosphate incorporation within ferric oxides has been shown to be 2:1 (Fe: P) (Thibault et al., 2009), it has been suggested that much higher solid Fe : P of 15 (Jensen et al., 1992) to > 20(Phillips et al., 1994) may be necessary to control phosphorus mobility under oxic conditions. Results from bioreactor experiments suggest that phosphorus is retained in the solid phase under oxic conditions at total Fe : P ratios of just 4.1 : 1, potentially due to the association of P with other solid sedimentary pools, particularly P<sub>Hum</sub>. Fe : P ratios below the stoichiometric limitation of 2:1, measured in the aqueous phase (1.5-1.9), during anoxic conditions are therefore likely due to the removal of  $Fe^{2+}$  from solution by secondary sorption and precipitation processes, subsequent to reductive dissolution. Probable secondary Fe<sup>2+</sup> removal processes include the formation of amorphous FeS (SI of up to +2.27 for mackinawite) (Rickard and Morse, 2005) and sorption of  $Fe^{2+}$  to clays (Gehin et al., 2007; Klein et al., 2010). This is supported by increases to  $Fe^{2+}$  /  $Fe^{3+}$  ratios in the 0.5 M HCl extractable fraction during anoxic conditions. Frequent rapid reoxidation of ferrous sulfide due to air sparging in the reactor experiment, and extensive in situ bioturbation, likely prevents formation of more recalcitrant and slow-forming iron sulfide minerals such as pyrite, despite strong thermodynamic favorability for pyrite formation (SI up to +13.13) (Caldeira et al., 2010; Peiffer et al., 2015). This is consistent with the results of XRD analysis, which did not identify pyrite (Fig. 2c). A similar effect has been previously demonstrated in lake sediments (Gächter and Müller, 2003).

#### 3.5 Hydrolytic enzyme activities

The activities of model phosphomonoesterase, phosphodiesterase and pyrophosphatase were found to be systemati-



**Figure 6. (a)** P speciation determined by <sup>31</sup>P NMR for the initial suspension (top left), the final suspension (bottom left) and the average of samples from oxic conditions n = 2 (top right), the average of samples from anoxic conditions n = 3 (bottom right). Pho-P is phosphonates, Di-P is diester P and Mon-P is monoester P. Polyphosphate was not detected at concentrations > 1% in any of the samples analysed. (b) Average extracellular enzyme activities under oxic and anoxic conditions for MUP, DiMUP, PYRO-P and MUGb (n = 5).

cally higher under oxic conditions compared to anoxic conditions by 37 % (p < 0.005), 8 % (not significant) and 24 % (p=0.08) respectively (Fig. 6b). Phosphomonoesterases were found to have the highest activities despite the inherent overestimation of phosphodiesterase activities when using MUF-tagged substrates (Sirová et al., 2013). The opposite trend was observed for glycopyranoside, part of the cellulose degradation pathway (Dunn et al., 2013), which showed consistently higher activity (69 % p = < 0.05) under anoxic conditions (Fig. 6b). The different trends exhibited by cellulose and phosphatase enzymes indicate that changes in activity were not universal but specific to enzyme function. Phosphomonoesterase activities obtained in the current study  $(1.76-2.4 \text{ mmol } \text{h}^{-1} \text{kg}^{-1})$  are similar to those previously reported in wetland sediments (Kang and Freeman, 1999) and suggest the capacity for rapid hydrolysis of Po species in necromass. Lowering of the water table in wetlands has previously been shown to increase the activity of phosphatase enzymes and the hydrolysis of Po species (Song et al., 2007). However, water table fluctuation results in concomitant changes to moisture content and redox conditions, which prevents isolation of the causal variable in field investigations (Rezanezhad et al., 2014). Therefore, this is, to the best of our knowledge, the first direct demonstration of phosphatase activity changes in response to changing redox conditions. We postulate that under anoxic conditions, when phosphorus availability in the aqueous phase is high, production of extracellular phosphatase enzymes by the microbial community is down-regulated. Conversely, when bioavailable phosphorus is removed from solution under oxic conditions, extracellular phosphatase production is up-regulated in response. Adjustments to enzyme production in response to changes in phosphate availability must occur on short timescales (hours/days) for such trends to be observable during the bioreactor experiment. An inverse relationship between phosphatase activities and phosphate concentration has previously been shown spatially in wetlands by Kang and Freeman (1999) but to our knowledge never temporally in sediments.

#### 3.6 <sup>31</sup>P NMR

Results from <sup>31</sup>P NMR analyses (Fig. 6a) show that the majority of phosphorus was present in the solid phase as orthophosphate (84-91 %) with 4-8 % monoester P, 3-8 % diester P and < 1% phosphonates and polyphosphates with no clear trend in relative abundance emerging during the experiment. The NaOH-EDTA extraction resulted in a recovery of  $\sim 27$  % of TP, which is comparable to previous studies with carbonate-buffered soils and sediments (Hansen et al., 2004; Turner et al., 2003a). Alpha and beta glycerophosphates are commonly identified in monoester spectral regions and have been demonstrated to be products of diesters degraded during analysis (Doolette et al., 2009; Jørgensen et al., 2015; Paraskova et al., 2015). As no glycerophosphates were identified in any of the analysed samples, recalculation of monoester/diester ratios was not performed. A higher mean monoester / diester ratio (2.31) was found in reduced samples than oxidized samples (0.97) (a statistically significant difference, p = 0.04). This difference could indicate either that monoester P was less efficiently extracted under oxic conditions due to sorption to metal oxides or that monoesterase/diesterase activity decreased under anoxic conditions, which is consistent with enzymatic activity assays (Fig. 6b). Total P<sub>o</sub> determined by <sup>31</sup>P NMR varied between 9 and 16% over time compared to 5–11% in the P<sub>resi</sub> from sequential extractions, indicating that not all P<sub>o</sub> was extracted in the P<sub>resi</sub> fraction, which is commonly referred to as the organic-P fraction. We postulate that the remaining ~ 5% of total phosphorus, identified as P<sub>o</sub> by <sup>31</sup>P NMR, was extracted during previous steps in the sequential extraction scheme, particularly MgCl<sub>2</sub>, which has been shown to efficiently extract P associated with microbial biomass (Ruttenberg, 1992) and NaHCO<sub>3</sub>. The relative activities of phosphatase enzymes appear to correlate with the relative abundances of P<sub>o</sub> species identified by <sup>31</sup>P NMR, e.g. monoesters > diesters > pyro-P.

Significant polyphosphate (>1%) was not detected by <sup>31</sup>P NMR during experiments. Previous studies focusing on WWTP tertiary treatment for phosphate removal suggest that redox oscillating conditions promote intracellular poly-P accumulation during aerobic conditions to be used as an energy store under anoxic conditions in order to uptake short-chain fatty acids (SCFA) in the absence of an electron acceptor (Hupfer et al., 2007; Wentzel et al., 1991). Phosphate uptake during aerobic conditions therefore requires P availability in excess of what is required for growth and maintenance of the microbial community. However, phosphate availability under aerobic conditions is limited by sorption and co-precipitation with iron oxides, assuming sufficient Fe : P, despite high total phosphorus concentration in sediment. The P requirements by the microbial community are also likely to be high during the transition to aerobic conditions due to the availability of O2 as an energetically efficient electron acceptor and fermentation products (SCFA), further decreasing the probability of polyphosphate accumulation. Additionally, polyphosphate accumulation and release has shown to be inhibited by denitrification and sulfate reduction due to competition for SCFA (Kortstee et al., 1994; Yamamoto-Ikemoto et al., 1994).

#### 4 Implications

Our controlled laboratory simulation of highly dynamic redox conditions in eutrophic sediment demonstrates the importance of multiple coupled elemental cycles (C, N, Fe, S, P) when determining internal P loading potential and timing. Our results demonstrate that neither aqueous- nor solid-phase Fe: P ratios or even solid-phase PFe quantification are good predictors of potential P release to the water column under anoxic conditions due to extensive reversible redistribution of both reduced Fe and associated P within the solid phase. We show that 99.4 % of reduced Fe and 95.5 % of  $P_{Fe}$  are not released to the aqueous phase upon Fe reduction but reversibly redistributed within the solid phase upon short periods of iron reduction. Additionally, the apparent requirement for complete nitrate depletion prior to anoxia-promoted P release to the aqueous phase has potential implications for water bodies where significant iron-bound legacy P is present within sediments. Our results suggest that decreasing  $NO_3^$ concentrations in external loads, while ostensibly ecologically beneficial, may, in some cases, increase the frequency and magnitude of internal P loading during short periods of anoxia. In P limited systems, the apparent ecological benefits of decreased  $NO_3^-$  may be offset by increased P release and eutrophication. However, numerous additional processes exist in natural systems, which were not simulated during our reactor experiment and may influence internal loading mechanisms. Finally, we demonstrate that oscillatory redox conditions, even in sediments with diverse and active microbial communities, do not necessarily result in accumulation of polyphosphate due to mineralogical phosphate immobilization and scavenging of SCFA by anaerobic heterotrophic respiration.

*Data availability.* Full tabulated aqueous- and solid-phase chemistry data for the experimental time series is provided in the Supplement.

#### **Appendix A: Abbreviations**

SWIsediment-water interfaceDOCdissolved organic carbonPoorganic phosphorus speciesSRPsoluble reactive phosphorusICP-OESinductively coupled plasma optical emission spectrometryOMorganic matterXRDpowder X-ray diffractionMDLmethod detection limitTDPtotal dissolved phosphorusRSD %relative standard deviationMUF4-methylumbelliferylMUP4-Methylumbelliferyl phosphateDiMUPbis(4-methylumbelliferyl)phosphatePYRO-P4-methylumbelliferyl beta-D-glucopyranosideTGAthermogravimetric analysisTPtotal phosphorus
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### The Supplement related to this article is available online at https://doi.org/10.5194/bg-14-3585-2017-supplement.

*Competing interests.* The authors declare that they have no conflict of interest.

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