Phosphorus export from a restored wetland ecosystem in response to natural and experimental hydrologic fluctuations

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Wetland restoration is a commonly used approach to reduce nutrient loading to freshwater and coastal ecosystems, with many wetland restoration efforts occurring in former agricultural fields. Restored wetlands are expected to be effective at retaining or removing both nitrogen and phosphorus (P), yet restoring wetland hydrology to former agricultural fields can lead to the release of legacy fertilizer P. Here, we examined P cycling and export following rewetting of the Timberlake Restoration Project, a 440 ha restored riverine wetland complex in the coastal plain of North Carolina. We also compared P cycling within the restored wetland to two minimally disturbed nearby wetlands and an adjacent active agricultural field. In the restored wetland we observed increased soluble reactive phosphorus (SRP) concentrations following initial flooding, consistent with our expectations that P bound to iron would be released under reducing conditions. SRP concentrations in spring were 2.5 times higher leaving the restored wetland than a forested wetland and an agricultural field. During two large-scale drawdown and rewetting experiments we decreased the water depth by 1 m in ~10 ha of inundated wetland for 2 weeks, followed by reflooding. Rewetting following experimental drainage had no effect on SRP concentrations in winter, but SRP concentrations did increase when the experiment was repeated during summer. Our best estimates suggest that this restored wetland could release legacy fertilizer P for up to a decade following hydrologic restoration. The time lag between restoration and biogeochemical recovery should be incorporated into management strategies of restored wetlands.


1. Introduction

Wetland restoration is increasingly used to reduce nutrient loading of downstream freshwater and coastal ecosystems [Verhoeven et al., 2006], with much wetland restoration occurring in marginally productive agricultural fields [Zedler, 2003], and supported by government programs in both the United States [Zedler, 2003] and Europe [Hansson et al., 2005]. Restoring the ability of wetlands to retain or transform nutrients in agricultural lands presents various challenges due to the legacy of disturbances to soil and hydrology associated with agriculture, including loss of soil organic matter, soil compaction, fertilizer application, drainage and pumping [Zedler, 2003]. Moreover, wetland restoration in former agricultural fields can create biogeochemical conditions that mobilize nutrients, an unintended consequence of such programs.

Many wetland restoration projects seek to reduce both nitrogen (N) and phosphorus (P) in surface water [Verhoeven et al., 2006], however the biogeochemical conditions necessary to promote denitrification (which removes N from surface waters) might facilitate P release. P availability in wetlands is primarily controlled by geochemical sorption onto soil minerals and biotic uptake and mineralization [Richardson, 1985]. Adsorption of inorganic P onto noncrystalline Al and Fe oxides is the dominant mechanism of P removal in wetlands with acidic soils [Richardson, 1985]. Predicting the effect of flooding on P cycling is challenging because reducing conditions can affect P sorption capacity both positively and negatively. Flooding promotes the creation of noncrystalline forms of Fe and Al and organic matter-Al complexes, increasing P sorption capacity [Darke and Walbridge, 2000]. In contrast, flooding and anoxic conditions can lead to the reduction of
Fe(III) to Fe(II), which releases P bound to oxidized Fe minerals in the soil [Reddy et al., 1999]. Droughts, and associated oxic conditions, can enhance mineralization of organic forms of P, leading to mobilization of bioavailable forms of P [Van Dijk et al., 2004].

[4] Restoring wetland hydrology to former agricultural fields can lead to the mobilization of legacy P, also called memory P, accumulated from fertilizer application [Reddy and DeLaune, 2008]. The release of this legacy P can be due to both the reduction of Fe (III) during anoxic conditions leading to the release of iron-bound P and to mineralization of organic P under oxic conditions during dry periods. Various laboratory and short-term studies have reported increases in P dissolution from soils due to flooding and drainage cycles of former agricultural lands [Aldous et al., 2005, 2007; Bostic and White, 2007; Corstanje and Reddy, 2004; Pant and Reddy, 2003; Van Dijk et al., 2004] and in response to storms [Novak et al., 2007]. Much of this research has relied mostly on laboratory or lysimeter-scale measurements of P from soils. There has been limited research in situ on the relationships between fluctuating water tables in a restored wetland and P export (but see Ardón et al. [2010] and Duff et al. [2009]).

[5] We previously reported water and nutrient (NH₄⁺, NO₃⁻, TDN, DON, TP and SRP) budgets following the reflooding of a former agricultural wetland as part of stream and wetland restoration in the Timberlake Restoration Project (TLRP), in the coastal plain of North Carolina [Ardón et al., 2010]. In the first 2 years following hydrologic reconnection, a mass balance for P for this large restored wetland documented a release of 0.6 kg TP ha⁻¹ yr⁻¹ and an 8X increase in annual TP export relative to estimated prerestoration export [Ardón et al., 2010]. P export from this site is important because the downstream Albemarle Sound is considered to be P limited, with SRP concentrations usually being less than 0.04 mg/L [Lin et al., 2007; Richardson, 1983]. Here, we examined how natural and experimental hydrologic fluctuations determine P cycling within a restored wetland. We anticipated that: (1) we would measure release of P immediately upon initial reflooding a former agricultural field as part of restoration and during storm events; (2) concentrations and export of SRP and TP from the restored wetland would be higher than a forested wetland but lower than an agricultural field; (3) soil TP in restored wetland would decline over time due to increased P export; and (4) large drawdown experiments would increase P export from the restored wetland. To elucidate the recovery trajectory of this restored wetland we also compared soil parameters and soil solution P to two nearby minimally impacted wetlands and an active agricultural field. Due to the 20 years of agricultural activities in TLRP, we expected to see higher soil solution P in the restored wetland than the reference wetlands and soil parameters from TLRP to resemble more an active agricultural field than reference wetlands.

2. Site Description

2.1. Albemarle Peninsula

[6] Located in the northern coastal province of NC, the Albemarle Peninsula extends 5000 km², with 2700 km² under 1 m elevation [Poulter and Halpin, 2008]. The Peninsula is located between the Albemarle Sound (north), Croatan Sound (east) and Pamlico Sound (south). Mean annual temperature is 16.6°C and mean annual precipitation is 1330 mm/yr (Plymouth Weather Station, Washington County, NC). Much of this region historically was covered by pocosin wetlands (evergreen shrub-scrub), but by 1979 only 30% of the original wetlands remained [Richardson, 1983]. Much of the landscape was deforested at the beginning of the 20th century and drained for agriculture in the 1970s–1980s [Carter, 1975]. Agriculture continues to be the predominant land use, with 80% of the agricultural area considered marginal because it requires active pumping to allow agriculture [Neely, 2008]. A critical feature of this area is the bidirectional hydrology of rivers, streams and wetlands due to wind-driven tides [Poulter et al., 2008]. Farmers constructed one-direction pump stations at the end of their properties to actively pump and maintain low-water tables needed for agriculture. These pump stations also prevented the upstream movement of water due to wind tides. Restoring the hydrology to such an area is done by removing pump stations and allowing the bidirectional movement of water.

2.2. Timberlake Restoration Project

[7] Timberlake Restoration Project (TLRP), located in the Albemarle Peninsula in Tyrrell County, NC (35°54′22″ N 76°09′25″W), is part of Great Dismal Swamp Mitigation Bank, LLC. The main objective of the mitigation bank is “to establish self-sustaining, functioning aquatic systems to replace the function and acreage of wetlands and other aquatic resources anticipated to be adversely affected” (U.S. Army Corps of Engineers, Umbrella memorandum of agreement between bank sponsors, p. 27, 1997). The total 1704.2 ha compensatory mitigation site is composed of: 420 ha of mature forested wetland that was never in agriculture (F), 787 ha of forested wetland under a preservation easement (preservation area, PA), 57.2 ha of drained shrub-scrub, and 440 ha of former agricultural fields undergoing stream and wetland restoration (RW). The site drains to the Albemarle Sound via the Little Alligator River (3 km from the site). Elevation in the site ranges from −0.4 m to 5.1 m above sea level [Needham, 2006]. The two major soil series in the site are Ponzer muck (loamy, mixed, dysic, thermic Terric Haplosapristis) and Hyde loam (fine-silty, mixed active, thermic Typic Umbraquults, USDA SSURGO Database 2005). There are also areas of Roper muck, Weeksville silt loam, and Pungo muck [Needham, 2006].

[8] The focus of this study was the 440 ha former agricultural field (RW), which is being restored into stream, riverine and nonriverine wetlands (Figure 1). After the last corn harvest in August 2004 land movement was begun to restore the hydrology to its preagricultural state. This included filling 53 km (33 miles) of “vee” ditches (approximately 90–100 cm deep), plunging sections of the main canal (approximately 3 m deep), and creating a focused zone of preferential flow by connecting the lowest-elevation areas across the site (Figure 1). Restoration also included planting 750,000 trees from 10 native wetland species: Taxodium distichum, Nyssa sylvatica var. biflora, Nyssa aquatica, Fraxinus pennsylvanica, Salix nigra, Chamaecyparis thyoides, Quercus nigra, Quercus michauxii, Quercus phellos, and Quercus falcata var. pagodafolia
Disabling the downstream gate-pump system in February 2007 was the final step in the restoration process, reinstating the precipitation and wind tide hydrologic regime and associated groundwater fluctuations. In August 2007 a pump was installed in the upstream end of RW to allow the 2424 ha adjacent farm to pump excess water onto the site.

In addition to the work on RW, we studied two minimally impacted wetlands: the preservation forested wetland (PA) (which is dominated by Belhaven muck (loamy, mixed, dysic, thermic Terric Haplosapristis)) and an area of freshwater wetlands in the nearby (5 miles north of TLRP) Palmetto Peartree Preserve (PP, dominant soil type Dorovan muck, dysic, thermic Typic Haplosapristis). To contrast the reference wetlands, we also worked in an active agricultural field (Ag, dominant soil type Weeksville silt loam, coarse-silty, mixed semiaquve, thermic Typic Umbraquults) immediately south of TLRP (Figure 1).

3. Methods

3.1. Phosphorus in Surface and Rainwater

To examine P losses from the restored site (RW), a forested wetland (For) and an agricultural field (Ag) we collected water samples (one filtered with 0.7 μm GF/F Whatman filter, and one unfiltered) from a constrained outflow in each one (Figure 1) weekly for 2 years (February 2007 to February 2009). Samples were refrigerated and maintained at 4°C until analyzed (usually within 3 weeks). On the filtered samples we measured soluble reactive phosphorus (SRP) using the ascorbic acid and molybdenum blue method [American Public Health Association (APHA), 1998] on a Lachat QuickChem automated system (Lachat QuikChem 8000, Lachat Instruments, Milwaukee WI). On the unfiltered samples we measured total phosphorus (TP) using persulfate digestion followed by ascorbic acid and molybdenum blue method as above [APHA, 1998]. Water pH, dissolved oxygen, conductivity, and temperature were measured every time we collected samples using a handheld device (YSI Multiprobe 560, Yellow Springs, OH, United States).

3.2. Hydrologic Manipulations and Storm Sampling

Because of the rapid changes expected during the initial flooding, drawdown, and storm events, we also conducted high frequency sampling for specific windows of time. During the initial flooding of RW (28 February to 3 March 2007) we collected samples every 6 h using ISCO automated samplers (ISCO 6712, Teledyne ISCO, Ohio). We conducted two drawdown experiments in RW, one in winter (4–18 February 2008) and one during the summer (18 August to 2 September 2008). During the drawdown experiments we activated the pump (70,000 gallons min⁻¹) on the downstream end of RW to decrease water depth by 1 m to approximately 10 ha for 2 weeks. During experimental drawdown we collected samples every 4–12 h depending on how fast surface water elevation was changing. We also conducted high frequency (samples every 4–6 h) sampling for 10 storm events over the 2 years for the RW and eight storms for the For. We compared concentrations and fluxes of SRP and TP from an active agricultural field (Ag), a mature forested wetland (For), and the former agricultural field undergoing restoration (RW).

3.3. Soil Solution P

To examine potential P release from the sediment to surface water, we measured soil solution every 2–4 months on 33 sampling points across the elevation gradient at the site. We sampled pore water from a network of piezometers (wetter conditions) and lysimeters (drier conditions) at 15 cm depth. In seven sites we also installed piezometers at 30, 60 and 100 cm. Samples were collected first by purging the piezometers, or creating a vacuum (40 kPa) on the lysimeters, and then collecting water samples the next
day. Samples were collected in HDPE plastic bottles, one unfiltered and one filtered (Whatman GF/F, 0.7 μm) and refrigerated until analyzed as described above. If we did not have enough sample for both SRP and TP, we prioritized SRP. We also collected soil solution from 5 piezometers (15 cm deep) before, during, and after the summer drawdown experiment (18 August to 2 September 2008).

[13] Each soil solution sampling point was instrumented with 5 platinum electrode redox probes [Vepraskas et al., 2004] along with a water level recorder (Solinst Levelogger Gold, Canada) at 15 cm depth. To examine the effects of hydrology on soil solution within RW we classified the sampling sites according to the average water table depth (WTD), such that there were 3 groups: RW-Dry (mean WTD < –20 cm), RW-Int (mean WTD between –15 and 0.5 cm), and RW-Wet (mean WTD > 5 cm). In addition to the points within RW, we instrumented 5 sampling points in preservation wetlands (PA), Palmetto Peartree Preserve (PP) and the agricultural field (Ag).

3.4. Soil Analyses

[14] To examine changes in soil nutrient pools, we collected soil samples in 2006 (August and November), 2007 (June and October) and 2008 (October). Soil cores (5 cm internal diameter, 30 cm long) were collected from 12 points from the three hydrology groups within RW described above: RW-Dry (n = 4), RW-Int (n = 4), and RW-Wet (n = 4). In addition to sampling within RW, on June 2007 and October 2008 we collected soil samples from 5 sampling locations in two nearby forested wetlands (PA and PP, n = 5 each) and an active agricultural field (Ag, n = 5). Soil samples were stored on ice during the trip to the laboratory where they were stored at 4°C until processed. Each sample was sectioned into the top 15 cm and 15–30 cm, and then sieved by hand (2 mm sieve) to remove large plant and root material. Here, we report data from the top 15 cm, where most of the biological activity occurs. Soil carbon and nitrogen were measured on a Carlo Erba Elantech Model NC2100 elemental analyzer (ThermoQuest Italia, Milan, Italy). Total phosphorus was determined after nitric-perchloric digest of 0.6 g of dried soils [Carter, 1993] and analyzed on Lachat QuickChem as described above.

[15] We conducted single point P isotherms on the 2007 soil samples to determine the capacity of the soils to sorb P. We determined a single point P isotherm according to [Axt and Walbridge, 1999]. Sediments (2.0 g of dry weight equivalent) were shaken for 24 h in 25 ml of 0.01 M CaCl2 containing 130 mg P/L as KH2PO4. The samples were then centrifuged for 20 min at 3400 rpm, filtered (Whatman GF/F, 0.7 μm) and analyzed for SRP as described above. We estimated the phosphorus sorption index (PSI) as x/log c, where x = P adsorbed by the soil (mg P per 100 g of soil) and c = the equilibrium solution SRP concentration after 24 h of shaking (in μmol/L).

[16] Because Fe and Al have been shown to be important controls on P sorption we also measured oxalate-extractable Al (Alox) and Fe (Feo) on the 2007 samples. We extracted 0.4 g of soil (dry weight equivalent) using 0.2 M ammonium oxalate and 0.2 M oxalic acid, by shaking for 4h in darkness (soil solution ratio 1:100) [Darke and Walbridge, 1994]. Two drops of Superfloc 0.4% were added before being centrifuged for 5 min at 2000 rpm. Samples were filtered (Whatman no. 42) and stored at 4°C until analyzed using atomic absorption spectroscopy (AAS 5100PC; Perkin Elmer, Norwalk, CT, United States).

3.5. Fates of Legacy P

[17] To examine changes in P cycling over time in RW we used a space-for-time substitution to construct our best educated guess for P budgets under natural wetland conditions, under active agriculture, and following the hydrologic transition due to wetland restoration. At each stage we combined our own data with literature values to examine the major pools (soil and plant biomass) and fluxes (surface water export) of P. To estimate soil and plant biomass P pools of a natural wetland we used soil TP data from our two minimally impacted wetland sites (PA and PP) and literature values for plant biomass TP from Mitsch et al. [1979]. To estimate surface water export of P during the natural wetland condition we used our measured TP export from the For site, which drains a mature forested wetland. We combined estimates of P in loose sediments (floc), periphyton, and macrophytes from Noe and Childers [2007] to estimate the size of the actively cycled P pool. To approximate changes in conversion to agriculture we estimated gains in soil P due to fertilizer application by multiplying the annual fertilizer application rates used in the area [Crozier, 2000; K. Cherry, Cherry Farms, Ttryrell County, North Carolina, personal communication, 2009] by the 20 years the site was under agriculture, while accounting for losses due to crop removal (K. Cherry, personal communication, 2009). We estimated surface water P losses during agriculture from measured P concentrations and fluxes from Ag [Ardón et al., 2010]. To estimate plant biomass P due to the 750,000 trees planted as part of the restoration, we used measurements of tree diameter collected for mitigation monitoring reasons (R. Needham, unpublished data, 2009). We used annual measurements of tree diameter at breast height from 38 to 73 plots (0.04 ha) across RW taken in 2006, 2007 and 2008. We used annual growth that occurred between each of those years and allometric equations from Jenkins et al. [2003] and Schroeder et al. [1997] to estimate plant biomass C. We increased the total plant biomass carbon estimated through the allometric equations by 25% to estimate belowground biomass [Schlesinger, 1997]. We then converted estimates of plant biomass C to biomass P using a C:P molar ratio of 1378 based on published data of % P for some of the most common trees on the site (Taxodium distichum, Nyssa sylvatica, Nyssa aquatica and Quercus sp. [Bedford et al., 1999]). We used a weighted average of soil TP measurements derived from 2006 (preflooding), 2007, and 2008 from the three hydrologic groups according to their relative area (RW-Dry 41%, RW-Int 41%, and RW-Wet 18%). To get a “back of the envelope” estimate of the time it will take RW to return to preagricultural state we estimated the amount of legacy fertilizer TP that could be lost as the difference between the 2008 soil TP in RW and soil TP in our two reference wetlands. We then used our measurements of TP export from RW and estimates of plant biomass P uptake as the two means of losses of P from the soil into long-term storage or solute export. This calculation relies on three simplifying assumptions: (1) surface export will be constant over time; (2) plant P uptake will also be constant; and
3.6. Statistical Analysis

To compare the concentration and export of SRP and TP among sites and seasons, we used two-way analysis of variance (ANOVA) on log transformed data with site (For, RW and Ag) and season (spring, summer, fall, and winter) as factors, followed by post hoc Tukey HSD. Because the pump that drains the Ag was only activated 28 days over the 2 years, we did not include fluxes from the Ag field in statistical analyses (but we did include concentrations from Ag site in statistical analyses). We used linear regression to determine relationships between changes in water surface elevation and SRP or TP export. We used analysis of covariance (ANCOVA) to compare soil TP between the three hydrologic groups (RW-Dry, RW-Int and RW-Wet) and 3 sampling years (2006, 2007 and 2008). We used linear regression to examine relationships between Al and Fe and P sorption index. All analyses were conducted using JMP Statistical software (SAS, Cary, North Carolina).

4. Results

4.1. Hydrologic Losses and Drawdown Experiments

Over the 2 years we sampled, we found clear seasonal patterns in both TP and SRP concentrations (mg/L) and export (g ha\(^{-1}\) d\(^{-1}\)) leaving RW, For, and Ag (Figure 2). TP and SRP concentrations were higher during spring and summer months in both sites (TP ANOVA F = 8.03, df = 299, p < 0.001; SRP ANOVA F = 4.49, df = 299, p < 0.001, Figure 2). SRP concentrations were 2.5 times higher leaving RW (0.017 ± 0.0025 mg/L) than the For (0.007 ± 0.0025 mg/L) and Ag (0.006 ± 0.0025 mg/L) during spring (Tukey HSD p < 0.0001, Figure 2a). TP concentrations were 2.5 times higher leaving RW (0.017 ± 0.0025 mg/L) than the For (0.007 ± 0.0025 mg/L) and Ag (0.006 ± 0.0025 mg/L) during spring (Tukey HSD p < 0.0001, Figure 2a).

Figure 2. Mean (±standard error) concentrations (mg/L) and daily fluxes (g ha\(^{-1}\) d\(^{-1}\)) of (a and b) SRP and (d and e) TP for an active agricultural field (Ag), a forested wetland (For) and restored wetland (RW). (c) Relationships between maximum increase in water surface elevation due to rain events and SRP (solid circles) and TP (open circles) export (kg/storm event) over 2 years from RW. Regressions in Figure 2c: TP \(r^2 = 0.81, p < 0.001\), SRP \(r^2 = 0.62, p < 0.001\). Asterisks denote significant differences from post hoc Tukey HSD.

(3) changes in bulk density of the soils due to peat and sediment accumulation will not affect P release or retention.
There was a significant positive relationship between increases in water surface elevation and SRP and TP export from RW during storms (Figure 2c).

During the initial flooding in February 2007, water depth in RW increased by more than 1 m in a day, leading to increased SRP concentration (Figure 3a). The experimental drawdown in winter rapidly decreased water depth to prerestoration levels, but did not cause any changes in SRP concentrations (Figure 3b). Average water temperature during the low-water period was 11.5°C (range 2 to 21°C). The experimental drawdown in summer also decreased water depth to prerestoration levels, and caused a fourfold increase in SRP concentrations during the dry period (Figure 3c), followed by decreases during the reflooding. Average water temperature during the low-water period was 26.6°C (range 20.2°–36.1°C).

4.2. Soil Solution

We collected and analyzed 190 samples of soil solution over the 2 years from all sites. Soil solution SRP and TP varied across sites and hydrologic group within RW (Table 1 and Figure 4). SRP was higher in the RW–Wet sites than in the other sites, but the difference was not significant (Figure 4a). TP was significantly higher in RW–Wet than the RW-Dry sites (Figure 4b, df = 189, F = 7.24, p < 0.001). The SRP:TP ratio in the RW-Dry, RW-Int and Wet sites was similar to the ratio observed in surface water (0.1–0.17, Table 1). Due to drought and active pumping of the water table, we were unable to obtain soil solution from the active agricultural field. Soil solution SRP increased during the summer drawdown event (Figure 5). While all wells showed increases, the magnitude of increase was least in the highest-elevation well (Figure 5c).

### Table 1. Mean and Range (Minimum–Maximum, mg/L) Phosphorus Concentrations in Soil Solution in the Timberlake Restoration Project and Three Reference Sites

<table>
<thead>
<tr>
<th>Soil Solution</th>
<th>TP</th>
<th>SRP</th>
<th>SRP:TP</th>
</tr>
</thead>
<tbody>
<tr>
<td>RW-Dry</td>
<td>0.063</td>
<td>0.011</td>
<td>0.17</td>
</tr>
<tr>
<td>RW-Int</td>
<td>0.025–0.44</td>
<td>0.0025–0.089</td>
<td>0.16</td>
</tr>
<tr>
<td>RW-Wet</td>
<td>0.103</td>
<td>0.016</td>
<td>0.14</td>
</tr>
<tr>
<td>PA</td>
<td>0.153</td>
<td>0.021</td>
<td>0.14</td>
</tr>
<tr>
<td>PP</td>
<td>0.01–0.25</td>
<td>0.0025–0.121</td>
<td>0.16</td>
</tr>
<tr>
<td>Ag</td>
<td>0.016–0.48</td>
<td>0.0025–0.016</td>
<td>0.03</td>
</tr>
</tbody>
</table>

*RW-Dry, restored wetland dry; RW-Int, restored wetland intermittent; RW-Wet, restored wetland wet; Ag, agricultural field; PA, Timberlake preservation area; PP, Palmetto Peartree Preserve wetlands. See text for definitions of hydrologic groups within RW.*
4.3. Soil Pools and Assays

RW soils had significantly higher bulk density than the reference wetlands and lower than the active farm field (F = 48.1, df = 65, p < 0.001, Table 2). The concentrations (in both mol/kg and mol/m$^3$) in RW-Dry were similar to the Ag for C, N, TP, Al$^{\text{ox}}$ and Fe$^{\text{ox}}$ (Table 2). The soil characteristics of the RW-Wet tended to be more similar to the two reference wetlands than the RW-Dry or RW-Int (Table 2). The agricultural field had significantly higher TP in the soil than the other sites (F = 26.7, df = 65, p < 0.001, Table 2). Within soils in RW there was a significant difference in soil TP among the three hydrologic groups (F = 11.2, df = 35, p < 0.001, Table 2). Within soils in RW there was a significant decrease in soil TP with time (F = 4.2, df = 35, p < 0.05, Figure 6), but the interaction term was not significant (hydrology by time, p = 0.9). P sorption index (PSI) was significantly higher in RW-Wet, RW-Int and PA, and similar between the Ag and one of the reference sites (PP, Table 2). PSI was strongly correlated to Al$^{\text{ox}}$ but not to Fe$^{\text{ox}}$ for all sites combined (Figure 7).

4.4. Fates of Legacy P

Our best estimate of the P budget of a natural wetland in this area before agricultural conversion suggests that 2.9 kmol P ha$^{-1}$ are stored in plant biomass, 3.6 kmol P ha$^{-1}$ are stored in soil, and 0.003 kmol P ha$^{-1}$ yr$^{-1}$ are exported through surface water. Based on Noe and Childers [2007] we estimate there is approximately 0.28 kmol P ha$^{-1}$ in floc, periphyton and macrophytes. Due to the naturally low P in

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Figure 4. Soil solution (average ± standard deviation) (a) SRP and (b) TP in two reference wetlands (PA, preservation area; PP, Palmetto Peartree Preserve) and three hydrologic groups within Timberlake Restoration Project over 2 years. RW-Dry, restored wetland dry sites; RW-Int, intermittently flooded sites in restored wetland; RW-Wet, permanently flooded sites in the restored wetland. See text for details of determination of these hydrologic groups. Letters denote significant differences after post hoc Tukey HSD.

Figure 5. Soil solution SRP (mg/L) in three sampling wells during the summer drawdown experiment (18 August to 2 September 2008) in Timberlake Restoration Project. Wells spanned a hydrologic gradient: (a) permanently flooded, (b) intermittently flooded, and (c) permanently dry. The site in Figure 5c had the highest elevation, and thus was least affected by the drawdown experiment. Arrows indicate start and end of the pumping.
Table 2. Soil Characteristics (Average ± Standard Deviation) From the Three Hydrologic Groups Within Timberlake Restoration Project (RW-Dry, RW-Int, and RW-Wet) and Three Reference Sites

<table>
<thead>
<tr>
<th>Location</th>
<th>Bulk Density (kg/m$^3$)</th>
<th>Carbon Mass (mol/kg)</th>
<th>Carbon Volume (mol/m$^3$)</th>
<th>Nitrogen Mass (mol/kg)</th>
<th>Nitrogen Volume (mol/m$^3$)</th>
<th>Phosphorus Mass (mol/kg)</th>
<th>Phosphorus Volume (mol/m$^3$)</th>
<th>Al$^{3+}$ Mass (mol/kg)</th>
<th>Al$^{3+}$ Volume (mol/m$^3$)</th>
<th>Fe$^{3+}$ Mass (mol/kg)</th>
<th>Fe$^{3+}$ Volume (mol/m$^3$)</th>
<th>PSI</th>
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<tbody>
<tr>
<td>Ag</td>
<td>1306 ± 55 (a)</td>
<td>2.6 ± 2.0 (a)</td>
<td>3425 ± 924 (ab)</td>
<td>115 ± 97 (a)</td>
<td>150 ± 26 (bc)</td>
<td>14.3 ± 1.4 (b)</td>
<td>18.7 ± 0.7 (a)</td>
<td>86.1 ± 13.7 (a)</td>
<td>112 ± 7.6 (a)</td>
<td>27.2 ± 5.4 (a)</td>
<td>35.4 ± 2.8 (a)</td>
<td></td>
</tr>
<tr>
<td>RW-Dry</td>
<td>1062 ± 85 (b)</td>
<td>6.5 ± 1.4 (a)</td>
<td>5365 ± 501 (b)</td>
<td>229 ± 50 (a)</td>
<td>187 ± 13 (bc)</td>
<td>7.4 ± 1.0 (c)</td>
<td>6.5 ± 0.5 (b)</td>
<td>108.3 ± 7.9 (a)</td>
<td>103 ± 5.1 (a)</td>
<td>16.3 ± 3.1 (a)</td>
<td>15.3 ± 1.9 (b)</td>
<td></td>
</tr>
<tr>
<td>RW-Int</td>
<td>854 ± 173 (c)</td>
<td>10.9 ± 1.7 (a)</td>
<td>6255 ± 653 (bc)</td>
<td>349 ± 60 (a)</td>
<td>235 ± 16 (ab)</td>
<td>7.1 ± 1.2 (c)</td>
<td>5.0 ± 0.6 (bc)</td>
<td>126.8 ± 9.7 (a)</td>
<td>105 ± 5.1 (a)</td>
<td>26.4 ± 3.8 (a)</td>
<td>23.4 ± 1.9 (b)</td>
<td></td>
</tr>
<tr>
<td>RW-Wet</td>
<td>441 ± 97 (d)</td>
<td>23.0 ± 1.6 (b)</td>
<td>8685 ± 623 (c)</td>
<td>673 ± 56 (b)</td>
<td>264 ± 15 (a)</td>
<td>11.1 ± 0.9 (bc)</td>
<td>4.0 ± 0.5 (c)</td>
<td>176.9 ± 7.4 (b)</td>
<td>75.6 ± 4.7 (b)</td>
<td>47.7 ± 2.9 (b)</td>
<td>19.1 ± 1.6 (b)</td>
<td></td>
</tr>
<tr>
<td>PA</td>
<td>184 ± 98 (e)</td>
<td>24.9 ± 2.8 (b)</td>
<td>4139 ± 921 (ab)</td>
<td>1488 ± 97 (c)</td>
<td>248 ± 26 (ab)</td>
<td>28.0 ± 1.6 (a)</td>
<td>3.4 ± 0.7 (d)</td>
<td>251.8 ± 13 (c)</td>
<td>46.7 ± 7.4 (c)</td>
<td>27.1 ± 5.4 (a)</td>
<td>4.6 ± 2.8 (c)</td>
<td></td>
</tr>
<tr>
<td>PP</td>
<td>73 ± 17 (f)</td>
<td>29.4 ± 2.4 (b)</td>
<td>2112 ± 922 (a)</td>
<td>1584 ± 97 (c)</td>
<td>113 ± 28 (c)</td>
<td>24.5 ± 1.7 (a)</td>
<td>1.5 ± 0.8 (d)</td>
<td>122.2 ± 7.4 (a)</td>
<td>8.8 ± 7.2 (d)</td>
<td>51.7 ± 5.1 (b)</td>
<td>3.5 ± 2.7 (c)</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Ag, agricultural field; PA, Timberlake preservation area; PP, Palmetto Peartree Preserve wetlands. Concentrations of elements are shown both on a mass basis (mol/kg) and a volume basis (mol/m$^3$) due to differences in bulk density across sites. Different letters denote significant differences ($p < 0.05$) from post hoc Tukey HSD.

Figure 6. Soil TP (mean ± standard error) in three hydrologic groups within Timberlake Restoration Project. RW-Dry, restored wetland dry sites; RW-Int, intermittently flooded sites in restored wetland; RW-Wet, permanently flooded sites in the restored wetland. See text for details of hydrologic groups. Three linear regressions for each hydrologic group. ANOVA Tukey HSD $p < 0.05$.

Figure 7. Relationships between single point phosphorus sorption index (PSI) and (A) Al$^{3+}$ and (B) Fe$^{3+}$. $r^2 = 0.49$, $p < 0.0001$. PA, preservation area; PP, Palmetto Peartree Preserve; RW-Dry, restored wetland dry sites; RW-Int, intermittently flooded sites in restored wetland; RW-Wet, permanently flooded sites in the restored wetland.
situations of SRP and TP among soil, soil solution, and the overlying water [Reddy and DeLaune, 2008]. Similarly, it is unlikely that vegetation uptake will remain constant, it will likely increase exponentially and level off [Craft et al., 25]

5. Discussion

[24] We found consistent results that both natural and experimental hydrologic fluctuations can lead to increased P concentrations and export from a restored wetland ecosystem. Over the 2 years of this study, SRP concentrations of water leaving RW were 2.5 times higher than in a forested wetland (For) and an agricultural field (Ag) during the spring (Figure 2a). Higher concentrations, along with increased hydrologic flux, led to higher export of SRP and TP from the restored wetland (RW) than forested wetland (For) in spring and summer (Figures 2b and 2e). We found various lines of evidence that indicate that flooding can lead to release and export of various forms of P from RW. First the initial flooding as part of restoration increased SRP concentrations (Figure 3a), potentially due to the release of Fe-bound P due to anoxic conditions. Increases in water elevation due to precipitation events increased TP export more than SRP (Figure 2c), which suggests that storms mobilized particulate and organic forms of P. We found higher soil solution TP and lower soil TP in flooded than in dry soils (Figures 4 and 6), suggesting that flooded conditions facilitate soil P losses. We also found that a drawdown experiment during the summer increased SRP concentrations in both surface (Figure 3) and soil solution (Figure 5), suggesting that mineralization of organic P during oxic conditions can also be an important source of SRP.

[25] Our “back-of-the-envelope” calculation suggests that the Timberlake Restoration Project could continue to release legacy P for 3–16 years. We acknowledge that there is a lot of uncertainty around those estimates, and that our three key assumptions (constant plant uptake, constant hydrologic losses, and negligible changes in soil bulk density) are not likely to hold true over time. For example, it is unlikely that hydrologic export will be constant. During a year with high rainfall, we would expect that more SRP and TP would be exported in surface water. The export from TLRP will depend on hydrologic flux and the equilibrium concentrations of SRP and TP among soil, soil solution, and the overlying water [Reddy and DeLaune, 2008]. Similarly, it is unlikely that vegetation uptake will remain constant, it will likely increase exponentially and level off [Craft et al.,
Some of vegetation P uptake will be incorporated into woody biomass, while some will be recycled through litter decomposition [Noe and Childers, 2007]. Vegetation will also play an important indirect role in determining P export by decreasing water yield through increased evapotranspiration as the trees mature and add more leaf surface area, and by increasing sedimentation and peat accumulation. We also acknowledge that the differences in soil TP over time in RW should be interpreted with caution given the relatively low level of soil sampling for such a large site.

Considering these caveats, our estimates do provide a realistic benchmark to compare to previous studies that have documented temporary P release after reflooding former agricultural soils in the laboratory [Aldous et al., 2005, 2007; Pant and Reddy, 2003]. Previous reports of the duration of P release after flooding have ranged from months [Aldous et al., 2005, 2007] to 3 years [Pant and Reddy, 2003; Reddy and DeLaune, 2008]. Our estimates, based on field measurements, significantly extend that time frame and agree with other field studies in the literature. For example, a recent study found large P fluxes from a restored lake fringe wetland in Oregon 5–7 years after restoration [Duff et al., 2009]. The exact time period that TLRP will release legacy P is unknown, but our results illustrate the importance of hydrologic fluctuations and seasonal patterns in affecting both the timing and magnitude of P losses.

Due to the large size of RW (440 ha) we expect there will be both spatial and temporal variations in P release and retention from patches within the site. Higher concentrations of soil solution SRP and TP in the RW-Wet soils (Figure 4) suggest these areas could be sources of P. However, the soils across the site have the capacity to sorb P (Figure 7), primarily controlled by $A_{\text{ox}}$, as has been observed in other wetlands [Darke and Walbridge, 2000; Richardson, 1985]. We found higher concentrations of $A_{\text{ox}}$ and higher sorption capacity (PSI) in the RW-Wet than the RW-Dry soils (Table 2), suggesting the RW-Wet soils could also function as P sinks. Seasonal differences due to both biological activity and extent of flooding could change soils across RW from sources to sinks of P. The interplay between hydrologic fluctuations, leading to P release, and internal buffering by soils through sorption to noncrystalline forms of Al, will determine the period of time before P cycling in RW returns to its preagricultural condition.

### 5.1. Solute P Losses

Increased P export from restored wetlands may have negative implications for downstream ecosystems. The Albemarle Sound is 5 km downstream and has historically low P concentrations (SRP usually less than 0.04 mg/L) and N:P ratio of 20, both of which suggest P limitation of phytoplankton [Lin et al., 2007; Richardson, 1983]. The Albemarle-Pamlico sound has previously experienced problems with harmful algal blooms, hypoxia and massive fish kills in response to elevated nutrient delivery [Burkholder et al., 2004; Paerl et al., 2001]. SRP and TP concentrations leaving RW were 2.5 times higher than concentrations draining Ag and For in the spring. The low concentrations of SRP and TP leaving the agricultural field, even under fertilizer application, suggest that these soils tend to be P limited and have a high capacity to sorb P. As we found (Table 2 and Figure 7), our results suggest that the hydrologic reconnection of TLRP could lead to sustained releases of P in the spring of each year as well as in response to drying and reflooding cycles due to droughts and storms. The spring timing of P pulse could provide limiting P to phytoplankton during a critical time when their activity is increasing.

Higher TP concentrations in soil solution at RW-Wet compared to RW-Dry and RW-Int (Table 1 and Figure 4), suggest that P release is associated with reducing conditions in flooded areas. Our results agree with previous reports of P release after reflooding former agricultural fields: soils formerly in dairy production in Florida [Pant and Reddy, 2003], peat soils in the Netherlands [Van Dijk et al., 2004], and soils from a restored lake fringe wetland in Oregon [Aldous et al., 2005, 2007; Duff et al., 2009]. Pant and Reddy [2003] reported rapid release of P from solubilization of Fe-bound P after flooding soils that had been in dairy production, and the magnitude of flux decreased after subsequent flooding and drying cycles. In peat soils in the Netherlands, increases in soil and soil solution pH lead to increased P release due to stimulation of mineralization [Van Dijk et al., 2004]. Our results provide support for both of these mechanisms, as we observed high P concentrations under both reducing conditions and in response to an experimental drought during the summer.

### 5.2. Hydrologic Fluctuations

To our knowledge, our study is the first to use an ecosystem-scale hydrologic manipulation to examine P dynamics. Most studies on the effects of hydrologic fluctuations on wetland soil P have been either in the laboratory [Aldous et al., 2005, 2007; Bostic and White, 2007; Corstanje and Reddy, 2004; Pant and Reddy, 2003], or at the lysimeter/well scale in the field [Van Dijk et al., 2004; Duff et al., 2009]. In the first drawdown experiment in February we did not see any changes in SRP concentrations during the drought or reflooding period. The lack of SRP increases after reflooding in February 2008 suggests that there was not a large pool of Fe-bound P that could have been released after reflooding. The relatively cold temperature (median 11.6°C, range 2°–21°C) during this period meant there was low biological activity, thus mineralization was likely to be low during the drought period. During the summer drawdown experiment, we observed a fourfold increase in SRP concentrations during the low–water period (Figure 3c). The water temperature in the summer experiment was much higher (median 25.6°C, range 20.2°–36.1°C), suggesting that with the onset of oxic conditions, mineralization of organic P was stimulated. The increase in SRP concentration could have been due to decreasing the amount of water on the site. However, the increase in SRP concentration following the rainfall events after 28 August provide support for our hypothesis that mineralization in oxic sites in the upper layers of the soil led to the increased P. Increases in soil solution P during the dry period (Figure 5) also support our interpretation of increased mineralization during the dry period. Our drawdown experiments agreed with our weekly sampling in illustrating the importance of seasonal patterns in
P transformations, something we would have missed in laboratory experiments under optimal conditions.

5.3. Soil Changes

[31] We were initially surprised by the difference in soil TP (mol/m$^3$, Table 2) between RW and the Ag site, which was also reflected in the difference between the soil TP after 20 years of agriculture (Figure 8a) and the initial stage of wetland restoration (Figure 8b). We had expected these soils to be more similar given that TLRP was in agriculture for 20 years. We believe the large difference in soil TP between the Ag site and our initial sampling of RW (Table 2 and Figure 8) was due to earth moving that occurred prior to our sampling. We believe the filling of 53 km of vee ditches could have facilitated hydrologic and wind losses of TP and the redistribution of legacy fertilizer P to deeper horizons that were not covered by our soil sampling of the top 15 cm. Previous studies have suggested removing topsoil that has received fertilizer before restoration to prevent P release after reflooding [Bostie and White, 2007; Van Dijk et al., 2004]. It is possible that the earth moving at TLRP had a similar effect as removing the topsoil by decreasing soil TP available for mobilization. However, it is unclear if the earth movement led to the movement of P to deeper and more stable soil horizons [Reddy and DeLaune, 2008] or a P pulse to downstream ecosystems.

[32] We found consistent declines in soil TP in all three of the hydrologic groups within RW (Figure 6). Contrary to our expectations, all three hydrologic groups lost TP at a similar rate (Figure 6). Future research will continue to monitor TP losses from the three hydrologic groups to try to determine the fate of the legacy P. We were unable to account for the observed soil TP decline through surface water export and uptake by trees, suggesting that it might be entering an actively cycled pool. This actively cycled P can be in soil solution, loose sediments (floc), taken up by algae and nonwoody vegetation. Much of this rapidly cycled P pool is susceptible to hydrologic loss, potentially contributing to particulate P release during storms [Novak et al., 2007].

[33] Properties of the soils within RW were usually intermediate between the ecological end points we sampled (Table 2). RW-Dry soil tended to be more similar to the agricultural field, while RW-Wet was more similar to the reference wetlands. Various studies have reported differences in soil characteristics between natural and restored wetlands [Bruland and Richardson, 2005; Hogan et al., 2004; Meyer et al., 2008; Sundareshwar et al., 2009]. A recent study and literature review suggested it can take decades or centuries before restored wetlands recover the levels of soil properties, such as bulk density and soil organic matter, of natural wetlands [Ballantine and Schneider, 2009]. Bulk density in RW, particularly RW-Dry and RW-Int, reflected the ~20 years the site was under agriculture, as they had higher bulk density than found for either reference wetland sites (Table 2). Soil C concentrations per kg of soil were much higher in the reference wetlands, but when expressed per volume (m$^3$) the highest C was in the RW-Wet soil (Table 2). The lowest-elevation portions of the former agricultural field are most similar to reference wetlands, likely because the areas were poorly drained during agriculture (and thus lost less soil organic matter to oxidation during agriculture).

6. Conclusion

[34] Our results demonstrate the potential for heightened and prolonged P export resulting from converting a former agricultural field to a hydrologically connected riverine wetland. Our best estimates of P cycling suggest that the Timberlake Restoration Project could continue to release legacy fertilizer P for up to 16 years. The time lag between restoration and return to preagriculture conditions should be considered in planning future mitigation needs and developing management strategies of restored wetlands; monitoring programs based on improved water quality conditions, for instance, should incorporate such lag effects into their protocol for sampling timing. The exact time frame for RW to return to its preagricultural state is unknown, but our results indicate that hydrologic fluctuations will play an important role in determining the loss of legacy P. We found clear seasonal and storm effects on P export, suggesting that both the timing of restoration and active management of water levels might help alleviate these problems in future wetland restoration projects.

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