Impoundments Increase Potential for Phosphorus Retention and Remobilization in an Urban Stream

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Abstract

Weirs are conventional structures that control water level and velocity in streams to facilitate water resource management. Despite many weirs built in streams, there is little information how weirs change hydrology regime and how that translates to sediment and phosphorus (P) responses. This study evaluated the influence of weirs on P retention and mobilization in an urban tributary of the Han River in Korea. Total P concentrations in sediments upstream of weirs were higher than the downstream site, mainly due to the increase of potentially available fractions (labile P and aluminum- and iron-bound P) ($p < 0.05$). Equilibrium phosphorus concentrations ($\text{EPC}_\text{p}$) were lower than soluble reactive phosphorus (SRP) concentrations of stream waters, but there was an increasing trend of sediment $\text{EPC}_\text{p}$ upstream of weirs compared to the downstream site ($p < 0.001$) indicating a greater potential for P release upstream of weirs. Sediment core incubation showed that SRP release rates upstream of weirs were higher than the downstream site under anoxic conditions of the water column ($p < 0.01$), but not under oxic conditions. SRP release rates under anoxic conditions were greater than that measured under oxic conditions. Un-neutral pH and increased temperature could also enhance SRP release rates upstream of weirs. We conclude that weirs can increase P retention within stream sediments and potentially promote significant P releases into waters, which in turn cause eutrophication.

Keywords: Eutrophication, Phosphorus release, Urban stream, Weir

1. Introduction

Urbanization has led to an increase in phosphorus (P) loading into streams [1]. Typically, high percentage of the P loads is trapped and processed upstream of impounding structures in streams [2, 3]. The impoundment of streams, even with low-head structures, such as weirs, alters hydrology by slowing flow velocities (or increasing hydraulic retention time and hydroperiod), thereby affects sediment dynamics and P cycling [2, 3]. Such excessive P loading under the control of stream flows regulated by impounding structures is of primary concern as this can cause the seasonal eutrophication of streams and socioeconomic problems at regional scales [4-6].

In urban streams, weirs are usually installed for flood control as flooding can be exacerbated by increased amounts of impervious surface [7, 8]. As such, weirs have created stepped stream systems where the hydraulic energy is highly reduced by the reduction of the slope and allow more time for dissolved P to interact with sediments and for particle-bound P to settle out [9-11]. Since the early 1980s, sediment has been identified as the primary pollutant of concern for stream waters. To address this concern, many management practices have been promoted to enhance nutrient management in the United States [12]. Recently, the installation of weirs in agricultural drainage ditches is being advocated as an innovative strategy to improve water quality of downstream ecosystems in the Lower Mississippi Fluvial Valley, USA [13]. The intention of weir installation in agricultural drainage ditches is to reduce sediments and nutrients like P from entering aquatic systems (rivers and streams) by altering ditch hydrology [10]. Although promoting sedimentation and P retention through increased retention time is the premise under which weirs are installed, there are several uncertainties about the way weirs change hydrology regime and how that translates to P and sediment responses in aquatic systems [11, 14, 15]. Information in details regarding influences of weirs on sediment retention and the associated P dynamics is still scarce [13].

Large proportions of P sources into urban streams are in a soluble form as mainly derived from municipal wastewater treatment plants [16]. Highly bioavailable dissolved P may be converted into less-bioavailable particulate and organic fractions as the consequence of uptake onto sediments or by phytoplankton under low flow conditions [2]. Dissolved P retention by sediments, together with particulate P deposition, during low flows permanently contributes to large stores of P in sediments.
upstream of weirs, especially in streams with high point source P loads [11, 13]. A portion of these P pools will be flushed down the stream during subsequent high flows. However, some previous studies have particularly shown the lack of net remobilization of the P pool associated with sediments above the weirs under highest flows in urban streams [11, 14]. Sediments above the weirs of urban streams thus potentially constitute long-term sinks, and consequently long-term sources, of P for years, decades and even centuries [17]. This P pool has been referred to as legacy P, which is particularly problematic to the watershed conservation management [15, 18]. The spatial-temporal variations of hydraulic, hydrology, morphology and land management clearly exert interacting controls on the interaction between sediments and stream water and the lag between P storage and release [15]. Quantification of phosphorus uptake/release mechanisms along with the sedimentation processes above the weirs would be of value to better understand P dynamics and further help to control eutrophication.

Research suggests that weirs can induce not only hydrological changes but also physicochemical changes of stream water including temperature, dissolved oxygen and pH [19]. Changes in stream water chemistry in turn can affect sediment conditions; for example, by controlling the diffusion of dissolved oxygen into sediments, by determining oxygen demands for microbological activities, and the protonation state of iron and aluminum oxides and hydroxides [20-22]. Sediments are especially considered as long-term sources of P in aquatic systems after the external P inputs from the watershed have been reduced [5, 15]. The magnitude of the phosphorus flux across sediment-water interfaces is substantially variable under effects of many environmental factors, for example ranged from 2% to 28% of the retained P in stream sediments in the basin of Lake Okeechobee [23]. Generally, sediment phosphorus fluxes are attributed to changes in redox conditions at the sediment-water interface [24]. The limited P efflux of oxic sediments is due to the strong adsorption of dissolved P to solid iron oxyhydroxides, whereas under anoxic condition, iron oxyhydroxides reductively dissolve and release phosphorus into the water column [25, 26]. Over the last decades, a number of alternative P release controls have been suggested, including the sediment composition (ratio of FeOOH and Al(OH)₃), the decomposition of organic phosphorus and the regulation of bacterial phosphorus as well as pH condition [20, 27, 28]. Sulfate-controlled P release can also explain several apparent deviations from the classical paradigm in sulfate-rich sediments [29]. However, effects of sulfate on sediment P releases in freshwater systems are small compared to that in saline systems with high sulfate concentration [30]. The indirect influences of weirs on the fate of sediment P pools via controlling water chemistry may finally result in a positive feedback toward greater availability of P in streams.

This study tested the hypothesis that weirs significantly increase the amount of P accumulating in sediments of an urban stream associated with point sources and examined the effects of weirs on the potential mobility of P retained. We aim 1) to describe the variations in patterns of different P fractions in streambed sediments across weirs, 2) to describe the dynamics of P in streambed sediments deposited upstream of weirs, and 3) to discuss the environmental factors mediating the direction and magnitude of phosphorus fluxes between sediments and water body upstream of weirs.

2. Materials and Methods

2.1. Study Site and Sampling

The Tancheon Stream in South Korea was surveyed to examine the trends of P release from sediments upstream of weirs. Tancheon Stream is one of the contaminated tributaries in the basin of the Han River, the largest river in South Korea located near the center of the Korean peninsula. Sediment and water sampling was conducted in a 2-km stretch of Tancheon Stream downstream from Tuncheon Bridge (37° 25’ 41” N; 127° 07’ 02” E) (Fig. 1). This section has a high degree of impact from several sewage treatment works and its flow is controlled by two free overfall weirs, spaced approximately 100 m apart. The weirs have a relatively broad crest (1 m × 1 m; width × height) and made up of cement-based materials across the whole width of the channel (48–55 m). Further details of the weir construction are unfortunately unknown. The Tancheon Treatment Plant is a combined sewage treatment plant among four sewage treatment plants in Seoul city, serving a population of 1.720 × 10^3. The Tancheon Treatment Plant has the capacity to treat up to 890 × 10^3 of sewage water per day [31].

Samples were collected in May 2012 at two locations, about 50 m upstream of the first weir (W1) and downstream of the second weir (W2) (Fig. 1). During the sampling time, the mean discharge was about 7.5 m³/s. The water level was extremely low. In the middle of the stream, the water depth was about 1 m upstream of the weirs and decreased to half a meter downstream of

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the weirs. No discharge was observed from the sewage treatment plant during the sampling time. The sediments are loamy sandy upstream and sandy downstream of the weirs at the surface (0–10 cm) and are underlain with gravel.

Triplicate intact sediment cores (10 cm i.d.), consisting of the top 10 cm of streambed sediments, were collected at each sampling location for the P flux measurement. Other triplicate sediment cores (5 cm i.d.) were also collected at each sampling location for biological and chemical analysis. For the kinetics experiments, eighteen intact sediment cores (10 cm i.d.) and stream water (30 L) were sampled at one location upstream of the weirs, chosen randomly and different from the sampling locations to examine the interacting effects of water pH, dissolved oxygen (DO) and temperature to the P release from streambed sediments.

Triplicate water samples (2 L) were also taken at each station for the analysis of total suspended solid (TSS), soluble reactive phosphorus (SRP), and anion concentration (NO$_3^-$ and SO$_4^{2-}$). All samples were transported to the laboratory immediately within 6 hr after being taken. Water and sediment samples for biological and chemical analysis were transported at 4°C using ice boxes.

2.2. Biological and Physicochemical Measurements

On-site measurement of pH, dissolved oxygen, temperature, and redox potential were made using portable meters. Each sediment core (5 cm i.d.) was homogenized before the analysis of concentrations of inorganic phosphorus fractions. The homogenized sediment samples were also subjected to other physicochemical measurements, such as bulk density (D$_b$), pH, water content, organic matter content (SOM), total phosphorus (TP) concentration, cation exchange capacity (CEC), total Ca, Al, Fe, and Mg concentration, extracellular enzyme activities (β-glucosidase, N-acetylglucosaminidase, phosphatase, and arylsulfatase), and equilibrium phosphorus concentration (EPC$_e$).

The TSS (mg/L) was determined by dividing the oven dry weight of the solid retained onto a 0.45-μm filter, after filtering a portion of the river water, by the volume of water [30]. Nitrate (NO$_3^-$) and sulfate (SO$_4^{2-}$) concentrations were analyzed using ion chromatography ( Dionex, Sunnyvale, CA, USA) after filtering through a 0.20-μm filter. SRP concentration of river water was determined using the ascorbic acid method [32].

Dry bulk density (kg/L) was determined by dividing the oven dry weight of the sediment by its bulk volume. Sediment pH was determined in a solution of 1:2.5 sediment/water using a glass electrode (Orion 3-Star Plus Benchtop pH; Thermo Scientific, Waltham, MA, USA). SOM (% of dry sediment) was measured based on the weight loss in 2.5 g of sediment burned at 550°C for 24 hr, following oven drying at 105°C for 24 hr for determination of water content [33]. Sediment extracellular enzyme activities (β-glucosidase, N-acetylglucosaminidase, phosphatase, and arylsulfatase; nmol/g/min) were measured according to Kang et al. [34]. The CEC was determined by colorimetric method according to Aram et al. [35]. The CEC was determined by colorimetric method according to Aran et al. [35]. The 2.5 g of wet sediment was shaken with 40 mL cobaltihexamine chloride (0.05 N) in a 50 mL polypropylene tube for 1 hr on a rotating shaker at 60 rpm, and then centrifuged at 7,000 g for 10 min. The supernatant was filtered through a 0.22-μm filter and the absorbance was measured at 472 nm. The CEC$_{a472}$ (meq/100 g or cmol+/kg) was calculated as:

$$\text{CEC}_{a472} = \frac{(A472_{\text{meq}} - A472_{\text{meq}})}{A472_{\text{meq}}(\text{cmol+/kg})} \times 50 \times \frac{V}{m} \times 100$$  

(1)

where A472$_{\text{meq}}$ and A472$_{\text{meq}}$ are the absorbencies at 472 nm of 0.05 N (50 meq/L) cobaltihexamine chloride solution and of the sample supernatant; V is the volume in L of 0.05 N cobaltihexamine chloride solution added to the sediment sample (0.04 L); and m is the sediment dry mass (2 g).

Metal extraction was conducted using aqua regia solution (3 HCl:HNO$_3$). A portion of sediment was air-dried, passed through a 2-mm sieve, and ground using a grinding machine. The 3 g of dried sediment was boiled with a mixture of 21 mL HCl (12 M) and 7 mL HNO$_3$ (15.8 M) for 2 hr. The aliquot after digestion was filtered using ashless filters (Whatman No. 400; Whatman, Maidstone, UK). Total concentrations of metals (Fe, Al, Mg, and Ca) in aqua regia aliquots were determined by ICP-MS [36].

2.3. P Fractionation and Total P Measurement

The non-calcareous wet sediments collected from Tanchun Stream were subjected to a sequential inorganic P fractionation [37]. The same procedure was applied successfully to fractionate the inorganic P forms in the non-calcareous streambed sediments of Han River in a previous study [38]. To measure total P concentration (g/kg), the aliquot after digestion for metal measurements was diluted with distilled water by a factor of 10. After the dilution, the aliquot was subjected to the measurement of total P concentration using the ascorbic acid method [32].

2.4. Sediment Core Incubation Experiment for the Kinetics of P Release

P flux measurements were based on the increase in the SRP concentration of the overlying water when the cores (10 cm i.d.) were incubated oxically and anoxically for 10 to 15 days in the dark in plant growth chambers (VS-3DMC; Vision Scientific, Daejeon, Korea). The water volumes collected together with sediment cores were discharged and refilled with filtered river water immediately before the incubation. Care was taken to minimize any disturbance to the sediment cores. Each core contained a volume of 0.785 L of river water (ca. 10 cm depth). The temperature and pH of the water in each sediment core was controlled during the incubation to reflect the in situ conditions at the sampling time. In doing so, the water in each sediment core was bubbled with gas mixtures during two successional stages. Firstly, the water was bubbled with air/CO$_2$ mixture for 10 days at a position of 5 cm above the sediment-water interface. Second, the water was exchanged with new water and continuously bubbled with N$_2$/CO$_2$ for a further 10 days. The bubbles provided the mixing necessary to eliminate concentration gradients in the water column without physically disturbing the sediment-water interface. We used HCl (1 M) and NaOH (1 M), in addition to the adjustment of CO$_2$ bubble flows to control pH levels approximating the expected conditions. Water pH and DO were measured simultaneously using portable pH and DO meters inserted through a hole in the top cap of the core, which was stopped during the experiment and only opened when the monitoring was needed.

The initial SRP concentrations of the overlying water were determined after 5 hr of gas bubbling. A 4-mL aliquot of overlying water was removed twice a day for the first three days of the incubation and daily over the remaining incubation period using syringes via tubing inserted into the water column. The SRP concentration of the water removed was immediately determined
2.6. Statistical Analysis

One-way ANOVAs were run to test the impacts of weirs on water and sediment quality. Repeated-measures ANOVA analysis was used to analyze the differences in the SRP release rates of sediments across weirs. Variations between site (upstream or downstream), incubation condition (oxic or anoxic) and the interaction between incubation and site were analyzed for SRP release experiments. The relationships between variables were evaluated using correlation analysis. PASW Statistics ver. 18.0 software (SPSS Inc., Chicago, IL, USA) was used for all analyses.

3. Results

3.1. Water Chemistry

No significant difference was observed between water temperature after being filtered through a 0.45-µm filter [32]. The cumulative release rates of P (mg/day) were determined through a least square linear regression of the mass of P in the overlying water over time. A calculating correction was added to account for the phosphorus removed at each interval measurement [39]. Areal release rates (mg/m²/day) were calculated as the cumulative release rate divided by the surface area of the sediment in the core (m²). The calculation was based on the initial linear portion of two-phase curves of P releases and reported in this study as maximum release rates.

Kinetic experiments measuring the SRP release rates in sediment cores (10 cm i.d.) were also conducted similarly to investigate the interacting effects of temperature (15°C and 25°C), water oxic/anoxic condition (DO > 7.0 mg/L and DO < 0.5 mg/L), and water pH (pH 3.0, 7.0, and 11.0).

2.5. Equilibrium Phosphorus Concentration

EPCₒ was determined from batch equilibrium experiments [40]. A 0.75-g air dried sediments were equilibrated for 24 hr in 25 mL CaCl₂ solutions (0.01 M) containing different amounts of Ca(H₂PO₄)₂ (0, 0.2, 0.5, 1, and 3 mg/L of P). The change in the amount of SRP adsorbed (∆Pₕₑₜ) after the 24 hr-incubation was plotted against the concentration of SRP in solution ([Pₕₑₜ] after the 24 hr-incubation. The Freundlich isotherm was used to fit the data using a least squares method [41]. EPCₒ (mg/L) is the x-intercept of fitting curves indicating the SRP concentration of the suspension at which no net adsorption or desorption takes place (∆Pₕₑₜ = 0). The y-intercept of the fitting curves indicates the native adsorbed phosphorus (NAP). EPCₒ is an empirical reference point on the sorption curve, permitting a direct estimate of the capacity of the soil or sediment to adsorb or release phosphorus if the concentration is changed. The linear adsorption coefficient K_d (the buffer intensity at the EPCₒ) is estimated as the slope of the isotherm curve. K_f is the Freundlich coefficient, and n is the exponential factor (Fig. 2).

**Table 1. Physicochemical characteristics of stream water and sediments (n = 3)**

<table>
<thead>
<tr>
<th></th>
<th>Water</th>
<th></th>
<th>Sediment</th>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Upstream</td>
<td>Downstream</td>
<td>Upstream</td>
<td>Downstream</td>
</tr>
<tr>
<td>pH</td>
<td>7.93 ± 0.50</td>
<td>9.19 ± 0.02</td>
<td>7.11 ± 0.03</td>
<td>6.99 ± 0.12</td>
</tr>
<tr>
<td>Temp. (°C)</td>
<td>25.7 ± 0.4</td>
<td>26.2 ± 0.5</td>
<td>23.3 ± 0.9</td>
<td>25.2 ± 0.9</td>
</tr>
<tr>
<td>DO (mg/L)</td>
<td>11.4 ± 0.2</td>
<td>8.9 ± 0.3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>TSS (mg/L)</td>
<td>24.0 ± 5.0</td>
<td>9.3 ± 2.6</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NO₂ (mg/L)</td>
<td>0.518 ± 0.013</td>
<td>0.498 ± 0.017</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NO₃ (mg/L)</td>
<td>13.388 ± 0.282</td>
<td>14.602 ± 0.505</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>SO₄²⁻ (mg/L)</td>
<td>39.033 ± 0.300’</td>
<td>41.980 ± 0.916’</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>SRP (mg/L)</td>
<td>0.364 ± 0.011</td>
<td>0.391 ± 0.012</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Redox. (mV)</td>
<td>-</td>
<td>-</td>
<td>-316.5 ± 1.5</td>
<td>-145.0 ± 6.0</td>
</tr>
<tr>
<td>Dₘ (kg/L)</td>
<td>-</td>
<td>-</td>
<td>0.833 ± 0.062’</td>
<td>1.512 ± 0.008’’</td>
</tr>
<tr>
<td>CEC (cmol+/kg)</td>
<td>-</td>
<td>-</td>
<td>18.056 ± 2.945’</td>
<td>8.027 ± 0.703’</td>
</tr>
<tr>
<td>TP (mg/kg)</td>
<td>-</td>
<td>-</td>
<td>558.131 ± 30.251’’</td>
<td>203.489 ± 19.487’’</td>
</tr>
<tr>
<td>SOM (%)</td>
<td>-</td>
<td>-</td>
<td>5.05 ± 0.70’’</td>
<td>1.78 ± 0.10’’</td>
</tr>
</tbody>
</table>

Values are presented as mean ± SE.

DO: dissolved oxygen, TSS: total suspended solid, SRP: soluble reactive phosphorus, CEC: cation exchange capacity, TP: total phosphorus, SOM: organic matter content.

’p < 0.05, ’’p < 0.01, ’’’p < 0.001.
perature, NO₃⁻ and NO₂⁻ concentration over the weirs (Table 1). Water temperature ranged from 25°C to 26°C. The SRP concentrations increased over weirs without a statistical significance (from 0.364 ± 0.011 mg/L upstream to 0.391 ± 0.012 mg/L downstream of weirs). Significant variations over weirs were observed in pH, DO, TSS, and SO₄²⁻ concentration of the water (Table 1). The most striking difference was seen in the TSS profile across the weirs. TSS concentrations decreased from 24.0 ± 5.0 mg/L upstream to 9.3 ± 2.6 mg/L downstream of weirs. Water had a mean pH of 7.93 ± 0.50 upstream and of 9.19 ± 0.02 downstream of weirs. DO concentrations decreased from 11.4 ± 0.2 mg/L upstream to 8.9 ± 0.3 mg/L downstream of weirs. SO₄²⁻ concentrations, however, increased from upstream to downstream (Table 1).

3.2. Physicochemical Properties of Sediments

Overall, streambed sediments exhibited distinct characteristics over the weirs (Table 1). Sediments upstream of the weirs contained significantly high TP concentrations (558.131 ± 30.251 mg/kg) and high SOM contents (5.05% ± 0.70%) compared to sediments downstream of the weirs, which contained 203.489 ± 19.467 mg/kg and 1.78% ± 0.10% of TP and SOM, respectively. The CEC of sediments also decreased significantly downstream. In contrast, the redox potential and bulk density of sediments increased significantly downstream in which redox potentials were negative for both upstream and downstream sediments. The total concentrations of metals (Fe, Al, Mg, and Ca) decreased significantly downstream (Fig. 3(a)). In addition, sediments upstream of the weirs exhibited significantly higher activities of β-glucosidase, phosphatase, and sulfatase than sediments downstream of the weirs (Fig. 3(b)). However, no significant differences in pH and temperature were observed between sediments over the weirs. The mean temperature of sediments was from 23.3°C to 25.2°C and the mean pH was from 6.99 to 7.11 (Table 1).

Correlation analysis indicated that TSS negatively correlated with bulk density (r = −0.871, p < 0.5), whereas it positively correlated with SOM (r = 0.923; p < 0.01), TP (r = 0.913; p < 0.05), CEC (r = 0.846; p < 0.05), metal concentrations (r = 0.908, 0.960, 0.907, and 0.880 for Fe, Al, Mg, and Ca, respectively; p < 0.05) and extracellular enzyme activities (r = 0.861 and 0.899 for β-glucosidase and phosphatase, respectively; p < 0.05). In addition, TP significantly correlated with SOM (r = 0.912; p < 0.05), as well as with total Fe (r = 0.966; p < 0.01), Al (r = 0.945; p < 0.01), and Mg concentration (r = 0.953; p < 0.01) in sediments.

3.3. Phosphorus Fractions in Sediments

We found significantly higher concentrations of NH₄Cl-extractable P, NH₄F-extractable P, and NaOH-extractable P fractions in upstream sediments compared to downstream sediments of the weirs (Fig. 4). Moreover, NH₄Cl-extractable P, NH₄F-extractable P, and NaOH-extractable P fractions were significantly positively correlated to TSS (r = 0.954, 0.898, and 0.868, respectively; p < 0.05). NH₄Cl-extractable P or labile-P (the most available P fraction in sediments) only accounted for a small portion of the total P in sediments, which was about 4%–5% of the total P (Table 2). Phosphorus adsorbed and precipitated with Fe and Al (non-apatite P or NAIP), which was calculated as a sum of P fractions extracted with NH₄F, NaOH, and sodium citrate-dithionite-bicarbonate (CDB), was the dominant portion of the

![Fig. 3. Spatial distribution of (a) total concentration of metals and (b) extracellular enzyme activities in sediments across weirs (mean ± SE, n = 3). Asterisks denote significant differences between upstream and downstream sediments at *p < 0.05, **p < 0.01, ***p < 0.001.](http://eeer.org)

![Fig. 4. Distribution of P fractions in sediments across weirs (mean ± SE, n = 3). Asterisks denote significant differences between upstream and downstream sediments at *p < 0.05, **p < 0.01, ***p < 0.001.](http://eeer.org)

| Table 2. Percent contribution of P fractions to total P in sediments across weirs (n = 3) |
|---------------------------------|------------|------------|
|                                  | Upstream   | Downstream |
| Labile-P                         | 4.4 ± 0.5  | 5.3 ± 0.3  |
| NAIP                             | 59.8 ± 7.6 | 51.5 ± 1.7 |
| AIP                              | 25.5 ± 4.8 | 31.3 ± 2.7 |
| Residual-P                       | 10.2 ± 4.7 | 11.9 ± 3.0 |

Values are presented as mean ± SE. Labile-P = NH₄Cl-P; NAIP = NH₄F-P + NaOH-P + CDB-P; AIP = H₂SO₄-P; Residual-P = TP – (labile-P + NAIP + AIP).
fluxes between the sediments over the weirs did not occur under oxic condition (DO > 7 mg/L) (Fig. 5). The SRP release rates estimated under oxic conditions of the water columns showed significant relationships with $D_{b}$ ($r = –0.842; \ p < 0.05$), TP ($r = 0.822; \ p < 0.05$), and $H_2SO_4$-extractable P ($r = 0.973; \ p < 0.01$) (Table 5). On the other hand, in addition to the relations with bulk density ($r = –0.969; \ p < 0.01$) and TP ($r = 0.918; \ p < 0.01$), SRP release rates under anoxic condition also showed significant relationships with $NaOH$-extractable P ($r = 0.817; \ p < 0.05$), Fe ($r = 0.845; \ p < 0.05$), Al ($r = 0.887; \ p < 0.05$), and Mg concentrations ($r = 0.886; \ p < 0.05$) (Table 5).

The SRP release rates under different water pH, temperature, and DO conditions in 18 sediment cores collected from one randomly-chosen location are shown in Fig. 6. SRP release rates were greater under anoxic condition at higher temperature and un-neutral pH. Specifically, the impacts of DO and temperature were illustrated clearly with the incubation experiments at pH 7.0 (Fig. 6(b)). Under oxic condition and neutral pH, P was sufficiently trapped within the sediments at 15°C, while an immediate release of SRP occurred when the temperature increased to 25°C. The SRP release rates at 25°C under anoxic condition were also discernibly higher than those under oxic condition.

Fig. 5. (a) Cumulative soluble reactive phosphorus (SRP) releases and (b) SRP release rates in sediments across weirs at 25°C, water pH 7.0 (±0.5) under oxic/anoxic conditions. The dashed line indicates the switch from oxic (DO > 7 mg/L) to anoxic condition (DO < 0.5 mg/L) in water columns of sediment cores.

Fig. 6. Soluble reactive phosphorus (SRP) release rates (mean ± SE, n = 2) at different oxic/anoxic conditions, temperatures (15°C and 25°C), and pH: (a) pH 3.0, (b), pH 7.0, and (c) pH 11.0.
Comparison of SRP release rates at un-neutral pH

The variations of SRP release rates at un-neutral pH strongly depended on the interacting effects of temperature and DO (Fig. 6(a) and 6(c)).

Table 3. Empirical determination factors of the Freundlich isotherm for sediments across weirs (n = 3)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Upstream</th>
<th>Downstream</th>
</tr>
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<tbody>
<tr>
<td>$K_c$ (L/g)</td>
<td>0.053 ± 0.006**</td>
<td>0.145 ± 0.002**</td>
</tr>
<tr>
<td>n</td>
<td>0.879 ± 0.032*</td>
<td>0.567 ± 0.057*</td>
</tr>
<tr>
<td>$K_s$ (L/g)</td>
<td>0.022 ± 0.002</td>
<td>0.368 ± 0.195</td>
</tr>
<tr>
<td>NAP (µg/g)</td>
<td>6.715 ± 0.583*</td>
<td>1.026 ± 0.803*</td>
</tr>
<tr>
<td>EPCn (mg/L)</td>
<td>0.273 ± 0.011***</td>
<td>0.003 ± 0.001***</td>
</tr>
<tr>
<td>SRP (mg/L)</td>
<td>0.364 ± 0.011</td>
<td>0.391 ± 0.012</td>
</tr>
</tbody>
</table>

* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. 

Table 4. Repeated-measures ANOVA for SRP release rates in sediments across weirs

<table>
<thead>
<tr>
<th>Source</th>
<th>F (repeated measures)</th>
</tr>
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<tr>
<td>Between subjects</td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td>65.993**</td>
</tr>
<tr>
<td>Within subjects</td>
<td></td>
</tr>
<tr>
<td>Incubation (oxic/anoxic)</td>
<td>69.634**</td>
</tr>
<tr>
<td>Incubation × Site</td>
<td>25.791**</td>
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</tbody>
</table>

SRP: soluble reactive phosphorus. 
* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. 

Table 5. Pearson correlation between SRP release rates and other parameters

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Oxic P release</th>
<th>Anoxic P release</th>
</tr>
</thead>
<tbody>
<tr>
<td>$D_s$</td>
<td>-0.842*</td>
<td>-0.969**</td>
</tr>
<tr>
<td>TP</td>
<td>0.822*</td>
<td>0.918**</td>
</tr>
<tr>
<td>NaOH-P</td>
<td>0.571</td>
<td>0.817</td>
</tr>
<tr>
<td>$H_2SO_4-P$</td>
<td>0.973**</td>
<td>0.807</td>
</tr>
<tr>
<td>Fe</td>
<td>0.667</td>
<td>0.845*</td>
</tr>
<tr>
<td>Mg</td>
<td>0.666</td>
<td>0.886*</td>
</tr>
<tr>
<td>Al</td>
<td>0.618</td>
<td>0.887</td>
</tr>
</tbody>
</table>

$D_s$: bulk density, TP: total phosphorus, SRP: soluble reactive phosphorus. 
* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. 

4. Discussion

SRP concentrations in stream water were consistently high (> 0.3 mg/L) and did not vary across the weirs in Tanchun Stream. Urban streams are usually described with elevated concentrations of nutrients and contaminants [42]. Phosphorus concentrations approaching 0.3 mg/L greatly enhance the growth of macrophytes and algal communities [43]. Thus, potentially increasing internal loading of P from streambed sediments upstream of weirs could lead to streams becoming eutrophic.

The lower $D_s$ of sediments upstream of weirs, compared to downstream sediments, indicated that the upstream sediments might be more susceptible to resuspension than downstream sediments; thus necessarily enabling P release into the water column [44]. Organic-rich sediments usually have lower $D_s$ and contain higher phosphorus concentration [45]. The Fe, Al, Mg, and Ca ions associated the organic matter in sediments tend to retain P by adsorption on surfaces or by forming insoluble complexes and precipitates [45]. We found that the amounts of organic matter and TP, as well as total metals (Fe, Al, Mg, and Ca), were significantly greater in sediments upstream of weirs. Correlation analysis also demonstrated that TP concentrations were associated with organic matter and metallic ions in sediments. Furthermore, the significant positive correlation between TSS and SOM, as well as TP, suggests that weirs acted as important physical barriers to the downstream transport of P in streams.

The significant difference of total P concentrations between sediments upstream and downstream of weirs was mainly due to the differences in P adsorbed and precipitated with Fe and Al in sediments. Labile-P contributed only a small portion of the variation of TP concentrations in sediments across the weirs. Frequent transports of sediments occur during moderate flows in streams [46]. In addition, Fe and Al-bound P fractions in a wide range of sediments are most concentrated near the sediment surface [47-49]. Therefore, any change of stream flows, such as those across weirs, could strongly affect the transport of surface sediments, thus of Fe and Al-bound P fractions. TSS concentrations significantly correlated with NH$_4$F-extractable P and NaOH-extractable P fractions, providing further evidence that the retention of P in streambed sediments were strongly depending on the sediment transport and hydrological regime in streams. These results also implied that the alteration of hydrology and consequent sediment transport could lead to the enrichment of potentially-labile P fractions in streambed sediments of urban streams. If phosphorus is retained in sediments in the potentially available forms, such as phosphorus associated with Fe and Al, it may constitute a significant source to the water column under certain changes of redox potential and pH [22, 50]. Thus, the majority of P pools in sediments trapped above the weirs, which was composed of labile, Fe and Al-bound P forms, would be fairly unstable and potentially affect the stream water quality. Potential release of P from sediments is most likely to occur in the summer, when oxygen demand and primary productivity are highest [22], or at stream sites with elevated organic matter during low (or pulse) flow conditions [51].

Extracellular enzyme activities were often used to measure the limitation of nutrients in ecosystems [52, 53]. In this study, we used these parameters as indicators of the microbial processing of organic matter in sediments of urban streams. The significantly higher activities of β-glucosidase and phosphatase in sediments upstream of the weirs, compared to downstream sediments, indicated the high oxidation potential of organic matter and reduction of various oxidants. The SRP concentrations in the interstitial water and the water column are assumed to be controlled partly by the mineralization of organic matter in sediments. During the mineralization of organic matter in sediments, bacteria consumed $O_2$, $NO_3^-$, and $SO_4^{2-}$ and provided the necessary conditions for the abiotic or biotic reduction of ferric ion (Fe$^{3+}$), the subsequent release of P, and precipitation of iron sulfide (FeS) [54].
The relative contribution of abiotic or biotic processes to the P retention and remobilization in sediments upstream of weirs might also be changed compared to natural streambed sediments (in sensu Lottig and Stanley [55]). The role of biotic processes controlling the SRP concentration in the interstitial water of sediments is often assumed to be limited [54]. However, Lottig and Stanley [55] demonstrate that the type and strength of sediment-associated processes determining the SRP concentrations in sediments are potentially altered due to changes in sediment composition (i.e., fine versus coarse particles). Accordingly, we propose that the P remobilization of sediments with higher silt and clay contents above the weirs will be dominated by abiotic processes, whereas biotic processes will mainly account for the P remobilization in naturally sandy sediments. Over time, if the abiotic P sink of sediments above the weirs is filled up due to the increase of P loads and the absence of scouring floods, the sediments above the weirs will shift to release P. The biotic P sink in natural sediments, however, can be refreshed with frequently new sediment input. Therefore, as the weirs can limit the sediment turnover in streams, they can also diminish long-term P buffering capacity of streams. This hypothesis needs to be evaluated in further experiments.

The EPC of sediments lower than the ambient SRP concentrations demonstrated that the sediments were still able to retain phosphorus. However, the buffer intensities of sediments with respect to any external loading of dissolved phosphorus significantly changed across the weirs. The lower Kₑ (or lower Kᵢ) and the higher EPCₑ of sediments upstream of the weirs indicated that sediments above weirs served as a less sustained sink of phosphorus to the water column. Sorption behavior of sediments in respect to Kₑ (or Kᵢ) and EPC, was argued to relate to the total exchangeable P pool of sediments [44]. It is assumed that organic P contributed less to the exchangeable P pool of our sediments, which is possibly reasonable in this study because the residual P has accounted for only 10% of the total P of sediments. The total amount of exchangeable P in our sediments can be calculated as a sum of NH₄Cl-, NH₄F-, NaOH- and CDB-extractable P fractions. As a result, the mean exchangeable P concentration is 357.966 ± 44.383 mg/kg of upstream sediments, which is higher than that of downstream sediments (116.029 ± 13.295 mg/kg). Furthermore, we found from the batch experiments that NAP concentrations were significantly higher in sediments upstream of weirs compared to downstream sediments. Thus, our results support the idea that, rather than Fe-P appearing predominantly in calcareous sediments, Fe-Al-P was predominantly in non-calcareous sediments [64]. Compared to Al-bound P, Fe-bound P was considered as more potentially labile [65]. High concentrations of aluminum hydroxides in sediments could even prevent the P release [66]. This implies that if the sediments trapped above weirs are calcareous sediments containing the same amounts of P, they may contribute to a higher risk of eutrophication in streams.

5. Conclusions

Weirs could increase P retention in urban streams. Phosphorus was mostly retained in sediments above weirs in both exchangeable and non-exchangeable forms, which potentially constituted a significant P source to waters under changing environmental conditions. Remobilization of sediment P above weirs thus can promote eutrophication in urban streams. Since interactions between sediments and stream water are highly heterogeneous, quantifying the significance of impoundment effects on P retention and remobilization needs to be intensified to gain valuable insights into the P transport and transformation as moving through watersheds.
Acknowledgments

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