

1 **Intensive agricultural management-induced**  
2 **subsurface accumulation of water extractable**  
3 **colloidal P in a Vertisol**

4 Shouhao Li<sup>a,b</sup>, Shuiqing Chen<sup>a</sup>, Shanshan Bai<sup>a</sup>, Jinfang Tan<sup>a,b\*</sup>, Xiaoqian Jiang<sup>a,b\*</sup>

5 <sup>a</sup> School of Agriculture, Sun Yat-sen University, Guangzhou, Guangdong 510275, PR  
6 China

7 <sup>b</sup>Modern Agricultural Innovation Center, Henan Institute of Sun Yat-sen University, PR  
8 China

9 Corresponding to: [jiangxq7@mail.sysu.edu.cn](mailto:jiangxq7@mail.sysu.edu.cn) and [tanjf7@mail.sysu.edu.cn](mailto:tanjf7@mail.sysu.edu.cn)

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11 **ABSTRACT:**

12 Long-term excessive application of mineral fertilizer leads to phosphorus (P)  
13 accumulation, increasing the risk of P migration and loss from soil profile. The colloids  
14 in the soil profile are important carriers for P migration due to their high P adsorption  
15 and transport capacity. It is not clearly understood how colloidal P (CP) is distributed  
16 in subsoils (<1.2 m) of a Vertisol, contributing to subsurface P loss. Understanding the  
17 depth sequence distribution and speciation of colloidal P in the soil profile is critical for  
18 a comprehensive assessment of P loss. In this study, water extractable colloids (WECs)  
19 with the size of 0.35-2  $\mu\text{m}$  were obtained from 0-120 cm soil profile by sedimentation  
20 and centrifugation scheme. The dissolved reactive P (DRP) and dissolved total P (DTP)  
21 in soil supernatant with particle sizes <0.35  $\mu\text{m}$  were measured by molybdate blue  
22 colorimetry. Solution <sup>31</sup>P nuclear magnetic resonance (NMR) and P K-edge XANES  
23 were used to characterize the species and distribution of CP in the soil profile of  
24 fertilized farmland. Total and available P in bulk soil and colloids decreased with soil  
25 depth. The organic P (OP) contained 97-344 mg kg<sup>-1</sup> per bulk soil and 110-630 mg/kg  
26 per WECs. The OP in soil profile consists of orthophosphate monoesters and diesters  
27 primarily according to NMR results. It suggested that OP in WECs from subsoils might

28 be affected by the translocation of CP from surface soils, probably due to soil  
29 acidification and preferential flow caused by swelling-shrinkage clays including  
30 montmorillonite and nontronite detected by X-ray powder diffractometer (XRD) result.  
31 Additionally, the more negative zeta potential of surface soil colloids suggests the high  
32 mobility of colloidal P towards the subsoils. The CP concentrations for  $<2 \mu\text{m}$  was  
33 about 38-93 mg P /kg per bulk soil, which is 6-37 folds of DRP concentrations,  
34 suggesting that CP plays a dominant role in P transport within the soil profile. The  
35 relatively small fraction of orthophosphate diesters suggests limited P assimilation by  
36 microorganisms for the accumulation of WECs containing organically-bound P in  
37 subsoils. The P K-edge XANES results indicated that the proportions of Al-P, Fe-P, and  
38 inositol hexakisphosphate (IHP) of WECs decreased but hydroxyapatite (HAP)  
39 increased with soil depth. This study showed that inorganic and organic P migrated  
40 from surface to deeper layer along the soil profile, with soil colloids having a significant  
41 effect on P migration from both surface and subsurface layers. The findings have an  
42 important significance for soil P migration evaluation and agricultural non-point source  
43 pollution control in Vertisols.

44 **Keywords:** Subsurface soil; Colloidal P, Organic P, Solution  $^{31}\text{P}$ -NMR, P K-edge  
45 XANES

## 46 **INTRODUCTION:**

47 Phosphorus (P) as one of essential macronutrients for crops is strongly immobilized  
48 with inorganic and organic soil components (Arai and Sparks, 2007). Vertisol (locally  
49 known as Shajiang black soil) covers approximately 3.13 million hectares in the Huang-  
50 Huai-Hai Plain of China (Wei et al., 2018). It is characterized by low soil organic matter  
51 (SOM), low water-air permeability, poor fertility, and strong swelling-shrinkage  
52 properties, thus contributing to low and medium crop yields (Chen et al., 2020).  
53 Therefore, high-intensity agriculture practices such as excessive fertilization have been  
54 applied for decades to maintain grain yields. Additionally, artificial ditches ( $\sim 1$ -1.5 m  
55 depth) are usually dug out in the edge of the field to facilitate drainage. However, long-  
56 term excessive input of mineral fertilizers may result in considerable P accumulation in

57 agricultural soils and the artificial ditches bring increasing risk of P losses into surface  
58 water and groundwater, causing serious environmental issues such as outbreaks of  
59 cyanobacteria and eutrophication (Whalen and Chang, 2001; Koopmans *et al.*, 2003;  
60 Hansen *et al.*, 2004).

61 In addition to the transport of soil surface P (Pote *et al.*, 1996), transport of soil  
62 subsurface P has also been considered as a crucial pathway to waterways (Jorgensen  
63 and Fredericia, 1992; Kronvang *et al.*, 1997; Xue *et al.*, 1998; Hens and Merckx, 2001;  
64 Williams *et al.*, 2016; Jiang *et al.*, 2021). Some researchers have reported that high  
65 rainfall events promote the losses of P from tile drainage (1.0-1.4 m depth) (Royer *et al.*,  
66 2006). In previous study in USA, colloidal P was the dominant P fraction of total P  
67 in tile water during high rainfall events (Jiang *et al.*, 2021). The colloids were found to  
68 carry over 1000 mg of P kg<sup>-1</sup>, which was dominated in the transported P from subsurface  
69 soil (Jiang *et al.*, 2021).

70 Currently, only a few papers have investigated dissolved and colloidal P distribution in  
71 subsoils which involved forest soils and Mollisols (Gentry *et al.*, 2007; Wang *et al.*,  
72 2020; Li *et al.*, 2022). However, it is still not clear about the distribution of colloidal P  
73 as well as their speciation in the soil profile of Vertisol. Furthermore, whether the  
74 transport of colloidal P from topsoils to subsoils occurs and contributes to total P in  
75 subsurface flow is not clearly understood. Considering that the presence of abundant  
76 shrink and swell clays in Vertisol, the translocation of dissolved and colloidal P from  
77 surface soil to subsoils by preferential flow is expected as an important P transport  
78 pathway and leads to spatial redistribution within soil profiles. Additionally, the  
79 artificial ditches could also facilitate the transport of dissolved and colloidal P from  
80 both surface and subsurface soil to surface waters, which is an important non-point  
81 source of eutrophication. Furthermore, the intensive input of fertilizer (~600 kg per  
82 hectare compound fertilizer, N-P-K: 28-6-6) was also expected to facilitate the release  
83 of colloidal P (Liang *et al.*, 2016).

84 It is expected that colloidal P could be derived from surface soils if the colloids in  
85 subsurface soil contains organic P (Li *et al.*, 2022). The aim of this research was to

86 explore (1) the physicochemical properties and speciation of colloidal P in the soil  
87 profile and (2) whether the colloidal organic and inorganic P exist in the subsoil and  
88 play important roles for the transport of P in intensively managed Vertisol. We assumed  
89 that the transport of colloidal P with inorganic and organic forms from the topsoil to  
90 subsoil of Vertisol is certainly possible as a result of long-term intensive agricultural  
91 management, which could bring an unavoidable risk on transport and loss of subsurface  
92 P in agricultural soil. This information is the first for the assessment of colloidal P  
93 release potential from subsoils in a Vertisol and is valuable for the construction of  
94 sustainable strategies to control agricultural P loss.

## 95 **2. MATERIALS AND METHODS**

### 96 **2.1 Site description and soil sampling**

97 The selected site was located at the agricultural study site in Pingyu County, Henan  
98 Provinces, China with precipitation of 904.3 mm and annual mean temperature of 15 °C,  
99 respectively. The soil is classified as a Vertisol according to USDA soil Taxonomy (Soil  
100 Taxonomy, USDA , 2010). Soil samples defined as A, B, C, and D were collected from  
101 four sites: Chenji Village (33°00'N and 114°51'E), Dingying Village (33°09'N and 114°  
102 49'E), Xinggang Village (32°99'N and 114°84'E), and Hanqiao Village (32°56'N and  
103 114°49'E). Samples were taken in Autumn 2021 along four vertical profiles at five or  
104 six different depths, i.e., 0-20, 20-40, 40-60, 60-80, 80-120 cm (or 80-100 and 100-120  
105 cm), which are denoted as depth 1, 2, 3, 4, 5 (or 5 and 6), respectively. The cultivation  
106 system in the region involved rotating winter wheat and summer maize crops. The  
107 fertilizers for winter wheat contained 750 kg/hm<sup>2</sup> compound fertilizer (N-P-K: 15-15-  
108 15) and 300 kg/hm<sup>2</sup> urea. Those for summer maize included 600 kg/hm<sup>2</sup> compound  
109 fertilizer and 225 kg/hm<sup>2</sup> urea. The samples were air-dried, ground, and passed through  
110 2-mm sieve before analysis.

### 111 **2.2 Physicochemical characterization of the soils**

112 Soil pH was assessed using a pH meter with a soil-to-water ratio of 1:2.5. Soil total

113 carbon (C) and nitrogen (N) levels were analyzed with an elemental analyzer (Jin *et al.*,  
114 2016). The equipment utilized for measuring total organic carbon (TOC) is TOC-VCPH,  
115 which employs the combustion oxidation and non-dispersive infrared absorption  
116 method. Oxalate extracted P (i.e. OA-P) was extracted by ammonium oxalate and oxalic  
117 acid (Jiang *et al.*, 2015) and was considered as the P bonded to amorphous, poorly  
118 crystalline and organo-Fe/Al (hydr)oxides (Masiello *et al.*, 2004; Kleber *et al.*, 2005;  
119 Neubauer *et al.*, 2013). The dithionite-citrate-bicarbonate (i.e. DCB) extracted P was  
120 extracted by sodium citrate solution, sodium hydrogen carbonate solution and sodium  
121 dithionite (Jiang *et al.*, 2015) and was considered as P associated with organically bound,  
122 amorphous and crystalline Fe oxides. The activities of acid and alkaline phosphatase,  
123 denoted as ACP and ALP respectively, were determined by performing *p*-nitrophenyl  
124 phosphate assays on moist soil samples at two pH conditions, 6.5 and 11 respectively  
125 (Tabatabai *et al.*, 1969). Soil available P was extracted with 0.5 M NaHCO<sub>3</sub> at pH 8.5  
126 (Van Lierop, 1988) and measured by molybdate blue colorimetry (Murphy and Riley,  
127 1962). Total inorganic P (TIP) was extracted by sulfuric acid and dilute sodium  
128 hydroxide solution separately via the sequential extraction method by Kronvang *et al.*  
129 (1997) and measured by molybdate blue colorimetry. For total P (TP), the extracts  
130 that were acquired from the acid and base treatments were treated with potassium  
131 persulfate and sulfuric acid before molybdate blue colorimetry and the sum of digested  
132 extracts was defined as TP. The total organic P (TOP) was the difference between TP  
133 and TIP.

### 134 **2.3 Soil particle separations and characterization of water extractable** 135 **colloids**

136 The soil samples of A and B with different depths were fractionated by the method  
137 described by Séquaris *et al.* (2003). Briefly, 100 g of soil was dispersed in 200 mL Milli-  
138 Q water in a 1 L Duran bottle and shaken for 6 h at 150 rpm. Then 600 mL Milli-Q  
139 water was added and mixed to settle. The particles >20 µm and 2-20 µm were obtained  
140 by removing the supernatant after settling for 6 min and 12 h, respectively. The  
141 supernatant was subsequently spun at 3500×g for 5 min to obtain the water extractable

142 colloids with size of 0.35-2  $\mu\text{m}$  (calculated based on Stokes' law and the density of  
143 particle is assumed as 2.65  $\text{g cm}^{-3}$ ). The final supernatant only contained the remaining  
144 fine colloids with an average hydrodynamic diameter of  $351.3 \pm 6.6$  nm (according to  
145 dynamic light scattering [DLS] result) as well as the electrolyte phase. The dissolved  
146 reactive P (DRP) and dissolved total P (DTP) were measured by molybdate blue  
147 colorimetry (Murphy and Riley, 1962) before and after the digestion of potassium  
148 persulfate and sulfuric acid for the final supernatant.

149 To elucidate the inorganic and organic P species, the extracted colloidal samples after  
150 lyophilization from sample A and B with different depths were selected for the NMR  
151 analysis. The colloid samples (1 g) were mixed with 10 mL of solution containing 0.25  
152 M NaOH and 0.05 M  $\text{Na}_2\text{EDTA}$  for 4 h (Jiang *et al.*, 2017). After that, extracts were  
153 centrifuged at  $10000 \times g$  for 30 minutes. The P, Fe, and Mn contents in the supernatant  
154 were measured by inductively coupled plasma optical emission spectroscopy (ICP-  
155 OES). The rest supernatant was freeze-dried and then 100 mg freeze-dried extracts were  
156 dissolved in 0.1 mL  $\text{D}_2\text{O}$  and 0.9 mL of a solution containing 1.0 M NaOH and 0.1 M  
157  $\text{Na}_2\text{EDTA}$ . After being prepared, the samples underwent centrifugation for 20 minutes  
158 at  $10000 \times g$ . The resulting supernatant was subsequently analyzed using a Bruker 500-  
159 MHz spectrometer with a 5 mm NMR tube. The NMR parameters contained 0.68 s  
160 acquisition time,  $90^\circ$  pulse width, 8000 scans, and proton decoupling. The relaxation  
161 time, which fell within the range of 3-6 seconds, was estimated by correlating  
162  $\text{P}/(\text{Mn}+\text{Fe})$  with spin-lattice relaxation times (McDowell *et al.*, 2006). The spiking  
163 samples with myo-inositol hexakisphosphate (myo-IHP),  $\alpha$ - and  $\beta$ - glycerophosphate,  
164 and adenosine monophosphate were cited to facilitate peak identification (Bai *et al.*,  
165 2023). The  $\alpha$ -,  $\beta$ -glycerophosphates and mononucleotides (Glyc+nucl) were classified  
166 as orthophosphate diesters rather than to monoesters (Young *et al.*, 2013; Liu *et al.*,  
167 2018). The area under each peak was determined by integrating spectra that were  
168 processed with a line broadening of 2 and 7 Hz. Mestrenova 10.0.2 software was used  
169 to process all spectra. The concentrations of individual P species were calculated by  
170 multiplying  $^{31}\text{P}$ -NMR proportions by the total NaOH- $\text{Na}_2\text{EDTA}$  extractable P

171 concentration. Additionally, zeta potential of colloids from sample A and B with  
172 different depths were measured using a Zetasizer (Malvern). The X-ray powder  
173 diffractometer (XRD, Empyrean) was selected to identify mineral compositions for soil  
174 colloids in the  $2\theta$  range from  $3^\circ$  to  $90^\circ$  with the scan step size of  $0.026^\circ$  and the scan  
175 rate of  $10^\circ \text{ min}^{-1}$ .

176 To elucidate the P bonding fractions in WECs, the P K-edge X-ray absorptions near-  
177 edge structure (XANES) measurements were performed at Beamline 4B7A of the  
178 Beijing Synchrotron Radiation Facility, Beijing, China. The following standard samples  
179 were chosen: aluminum phosphate (Al-P,  $\text{AlPO}_4$ ), iron phosphate dihydrate (Fe-P,  
180  $\text{FePO}_4 \cdot 2\text{H}_2\text{O}$ ), inositol hexakisphosphate (IHP), and hydroxyapatite (HAP,  
181  $\text{Ca}_5(\text{PO}_4)_3\text{OH}$ ). The soil spectra were collected using a SiLi detector in PFY mode,  
182 providing detailed information about the fluorescence yield. The spectra of P standard  
183 samples were measured in total electron mode without self-absorption. Multiple spectra  
184 (three repetitions for soil samples) were collected and averaged. All XANES spectra  
185 were measured by Athena (0.9.26). The absolute energy scale was calibrated to 2149  
186 eV (E0) as the maximum energy of the first peak in the first derivative spectrum of  
187  $\text{AlPO}_4$  (Beauchemin *et al.*, 2003). Linear combination fitting (LCF) of the soil P spectra  
188 was conducted in the relative energy range between -10 and 30 eV. The goodness of fit  
189 was judged by the chi-squared values and R values, and P standard samples yielding  
190 the best fit were considered as the most possible P species in the investigated soil  
191 samples.

## 192 **2.4 Statistical analysis**

193 One way analysis of variance (ANOVA) as a single factor analysis of variance was used  
194 to test significant differences of soil indicators with different soil profiles. Tukey's  
195 honesty significance difference (HSD) test was used, and the significance level was  
196 0.05. SPSS 25.0 and Excel software were used for statistical analysis. These data were  
197 created using OriginPro 2021 (OriginLab Corp., Northampton, MA, USA).

## 198 **3. RESULTS AND DISCUSSION**

### 199 **3.1 Physicochemical characterization and P distribution of Soil profile**

200 With the increase of soil depth in all samples, the soil pH exhibited a significant overall  
201 increase from acidic to alkaline conditions, ranging from 4.88 to 8.37 (Table 1). The  
202 serious acidity of the topsoil (0-20 cm) is probably due to increasing release of protons  
203 by nitrification processes after excessive application of nitrogen fertilizers (750 kg/hm<sup>2</sup>  
204 compound fertilizer and 300 kg/hm<sup>2</sup> urea per year in the studied area) and the  
205 continuous removal of base cations by crop harvest (Guo *et al.*, 2010). The calcareous  
206 nature of the studied soil contributed to subsoils with pH values of slight alkalinity. The  
207 contents of total C (from 1.17 to 0.25%) and total N (from 0.14 to 0.03%) decreased  
208 significantly from topsoil to subsoil (Table 1). The accumulation of total C and N in  
209 surface soil (0-20 cm) could be related to the increased organic matter contents by  
210 biomass inputs from crop residue and N fertilization. The deepest layer soil (80-120 cm)  
211 contained the highest pH values and lowest total C and N contents, suggesting that the  
212 subsurface soil (20-80 cm) was also influenced by these intensive agricultural  
213 managements to some extent. The activities of acid phosphatase (ACP) was as high as  
214 1177 µg/(g\*h) in topsoil and decreased with the depth of soil layer as low as 96.2  
215 µg/(g\*h) in subsoil (Table 1). The activities of both acid and alkaline (ALP)  
216 phosphatase decreased significantly with depth for all the samples. The ACP was higher  
217 in the surface soil (0-20 cm) but was lower in the subsurface soil (20-120 cm) compared  
218 to ALP. Acid phosphatase activity, mainly released by plant roots and fungi, is  
219 predominant in acidic soils (Eivazi *et al.*, 1977; Juma and Tabatabai, 1977; Arenberg *et*  
220 *al.*, 2020). Alkaline phosphatase activity, primarily produced by soil microbes, is  
221 optimized in neutral and alkaline conditions (Juma and Tabatabai, 1988; Dick *et al.*,  
222 2000; Krämer and Green, 2000). The higher ACP in surface soil may be attributed to  
223 acidic pH and the rhizosphere effect, where plant roots and fungi are easily concentrated  
224 (Häussling *et al.*, 1989). Thus, lower ACP in the subsurface soil (> 20 cm) was due to  
225 increasing pH with depth.

226 Oxalate extractable P concentration ranged from ~30-162 mg kg<sup>-1</sup> (~7-45% of TP) in  
227 all soil sample (Table 1). The oxalate extractable P concentration in the surface soils  
228 was ~127 to 162 mg/kg and it accounted for 19-27% of TP and ~76 to 98 % of DCB

229 extractable P, suggesting that amorphous Fe/Al oxides bounded P was dominated for  
230 the Fe-P in the surface soils compared to crystalline Fe/Al oxides. Many studies have  
231 reported that the majority of P was associated with amorphous Fe /Al oxides fractions  
232 in many soil types (Rick and Arai, 2011; Jiang *et al.*, 2015). It has been reported that  
233 specific anion adsorption such as P suppressed the transformation of amorphous Fe  
234 oxides to crystalline Fe oxides (Biber *et al.*, 1994), supporting the lower amount of P  
235 associated with crystalline Fe oxides.

236 The concentrations for TIP, TOP, and TP are included in Fig.1. The total P  
237 concentrations in all soil samples ranged from approximately 230 to 670 mg kg<sup>-1</sup>, and  
238 exhibited a generally decreasing trend with the increase of soil depth (Fig. 1). TIP and  
239 TOP accounted for ~48 to 65% and ~35 to 52% of TP, respectively. Organic P  
240 accumulation (~97 to 344 mg kg<sup>-1</sup>) was measured in all soil depths. The OC/OP ratio  
241 of > 200:1 is favorable for P immobilization (Dalal and Hallsworth, 1977; Sanyal and  
242 De Datta, 1991). The OC/OP ratio of soil samples in this study was about 4.9 to 49.6,  
243 which promoted P mineralization. Thus, the accumulation of organic P in the surface  
244 soil would be due to the increasing organic matter contents, rather than the biological  
245 immobilization of P. Gradual transport of OP from surface to subsoils could contribute  
246 to the accumulation of OP in the subsoils. Available P was very high with ~20 to 40 mg  
247 kg<sup>-1</sup> in the surface soils but decreased dramatically with the increase of soil depth in all  
248 samples (Fig. 1). The available P content in the topsoil has been classified as "high",  
249 and a threshold of 20 mg/kg has been regarded as optimal growth level for crops (Li *et*  
250 *al.*, 2015). This implies that the surface soils (0-20 cm) contained enough available P  
251 for the growth of winter wheat and summer maize.

### 252 **3.2 Physicochemical properties of water extracted colloids (WECs)**

253 Considering that all four samples showed similar physicochemical properties and P  
254 distribution, we further investigated the colloids of samples A and B with different  
255 depths. It is crucial to mention that the WECs are colloids that are well-defined with  
256 size of 0.35-2 μm operationally. The major soil series are sandy loam and loam, with  
257 the colloidal mass of 5.3 to 8.3% (Table 2). The colloidal mass increased with depths

258 for both samples and previous researcher has reported a similar tendency for the  
259 recovery rate of WECs in dark-colored Mollisols (Li *et al.*, 2022). Notably, no colloids  
260 existed in the depth > 60 cm for sample B, suggesting that the generation of colloids  
261 from weathered minerals in subsoils was limited. It suggested that the colloids in the  
262 depth >60 cm for sample A originated from the transport of upper soil colloids mainly.  
263 The XRD results further verified it considering that the colloids of sample A and B with  
264 all depths contained the same mineral composition (Fig. 3). The minerals in all colloids  
265 included montmorillonite, nontronite, and illite, indicating the existence of swelling-  
266 shrinkage clays. These secondary minerals all have a significant adsorption capacity for  
267 P due to great surface area and the structural binding sites of clay mineral edges (Gérard,  
268 2016; Chen *et al.*, 2020). Values of zeta potential for WECs were approximately -20  
269 mV at the surface soil, but the values increased with depth (Table 2). A higher P  
270 concentration in surface soils could facilitate P special adsorption to minerals of WECs  
271 such as iron oxyhydroxide (e.g. hematite, goethite, and ferrihydrite) and aluminosilicate  
272 minerals (Arai and Sparks, 2001; Celi *et al.*, 2001; Jiang *et al.*, 2015), causing the  
273 surface charge lower compared to that of WECs in subsurface soils. The more negative  
274 values of zeta potential at surface soil suggests the higher mobility of colloidal P from  
275 the surface soil to the subsoils. The cation eluviation such as  $\text{Ca}^{2+}$  from the topsoil to  
276 subsoil and the subsequence illuviation in the subsoil probably affect cation-particle  
277 interactions and contribute to the more positive zeta potential in subsoil, as shown by  
278 the increase of hydroxyapatite with depth detected by P K-edge XANES results (see  
279 section 3.4). It is important to note that DRP accounted for only 29-51% of DTP (Table  
280 2). Furthermore, the sum of colloidal P concentration for <0.35  $\mu\text{m}$  (i.e. the difference  
281 between DTP and DRP) and 0.35-2  $\mu\text{m}$  was about 38-93 mg/kg soil, which is 6-37 folds  
282 of DRP. It suggested that the potential contribution of CP to transport P in both surface  
283 and subsurface soil environment is important and should not be overlooked. However,  
284 it is not excluded the transport of dissolved P with subsequent sorption to the colloidal  
285 fraction of subsoils.

### 286 **3.3 Inorganic and organic P distribution in WECs with soil depth**

287 At each sampling site, the concentration of TP in WECs was found to be the highest in  
288 the topsoil layer (0-20 cm), with a range of 1150 to 1300 mg kg<sup>-1</sup> (Fig. 3), which were  
289 significantly higher than the TP in bulk soils (~600-700 mg kg<sup>-1</sup>). The soil colloids  
290 enriched in secondary clays such as montmorillonite, nontronite, and illite as shown in  
291 XRD results could readily immobilize P. Generally, the TP in WECs at each site  
292 decreased with the increase of soil depth. The concentration of total IP in WECs was  
293 found to be high (ranging from 680 to 730 mg kg<sup>-1</sup>) in the topsoil and displayed a  
294 decreasing trend with the increase of soil depth across all sites. As predicted, total OP  
295 concentration was high (430-630 mg kg<sup>-1</sup>) in the colloids of surface soil. This is  
296 associated with the high OC concentration in the surface soil, and P was probably  
297 immobilized in the organic matter of soil. The declined tendency of OP concentrations  
298 with soil depth was probably due to the decreasing OM contents. It is noteworthy that  
299 OP was still presented in the WECs of subsoils where the OC/OP is less than 300 that  
300 could not support immobilization for the accumulation of OP (Table 1 and 3). This  
301 finding implies that colloidal-bound OP has the potential to be transported from topsoil  
302 to subsoil layers.

### 303 **3.4 Solution <sup>31</sup>P NMR analysis and P K-edge XANES analysis**

304 The <sup>31</sup>P NMR spectroscopy spectra of WECs were presented in Table 3. It is worth  
305 mentioning that NaOH-EDTA extracted P in the NMR analysis was below 100 %.  
306 Therefore, the concentrations of OP and IP did not correspond with the chemical  
307 digestion data presented in Fig. 2. For inorganic P, orthophosphate was found in all  
308 samples but pyrophosphate was only found in the surface soil. Pyrophosphate was  
309 present in live fungal tissue and was easily decomposed (Koukol *et al.*, 2008). In  
310 addition to inorganic P, the OP of WECs in all depths for sample A and B contained  
311 similar species, including orthophosphate monoesters (36-128 mg kg<sup>-1</sup>) and diesters (0-  
312 89 mg kg<sup>-1</sup>) according to NMR results (Table 3). It was clear that OP existed in colloids  
313 for both surface and subsoils, suggesting that OP in WECs from subsoils was affected  
314 by the translocation of CP from surface soils (Li *et al.*, 2022). Colloids containing clay  
315 minerals in the soil profile could retain orthophosphate monoesters and diesters.

316 Inositol phosphate has been found to be adsorbed on amorphous metal oxides and clay  
317 minerals (e.g. montmorillonite, illite, and kaolinite) (Barka and Anderson, 1962; Celi  
318 et al., 1999).

319 The normalized XANES spectra of WECs in sample A and B with soil profile are shown  
320 in Table 4 and Fig. 4. Aluminum phosphate (Al-P), iron phosphate dihydrate (Fe-P),  
321 hydroxyapatite (HAP), and inositol hexakisphosphate (IHP) were detected in WECs for  
322 all the samples. The XANES results of WECs showed that the proportions of Al-P, Fe-  
323 P, and IHP of WECs decreased but HAP increased with soil depth. As a typical alkaline  
324 soil containing carbonate concretions, the Vertisol facilitated the formation of Ca-P  
325 minerals, thus causing low P availability for crops (Westermann, 1992; Iyamuremye et  
326 al., 1996; Li et al., 2011). The decrease of soil pH in the surface soils accelerated the  
327 dissolution of Ca mineral phases, the release of associated colloidal P, and the  
328 transformation of Ca-P to Al-P and Fe-P, thus increasing the concentrations and  
329 proportions of higher activity inorganic P fractions (Zhao *et al.*, 2019). The dominance  
330 of hydroxyapatite in the subsoils might point to the relevance of leaching of dissolved  
331 ortho-P from topsoils into subsoils with subsequent precipitation of apatite due to the  
332 increase in pH and Ca<sup>2+</sup> concentrations. On the other hand, the dominance of  
333 hydroxyapatite in the WEC of subsoils may also reflect the presence of apatite in the  
334 unweathered parent material of soil formation. The proportions of IHP decreased with  
335 soil depth but IHP still existed as certain amounts in the subsoil, which was further  
336 confirmed by the results of NMR.

337 In principle, there are four processes that could lead to the presence of organically-  
338 bound colloidal P in subsoils: i) the mobilization and subsequent transport of colloids  
339 from topsoils into subsoils, ii) the leaching of dissolved organic P and their subsequent  
340 sorption to surfaces of WDC in subsoils, iii) direct input of organic P into subsoils due  
341 to root exudation or root decay, and their subsequent sorption to surfaces of WDC in  
342 subsoils, and iv) the leaching of P from subsoils and its subsequent assimilation by  
343 microorganisms colonizing mineral surfaces in subsoils. As discussed above, the first  
344 process was probably dominated because more negative values of zeta potential at

345 surface soil colloids are beneficial for the transport of colloids from topsoil to subsoil  
346 by increasing repulsion force with soil clay particles. Additionally, the presence of  
347 abundant swelling-shrinkage clays increases preferential flow which also facilitates the  
348 downward transport of colloids. The process ii) and iii) were also possible but their  
349 contributions were not clear. The process iv) seems not important because the OC/OP  
350 in all soil samples ranged from 4.9 to 49.6, which did not support the immobilization  
351 of P. Additionally, microbial phosphorus is mainly phosphate diesters (Turner et al.,  
352 2005). The relatively small fraction of orthophosphate diester indicated by  $^{31}\text{P}$ -NMR  
353 measurements in WEC of subsoil suggested that the process iv) was probably not the  
354 most relevant process explaining the accumulation of WDC containing organically-  
355 bound P in subsoils.

#### 356 **4. CONCLUSIONS**

357 In this study, the distribution of WECs with soil profiles (0-120 cm) was investigated  
358 in a Vertisol under high intensity agricultural management. The P-rich WECs (0.35-2  
359  $\mu\text{m}$ ) with the proportion of 5.3-8.3% of bulk soil was dispersed and transported from  
360 surface soils to subsoils. The TP concentration in the WECs was as high as 1150-1300  
361  $\text{mg kg}^{-1}$ . It is noteworthy that OP associated with WECs were found in subsurface soils,  
362 indicating the colloidal P was transported from surface soils to subsoils, resulting in the  
363 distribution of P-rich WECs throughout the entire soil profile. Soil colloids with a  
364 greater negative charge may be repulsed by negatively charged soil mineral surfaces,  
365 leading to the transportation of these colloidal P to the subsoil (Ilg et al., 2008). The soil  
366 acidification could facilitate deterioration of Ca-stabilized aggregates and accelerate the  
367 release of associated colloidal P. This process subsequently shifted the composition  
368 from Ca dominated colloids to Fe/Al oxides. The existence of swelling-shrinkage  
369 minerals such as montmorillonite and nontronite promoted preferential flow and the  
370 transport of colloidal P. The sum of colloidal P  $< 2 \mu\text{m}$  was 6-37 folds of DRP,  
371 suggesting that colloidal P contributed to P transport significantly in the whole soil  
372 profile. Thus, it is crucial to take into account the impact of colloidal P when predicting  
373 P loss from surface to subsurface flow in Vertisol.

374

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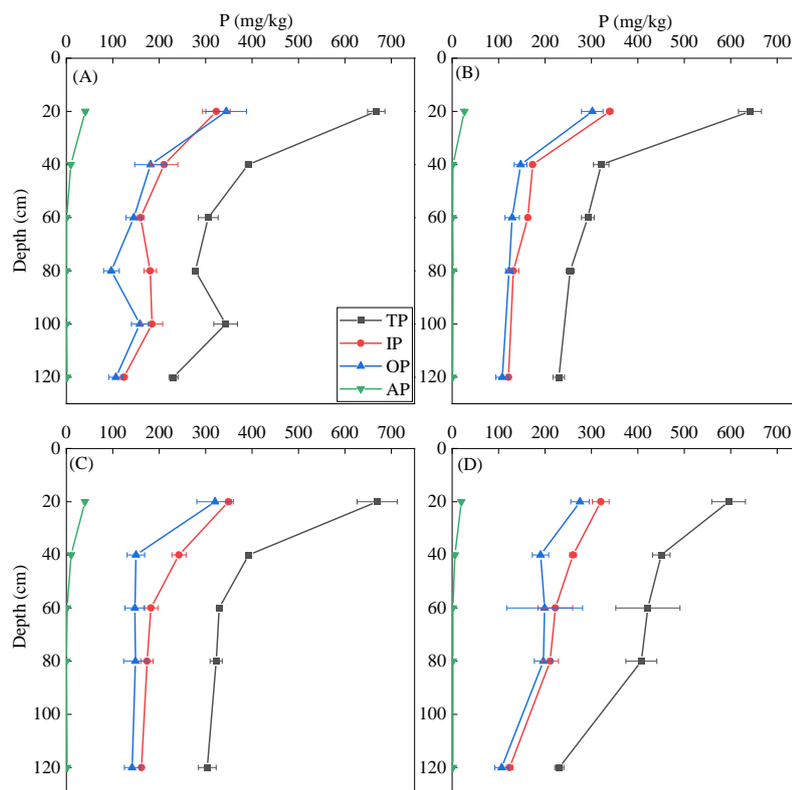
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590 Fig. 1. The concentrations of available phosphorus (AP), total phosphorus (TP), inorganic phosphorus (IP), and  
 591 organic phosphorus (OP) in soil profiles of sample A, B, C, and D.

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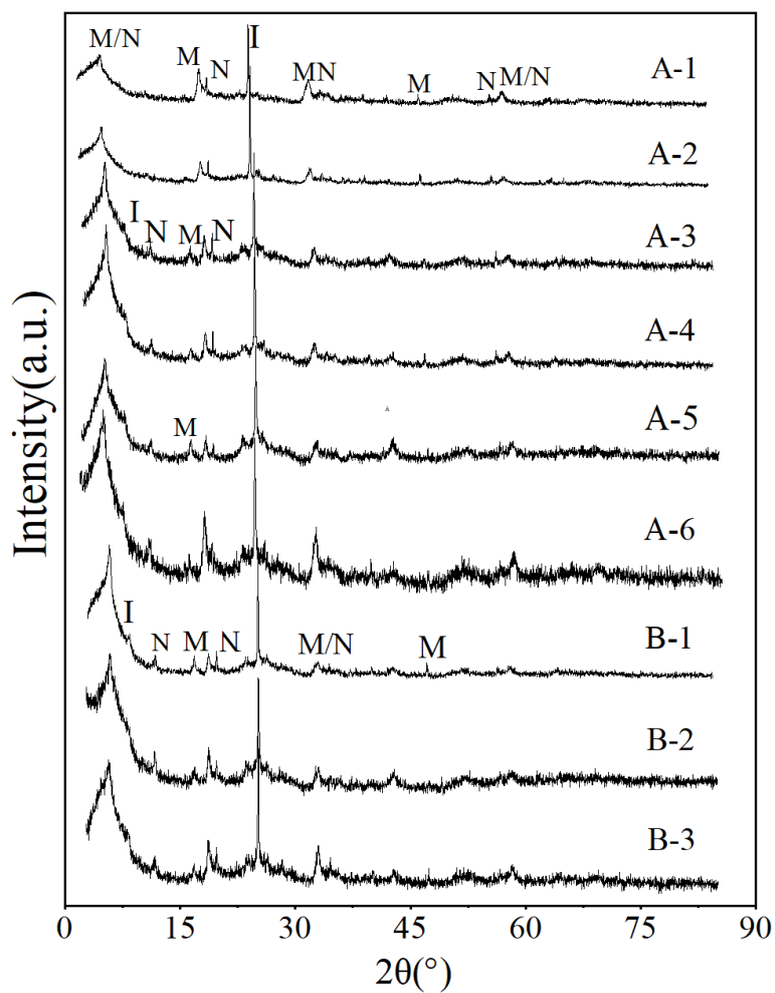
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607 Fig. 2. XRD patterns of water-extractable colloid (WEC) fractions for soil sample A and B with different depths

608 (N: nontronite; M: montmorillonite; I: illite).

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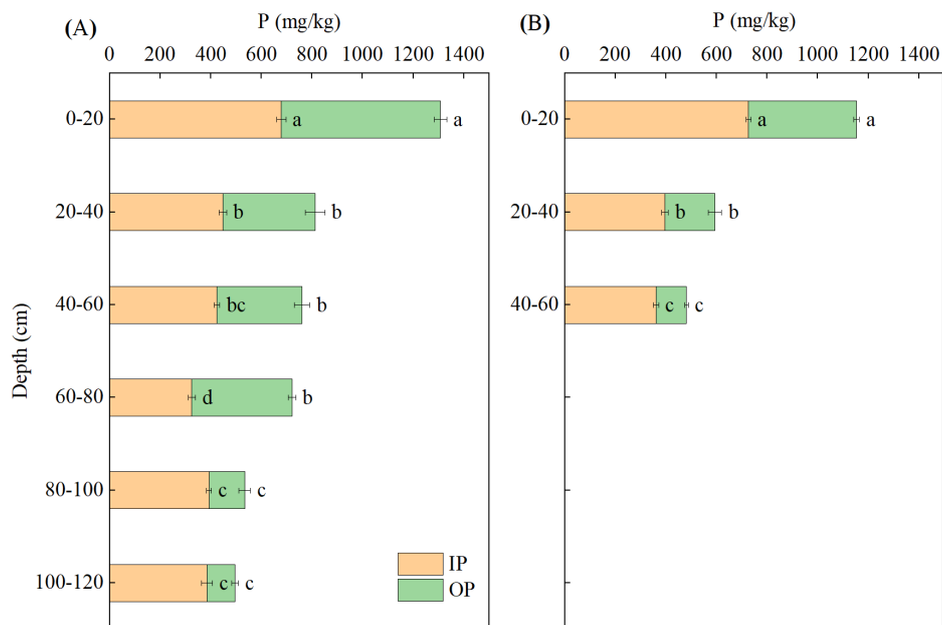
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Fig.3 Inorganic (IP) and organic P (OP) concentrations (mg P/kg colloids) for water-extractable colloid (WEC)

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fractions of sample A and B with different depths.

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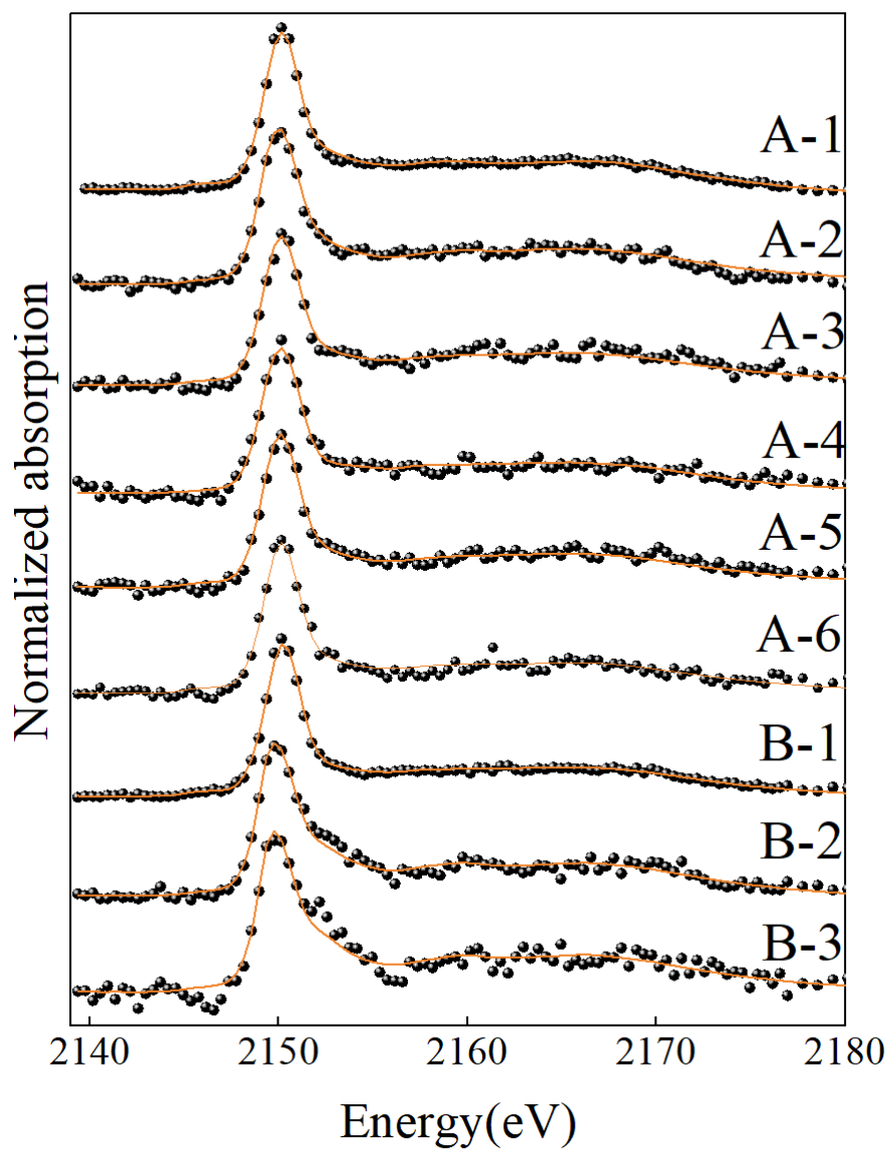
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642 Fig. 4 presents the results of the linear combination fitting (LCF) of the P K- edge XANES spectra for soil colloids

643 of samples A and B. The raw data is represented by points and the fitting results are shown by solid lines. The LCF

644 fitting results can be found in Table 4.

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Table 1 Physicochemical characteristics of bulk soils.

Samp le	Depth(cm)	pH	%TC	%TN	OC/OP	ACP ( $\mu\text{g}/(\text{g}\cdot\text{h})$ )	ALP ( $\mu\text{g}/(\text{g}\cdot\text{h})$ )	OA-P (mg/kg)	DCB-P (mg/kg)	DPS (%)
A	0-20	5.13±0.04e	0.91±0.00a	0.10±0.00a	40.9±3.4ab	647.8±13.5a	465.5±10.2a	142.9±0.8a	178.3±12.3a	4.6±0.0c
	20-40	6.43±0.10d	0.54±0.01d	0.07±0.01b	26.3±2.4b	280.5±4.1b	305.3±14.1d	121.4±5.5b	121.4±5.5bc	4.4±0.0c
	40-60	7.23±0.02c	0.52±0.00e	0.06±0.01bc	30.2±15.8ab	213.3±6.7c	358.6±4.9c	137.6±6.5b	137.6±6.2b	6.8±0.0b
	60-80	7.52±0.02b	0.67±0.01c	0.06±0.01bc	33.9±3.8ab	127.8±9.8d	390.3±8.1b	31.9±3.5c	98.8±22.1c	6.3±0.0b
	80-100	7.52±0.02b	0.75±0.01b	0.07±0.00b	49.6±5.3a	122.6±11.7d	414.0±11.1b	67.0±6.2b	103.1±16.7bc	10.0±0.0a
	100-120	7.70±0.02a	0.50±0.01f	0.06±0.01c	43.9±6.7ab	96.2±8.2c	346.4±16.6c	75.8±15.3b	100.6±5.5c	7.2±0.0b
B	0-20	4.88±0.07c	1.17±0.01a	0.14±0.00a	40.2±4.6a	997.2±14.6a	701.4±14.7a	141.70±9.5a	185.9±2.8a	6.5±0.0c
	20-40	6.28±0.02d	0.66±0.01b	0.08±0.00b	37.2±5.1a	413.0±4.7b	514.7±5.8b	84.7±1.3b	121.4±7.8b	17.4±0.0a
	40-60	7.41±0.01c	0.48±0.00d	0.06±0.00c	26.2±3.9b	186.6±3.8c	509.3±17.9c	59.7±12.7c	66.6±11.4d	16.1±0.0a
	60-80	8.03±0.01b	0.35±0.01e	0.05±0.00d	10.4±1.8c	147.2±14.3d	392.5±16.6d	73.7±3.2bc	99.8±2.8c	11.0±0.0b
	80-120	8.37±0.01a	0.52±0.01c	0.04±0.00e	17.5±1.4c	100.1±9.3e	326.2±14.3e	60.7±1.9c	79.1±0.9d	5.2±0.0c
	C	0-20	5.01±0.02c	0.94±0.00a	0.11±0.00a	19.7±2.2a	1177.6±31.8a	510.6±15.2b	161.7±5.6a	182.3±6.0a
20-40		6.23±0.03d	0.54±0.00b	0.07±0.00b	15.9±3.0a	332.4±12.8b	377.3±15.3d	33.7±0.7b	70.4±3.90b	15.7±0.0b
40-60		7.32±0.04c	0.48±0.00c	0.06±0.00c	9.3±1.0b	190.0±4.7c	578.1±7.6a	29.7±13.4b	45.6±8.8c	19.2±0.0a
60-80		7.60±0.03b	0.39±0.01d	0.04±0.01d	5.4±0.9b	176.6±3.2c	489.5±4.2b	41.7±6.2b	43.8±5.3c	14.3±0.0b
80-120		7.74±0.03a	0.27±0.00e	0.04±0.00d	4.9±1.0b	126.8±17.9d	448.1±7.4c	48.7±4.81b	50.1±4.3c	4.8±0.0c
D		0-20	5.17±0.04e	0.88±0.01a	0.10±0.01a	28.7±2.3a	749.6±9.6a	502.1±12.9bc	126.7±6.2a	129.2±6.30a
	20-40	6.97±0.06d	0.57±0.02b	0.07±0.00b	33.1±3.1a	435.4±3.2b	557.2±14.6a	33.7±2.7b	51.4±2.07c	19.4±0.0ab
	40-60	7.46±0.03c	0.50±0.00c	0.06±0.00c	32.7±3.8a	136.7±5.8c	526.0±16.3b	34.7±7.0b	65.6±2.28b	21.0±0.0a
	60-80	7.78±0.03b	0.36±0.01d	0.04±0.01cd	21.1±1.0b	175.9±6.5d	497.1±1.7c	42.7±5.7b	49.8±49.1c	13.7±0.0c
	80-120	8.04±0.01a	0.25±0.00e	0.03±0.01d	13.1±1.7c	97.9±2.3e	349.4±10.3d	37.7±1.7b	40.1±1.93c	17.2±0.0b

656 TC: total carbon, TN: total nitrogen, ACP: acid phosphatase activity, ALP: alkaline phosphatase  
657 activity, OA-P: oxalic acid extractable P, DCB-P: dithionite–citrate–bicarbonate extracted P. DPS:  
658 degrees of phosphorus saturation ( $\text{DPS} = [\text{OA-P}] / 0.5([\text{OA-Fe}] + [\text{OA-Al}])$ ). OA-Fe: oxalic acid  
659 extractable Fe. OA-Al: oxalic acid extractable Al.

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676 Table 2 Soil fractionation, physicochemical characteristics and P levels of water extracted colloids.

Sample	Depth (cm)	Soil texture	Soil sand fractions (>20 $\mu\text{m}$ , %)	Soil silt fractions (2-20 $\mu\text{m}$ , %)	Colloidal fraction (0.35-2 $\mu\text{m}$ , %)	Zeta potential (mV)	Colloidal P (0.35-2 $\mu\text{m}$ ) (mg/kg soil)	DTP (<0.35 $\mu\text{m}$ , mg/kg soil)	DRP (<0.35 $\mu\text{m}$ , mg/kg soil)	CP/TP (%)
A	0-20	sandy loam	57.0 $\pm$ 4.0ab	34.8 $\pm$ 3.8ab	6.2 $\pm$ 0.3c	-19.1 $\pm$ 2.7	80.7 $\pm$ 3.6a	22.1 $\pm$ 1.0a	10.0 $\pm$ 1.3a	15.6 $\pm$ 1.0ab
	20-40	sandy loam	55.7 $\pm$ 1.4ab	34.2 $\pm$ 1.2ab	6.3 $\pm$ 0.3bc	-18.5 $\pm$ 1.1	51.0 $\pm$ 1.7b	15.3 $\pm$ 0.9b	7.4 $\pm$ 0.4b	13.1 $\pm$ 0.7b
	40-60	sandy loam	57.3 $\pm$ 4.5 ab	33.6 $\pm$ 4.4ab	6.9 $\pm$ 0.2bc	-17.9 $\pm$ 0.6	52.4 $\pm$ 1.9b	12.3 $\pm$ 1.2c	4.6 $\pm$ 1.1c	14.4 $\pm$ 1.5b
	60-80	sandy loam	56.8 $\pm$ 2.7ab	33.0 $\pm$ 2.5ab	7.3 $\pm$ 0.6ab	-17.5 $\pm$ 2.9	52.9 $\pm$ 4.4b	6.7 $\pm$ 1.4d	2.6 $\pm$ 0.5cd	14.1 $\pm$ 1.9b
	80-100	sandy loam	59.7 $\pm$ 2.9 a	30.4 $\pm$ 3.3b	7.0 $\pm$ 0.5bc	-15.7 $\pm$ 2.7	37.3 $\pm$ 1.3c	6.0 $\pm$ 0.3d	2.0 $\pm$ 0.7d	12.0 $\pm$ 0.8b
	100-120	loam	50.3 $\pm$ 1.6 b	38.8 $\pm$ 0.9a	8.3 $\pm$ 0.5a	-15.8 $\pm$ 0.8	41.2 $\pm$ 1.4c	4.1 $\pm$ 0.8d	1.2 $\pm$ 0.3d	19.2 $\pm$ 1.7b
B	0-20	sandy loam	56.3 $\pm$ 3.4a	35.9 $\pm$ 3.6a	5.3 $\pm$ 0.3b	-22.0 $\pm$ 1.7	61.3 $\pm$ 2.7a	19.5 $\pm$ 1.1a	10.0 $\pm$ 1.3a	10.6 $\pm$ 1.1a
	20-40	sandy loam	56.5 $\pm$ 3.2a	34.8 $\pm$ 4.1a	5.3 $\pm$ 0.4b	-18.9 $\pm$ 1.0	31.6 $\pm$ 1.5b	14.4 $\pm$ 1.2b	6.3 $\pm$ 0.4b	10.1 $\pm$ 0.8a
	40-60	sandy loam	57.0 $\pm$ 1.9a	33.1 $\pm$ 1.2a	6.5 $\pm$ 0.3a	-9.0 $\pm$ 0.6	31.1 $\pm$ 1.1b	11.2 $\pm$ 0.9c	4.6 $\pm$ 0.3b	11.4 $\pm$ 0.7a

677 DTP: total P concentrations of dissolved fractions (<0.35  $\mu\text{m}$ ); DRP: reactive P concentrations of  
678 dissolved fractions (<0.35  $\mu\text{m}$ ); CP: colloidal P; TP: total P.

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698 Table 3. Concentrations of inorganic and organic P extracted by NaOH-Na<sub>2</sub>EDTA (mg /kg-1) in the water-extractable  
 699 colloid (WEC) fractions of sample A and B by solution <sup>31</sup>P-NMR.

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Sample	Depth(cm)	Inorganic P (mg/kg)		Organic P (mg/kg)					
		Orth	Pyro	Orthophosphate Monoesters				Orthophosphate Diesters	
				Monoesters*	Myo-IHP	Scyllo-IHP	Other Monoesters	Diesters*	Glyc+nuc l
A-1	0-20	202.4	10.1	50.6	8.1	3.5	39	89.1	81.0
A-2	20-40	46.8	5.1	62.0	1.4	0.0	60.6	15.4	1.8
A-3	40-60	14.1	0.14	42.2	0.6	2.0	39.6	11.4	0
A-4	60-80	14.0	0.84	38.5	0.4	0.00	38.1	0.0	0
A-5	80-100	15.3	5.5	39.6	9.2	0.00	30.4	8.1	3.5
A-6	100-120	12.5	0.0	36.3	3.8	0.00	32.5	7.5	3.8
B-1	0-20	427.5	8.6	128.3	4.3	17.6	106.4	21.4	8.5
B-2	20-40	121.0	7.3	128.1	3.7	0.00	124.4	12.2	7.3
B-3	40-60	57.5	10.4	52.9	3.5	0.00	49.4	1.7	1.7

701 \* after diesters means calculation by including diester degradation products (i.e. Glyc+nucl:  $\alpha/\beta$ - glycerophosphate,  
 702 and mononucleotides) with orthophosphate diesters (Diesters) rather than orthophosphate monoesters (Monoesters).

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Table 4 Phosphorus K-edge XANES fitting results showing the relative percent of each P species in water-extractable colloid (WEC) fractions (%) of sample A and B.

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Sample	Depth (cm)	Al-P (%)	Fe-P (%)	HAP (%)	IHP (%)
A-1	0-20	17.0±4.0	33.0±3.0a	10.0±2.0c	32.0±6.0a
A-2	20-40	13.0±2.0	30.0±5.0a	26.0±4.0bc	30.0±0.8ab
A-3	40-60	14.0±7.0	28.0±0.8a	32.0±7.0abc	26.0±5.0abc
A-4	60-80	12.0±1.0	29.0±3.0a	35.0±3.0abc	22.0±7.0abc
A-5	80-100	11.0±3.0	24.0±3.0ab	45.0±2.0ab	10.0±0.8c
A-6	100-120	9.0±7.0	16.0±3.0b	53.0±7.0a	13.0±5.0bc
B-1	0-20	25.0±7.0a	36.0±2.0a	13.0±3.0b	17.0±6.0
B-2	20-40	11.0±0.9b	19.0±6.0a	32.0±4.0ab	9.0±4.0
B-3	40-60	3.0±0.6b	20.0±5.0b	60.0±10.0a	7.0±4.0

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Al-P: aluminum phosphate ( $\text{AlPO}_4$ ), Fe-P: iron phosphate dihydrate ( $\text{FePO}_4 \cdot 2\text{H}_2\text{O}$ ), HAP: hydroxyapatite ( $\text{Ca}_5(\text{PO}_4)_3\text{OH}$ ), and IHP: inositol hexakisphosphate. Values in each column followed by the different lowercase letters indicate significant differences ( $P < 0.05$ ).

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