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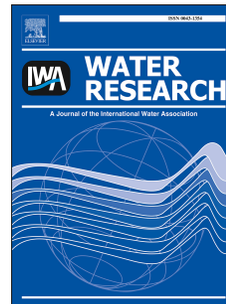
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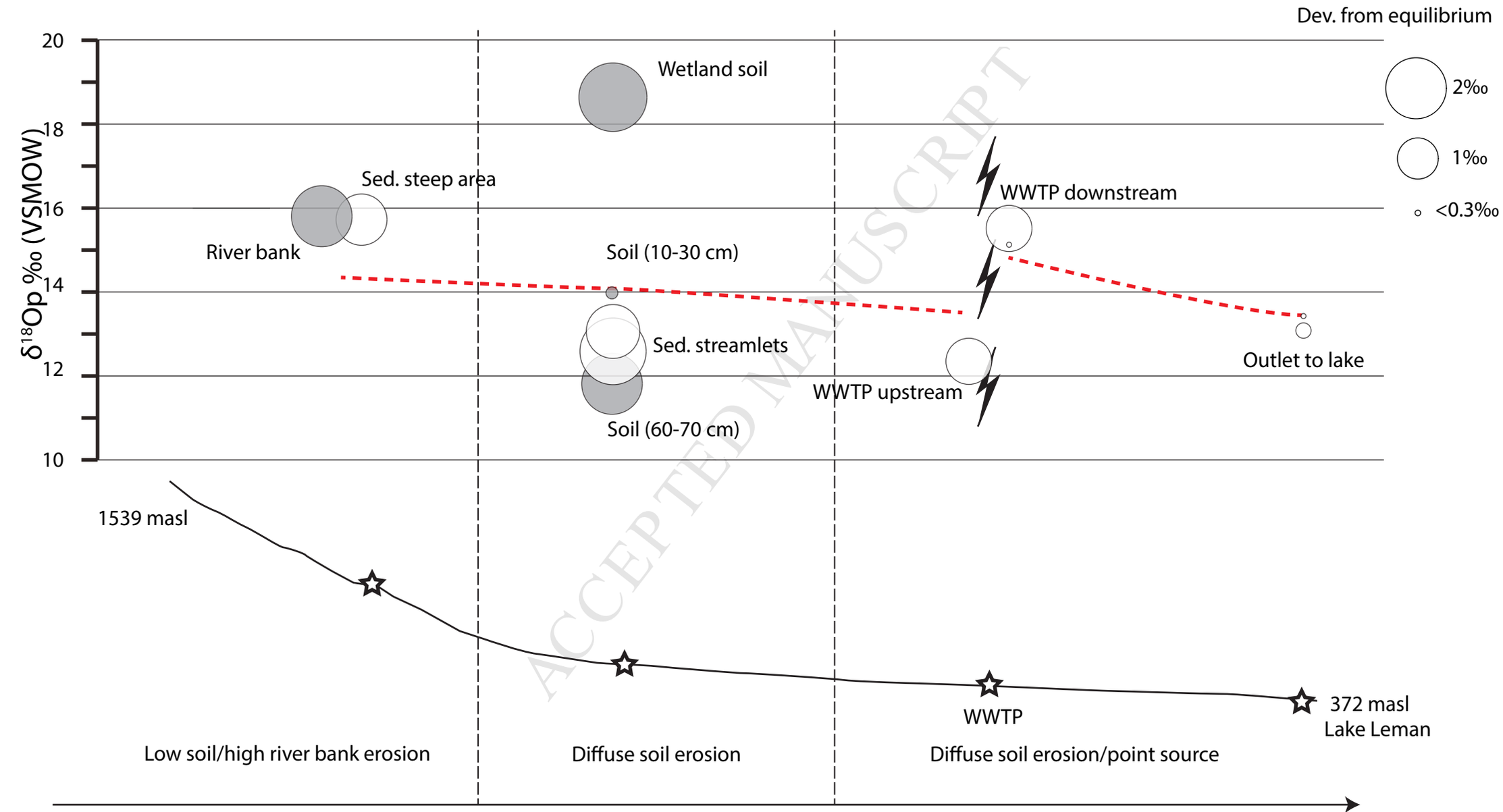
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1 **Tracing the sources and cycling of phosphorus in river sediments using oxygen isotopes:**
2 **methodological adaptations and first results from a case study in France**

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11

12 **Abstract**

13 An essential aspect of eutrophication studies is to trace the ultimate origin of phosphate ions (P-
14 PO₄) associated with the solid phase of river sediments, as certain processes can make these
15 ions available for algae. However, this is not a straightforward task because of the diversity of
16 allochthonous and autochthonous sources that can supply P-PO₄ to river sediments as well as
17 the existence of in-stream processes that can change the speciation of these inputs and obscure
18 the original sources. Here, we present the results of a study designed to explore the potentials,
19 limitations and conditions for the use of the oxygen isotope composition of phosphate ($\delta^{18}\text{O}_\text{p}$)
20 extracted from river sediments for this type of tracing. We first tested if the method commonly
21 applied to soils to purify P-PO₄ and to measure their $\delta^{18}\text{O}_\text{p}$ concentrations could be adapted to
22 sediments. We then applied this method to a set of sediments collected in a river along a
23 gradient of anthropogenic pressure and compared their isotopic signatures with those from

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24 samples that are representative of the potential P-PO₄ inputs to the river system (soils and
25 riverbank material).
26 The results showed that following some adaptations, the purification method could be
27 successfully transposed to river sediments with a high level of P-PO₄ purification (> 97%) and
28 high $\delta^{18}\text{O}_p$ measurement repeatability and accuracy (< 0.4‰). The values for the potential
29 allochthonous sources varied from 11.8 to 18.3‰, while the $\delta^{18}\text{O}_p$ value for the river sediments
30 ranged from 12.2 to 15.8‰. Moreover, a sharp increase (> 3‰) in the sediment $\delta^{18}\text{O}_p$ value
31 immediately downstream from the discharge point revealed the strong impact of municipal
32 wastewater. The calculation of the theoretical equilibrium $\delta^{18}\text{O}_p$ values using the river water
33 temperature and $\delta^{18}\text{O}_w$ showed that the downstream sediments were in equilibrium, which was
34 not the case for the upstream sediments. This difference could be related to the contrast
35 between the short residence time of the transfer system in the catchment head, which can
36 preserve the isotopic variability of the source materials, and the longer residence times and
37 higher P bioavailability in the lower catchment, possibly fostering the recycling of P-PO₄ by
38 the biota and the equilibration of the oxygen isotope signature in P-PO₄. These results
39 demonstrate the potential of the isotopic approach to assess the sources and in-stream turnover
40 of sedimentary P in river systems.

41
42 **KEYWORDS:** Phosphorus transfer, Agriculture, In-stream processes, Eutrophication,
43 Phosphate oxygen isotopes

44

45 **1. Introduction**

46

47 Phosphorus (P) associated with the solid phase and designated as “particulate-P” often accounts
48 for the major part of phosphorus transferred from terrestrial to aquatic ecosystems (Kleinman et

49 al., 2011; Kronvang et al., 2007; Sharpley et al., 1999). The impact of particulate-P on the
50 trophic state and eutrophication of the receiving aquatic ecosystems, especially lentic
51 ecosystems, depends on the size of its potentially bioavailable fraction. The amount of P that
52 occurs as inorganic phosphate ions (P-PO₄ pools), either in solution (dissolved P-PO₄), sorbed
53 onto, or precipitated within stream and river sediments, is thought to represent the most readily
54 bioavailable P pool (Moorlegghem et al., 2013). Therefore, an essential aspect of studies
55 dedicated to the restoration of eutrophicated aquatic ecosystems is to quantify P-PO₄ pools in
56 river water and sediments and to trace the ultimate origin of these pools (Kleinman et al., 2011;
57 Mellander et al., 2015).

58 However, this quantification is not a straightforward task because of the diversity of the
59 allochthonous and autochthonous sources that may supply P-PO₄ to river sediments. Several
60 allochthonous compartments in the catchment can supply particulate-P to river sediments,
61 including pristine and weathered basement rocks, forested or agricultural soils, riverbank
62 sediments, etc. For example, the autochthonous addition of P-PO₄ to the sediments can occur
63 through sorption onto the sediments of dissolved P-PO₄ ions supplied by sewage effluents or
64 the in-stream mineralization of sewage organic matter (Poulenard et al., 2008). The in-stream
65 retention of dissolved P-PO₄ by the sediments primarily occurs during low-flow periods when
66 the retention time of the sediment is long enough to allow the necessary physical-chemical
67 reactions to develop (Dorioz et al., 1998a, 1989; Ensign et al., 2006; Pistocchi et al., 2012) and
68 is particularly intense immediately downstream of the point source pollution inputs.

69 Autochthonous P-PO₄ pools can also come from the recycling of allochthonous particulate-P by
70 aquatic flora or by microorganisms in the sediment. The biological uptake by biofilm is another
71 important seasonal process that is involved in the retention of P-PO₄ ions by the sediment
72 (Wang et al., 1999). These in-stream recycling and retention processes, which help the P to turn
73 over within the river ecosystem, are expected to be triggered by the bioavailability of

74 allochthonous P and to regulate the internal productivity of the river system. Their
75 quantification can be used to assess the efficiency of a river system to recycle and transform
76 any particulate P into more bioavailable P-PO₄, but an efficient recycling can eventually
77 obscure the ultimate origin of the P-PO₄ exported at the outlet.

78 It is often impossible to quantify and trace the different sources of particulate-P that are
79 potentially involved in eutrophication using only the variability in P speciation as determined
80 by classical chemical extraction methods (Burrus et al., 1989; Pacini and Gächter, 1999;
81 Poulenard et al., 2009; Walling et al., 2008). As a result, there a need to develop tools or/and
82 strategies that can be reliably used to trace this important P compartment, particularly its
83 dominant inorganic P-PO₄ fraction, and to understand the extent and rate at which particulate-P
84 is biologically recycled and modified within streams and rivers.

85 Oxygen isotopes could provide one such possible tool. The oxygen isotopic composition of
86 phosphate ($\delta^{18}\text{Op}$) has already been successfully used to trace P sources and study the P-PO₄
87 cycle in soils, freshwater and marine sediments (Davies et al., 2014; Goldhammer et al., 2011;
88 Goody et al., 2016; Gruau et al., 2005; Melby et al., 2013; Tamburini et al., 2014). This
89 approach is based on: i) the existence of measurable differences in the abundance ratio for ¹⁸O
90 and ¹⁶O atoms in the phosphate group of potential P-PO₄ sources (e.g. primary P-rich minerals,
91 chemical fertilizers, chemical detergents, phosphate from wastewater, etc.); and ii) the fact that
92 biologically mediated processes have the possibility to modify $\delta^{18}\text{Op}$ in a predictable way
93 (Blake et al., 2005). The first property allows the oxygen isotope composition of P-PO₄ to be
94 potentially used as a source tracer in environmental systems, whereas the second property
95 allows it to be used to quantify the rate of biological recycling of P-PO₄. Two enzymatic
96 processes are mainly responsible for changes in the $\delta^{18}\text{Op}$ value. One is the isotope
97 equilibration between oxygen in P-PO₄ and oxygen in ambient water. This process, which is
98 caused by reversible phosphorylation/dephosphorylation reactions that are catalyzed by the

99 inorganic intracellular pyrophosphatase enzyme (Blake et al., 2005), is temperature dependent
100 and has been described by an empirical equation (Longinelli and Nuti, 1973) that was recently
101 revised by Chang and Blake (2015). The second is the kinetic isotope fractionation produced by
102 the enzymatic hydrolysis of phosphoesters (Liang and Blake, 2006; von Sperber et al., 2015,
103 2014), the magnitude of which depends on the type of enzyme involved (e.g. acid or alkaline
104 phosphatases) and the final effect of which on the isotopic signature of the recycled P-PO₄ also
105 depends on the $\delta^{18}\text{O}_p$ value for the original substrate, i.e. inheritance effect (Liang and Blake,
106 2006). Small kinetic isotope effects are associated with the initial stages of the abiotic reaction,
107 such as the sorption of P-PO₄ onto the solid phases (Jaisi et al., 2010; Melby et al., 2013b) or
108 biological uptake (Blake et al., 2005).

109 In freshwater ecosystem studies, the use of $\delta^{18}\text{O}_p$ is at an early stage (Davies et al., 2014). For
110 river sediments, the lack of a standardized method to extract, purify and analyze $\delta^{18}\text{O}_p$ is a
111 hinderance. In this study, we first adapted the method that is commonly used for soils to
112 sediments and validated the capability of this method to provide precise and accurate results.
113 We then applied this method to riverbed sediments from a river in the French Alps with the
114 following objectives: i) to characterize the variability of the $\delta^{18}\text{O}_p$ signatures in the riverbed
115 sediments and the potential external sediment sources (soils, riverbank materials) in an
116 exemplar river system; ii) to explore the potential, limitations and conditions of using the
117 isotopic tool to assess the respective role of external inputs and internal recycling processes to
118 build up the P-PO₄ budget of the riverbed sediments in this river.

119

120 **2. Materials and methods**

121

122 **2.1 Site description and sampling**

123 The Redon River is a small alpine torrential tributary (i.e. a shallow high-gradient, high-energy
124 stream with an annual average discharge of $0.5 \text{ m}^3/\text{s}$ at the outlet) of Lake Geneva (France).
125 The sediment dynamic of this river has been already intensively studied (Al-Charideh, 1988;
126 Dorioz et al., 1998b; Feuillade and Dorioz, 1992; Hamid et al., 1989; Poulenard et al., 2008)
127 making it a particularly appropriate site for the present exploratory study. The total catchment
128 surface area is 3,300 ha with a land use that consists mainly of forests (1,100 ha) in its upland
129 part and grasslands (1,200 ha) and cultivated farmlands (300 ha) in its lowland part. Several
130 wetlands (20 ha) are present, while urbanized areas are concentrated near the outlet, close to the
131 shore of Lake Geneva. The Redon River is impacted by both point and diffuse sources of P.
132 The main point source of P is located in the lower part of the river and consists of inputs of
133 predominantly dissolved P from a wastewater treatment plant (WWTP), whereas the main
134 diffuse sources occur in the upper part of the catchment, consisting mainly of particulate-P
135 from agricultural and forested lands (Fig. 1).
136 The soils are mainly Cambisols, formed from the weathering products of quaternary moraine
137 material consisting of a large range of mainly calcareous sedimentary, metamorphic and
138 igneous rocks from the Rhône glacier (Nicoud et al., 1993). The soil mineralogy is comprised
139 of a mixture of minerals inherited from the weathering of this heterogeneous parent material
140 and their secondary weathering products. Previous studies (Dorioz et al., 1998a, 1998b) have
141 shown that the Redon River sediments are CaCO_3 -rich (10 to 20% CaCO_3 , $\text{pH} > 7$). Their fine
142 ($< 200 \mu$) fraction is dominated by loamy and sandy materials, with an organic matter content
143 between 2 to 4.5%. The clay fractions (7-11%) consist mostly of soil-derived illite, biotite and
144 chlorite (Poulenard et al., 2008). The water in the Redon River is bicarbonated-calcic.

145

146 [Figure 1]

147

148 2.2 Sampling strategy and sample preparation

149 Details regarding the sampling sites and sampling conditions are given in Tables 1 and 2. Three
150 types of samples were collected: i) eight riverbed sediments collected at six locations (A, B, C,
151 D, E and F in Fig. 1) along a downstream gradient of land use that changes from forested to
152 mixed agricultural/urban areas, thus representing a gradient of increasing anthropogenic
153 pressure; this includes two samples collected in streamlets draining a small agricultural sub-
154 catchment (locations A and B in Fig. 1) that is representative of the type of agriculture
155 practiced in the Redon River catchment (Dorioz and Ferhi, 1994); ii) four samples
156 representative of the potential sources of river sediments, including cultivated topsoil and deep
157 horizon, riverbank material, and soil from a 2 ha wetland, hydraulically connected to the river;
158 and iii) two samples from sediments from Lake Geneva representing an average reference for
159 the local superficial lacustrine sediments (collected in 2012 and provided by the EDYTEM
160 laboratory).

161 Given that it was difficult to access the river bed, it was not possible to visit all six riverbed
162 sediment sampling locations at the same time. Three locations were first sampled at the end of
163 March 2012 (B, C and F), and five (A, B, D, E and F) were sampled (again) at the beginning of
164 September 2012. Both periods were low-flow periods, where the September period corresponds
165 to the lowest weekly discharge period of the year. Samples were taken in areas of preferential
166 accumulation of fine sediments. At each location, single samples of the entire sediment matrix
167 were taken manually at 3-5 different points and then mixed to obtain an average sample (0.5-1
168 kg dry weight). Simultaneously, the electric conductivity (EC) was measured and a water
169 sample was collected to determine the SRP (soluble reactive phosphorus) concentration and
170 water oxygen isotope composition ($\delta^{18}\text{O}_w$). Hourly water temperatures were obtained from
171 automatic recorders at the outlet. The riverbank sample was collected in the upstream, forested
172 part of the catchment (see Fig. 1). The cultivated soil samples were collected in the same small

173 sub-catchment used to sample the streamlet sediments (Fig. 1), as this sub-catchment is known
174 to represent zones of maximum soil erosion risk in the Redon catchment (Dorioz and Ferhi,
175 1994). In addition to taking a topsoil sample, a deeper soil sample was taken at a depth of
176 approximately -70 cm to account for the possible erosion of deep soil materials along ditches
177 or/and soil embankments. These two samples were obtained by pooling five subsamples (200-
178 300 g) collected in different locations within a single field or its embankment. The pooled
179 samples were then air-dried and sieved at 200 μm before analysis. No river water samples were
180 collected for the analysis of the $\delta^{18}\text{O}_\text{p}$ values for the dissolved P- PO_4 , although these could
181 have been useful in the interpretation of the riverbed sediment data. The decision not to sample
182 the river water was due to the very low concentrations of dissolved P- PO_4 recorded in this river
183 (Table 1), which makes it extremely difficult to analyze the $\delta^{18}\text{O}_\text{p}$ in the current state of the
184 analytical protocols.

185
186 [Table 1]

187
188 [Table 2]

189

190 **2.2 Sediment and soil P chemical analyses**

191

192 The total phosphorus (Pt) and Olsen P (P_{Olsen}) concentrations were determined in all of the
193 samples. P_{Olsen} (Olsen et al., 1954), which is considered as being representative of the
194 bioavailable fraction of P in soils, has been shown to correlate linearly with P extracted from
195 the sediment in bio-assays with algae (Jordan-Meille, 1998). The Pt concentrations were
196 measured after a hot sodium hydroxide digestion. The total inorganic and organic P in the
197 samples were determined according to Saunders and Williams (1955) as described by Fares et

198 al. (1974). The spectrophotometric determination of P-PO₄ in all of the extracts/digestates was
199 carried out using the molybdenum blue method (Murphy and Riley, 1962) at 882.0 nm on a
200 Cary 50 UV-Vis spectrometer (Agilent).

201

202 **2.3 Isotopic analyses and data treatment**

203 *2.3.1 Extraction and purification protocol*

204 We used the extraction and purification protocol initially developed by Tamburini et al. (2010)
205 for soils (hereafter referred as to Tb10), which we adapted for sediments. The Tb10 protocol
206 consists of the following six steps:

207 i) extraction of phosphate from the soil/sediment with 1 M HCl, which in sequential extraction
208 mainly targets the P-PO₄ pool (HCl-P) associated with Ca, but in our case (singular extraction)
209 also P sorbed onto Fe and Al oxides;

210 ii) dissolved organic matter (DOM) removal from the solution using an Amberlite resin,

211 iii) ammonium phospho-molybdate (APM) precipitation and subsequent dissolution,

212 iv) magnesium ammonium phosphate (MAP) precipitation and dissolution,

213 v) cation removal by a cation exchange resin (BioRad, AG50 x8),

214 vi) silver phosphate (Ag₃PO₄) precipitation through the addition of an Ag-ammine solution to
215 the purified phosphate solution.

216 The goal of these steps, after HCl extraction, is to avoid sources of oxygen (i.e. DOM, nitrate,
217 etc.) other than P-PO₄ and other contaminants, e.g. chloride, in the final silver phosphate
218 precipitate.

219 We modified the Tb10 protocol in several ways as follows (Fig. 2 in the supplementary
220 material). First, the samples were sieved at 200 µm instead of 2 mm, since this finer fraction
221 contains the bulk of the bio-geochemically active P in river sediments (Poulenard et al., 2008).
222 The liquid-to-solid ratio in the extraction step was modified to 100:1 instead of 50:1, since the

223 pre-tests showed that, with our highly carbonated sediments, the solution did not reach the
 224 targeted pH (1 ± 0.5). The quantity of sediment was adjusted to have at least 10 μ moles of P for
 225 the purification, corresponding to 10 to 20 g of sieved sediments. The DOM removal step using
 226 an Amberlite resin was not performed as the subsequent steps successfully eliminated the
 227 organic matter (see below). Finally, for samples with a high calcium carbonate content, an
 228 additional step had to be introduced to account for the formation of a calcium molybdate
 229 precipitate during the APM precipitation. To eliminate the precipitate, we repeated the
 230 subsequent MAP precipitation step twice.

231 The 1 M HCl extraction may hydrolyze some of the organic or condensed P (Turner et al.,
 232 2005) and the released Pi may bias the isotopic composition of the inorganic P pool targeted
 233 here. To assess the existence of such a possible artifact, extractions were run in duplicate using
 234 ^{18}O -labeled and unlabeled reagents. An ^{18}O -labelled 1M HCl solution was prepared by adding
 235 approximately 0.05 ml of ^{18}O -enriched water (97‰, Aldrich) to 1 liter of 1 M HCl solution.
 236 Labeled and unlabeled solutions were analyzed for $\delta^{18}\text{O}_w$. If hydrolysis occurred, it should
 237 result in different isotopic compositions depending on whether labeled or unlabeled reagents
 238 were used, i.e. a higher $\delta^{18}\text{O}_p$ for the labeled sample compared to the unlabeled one. In this case,
 239 we calculated the “real” isotopic value by a revised version of the mass balance equation used
 240 by McLaughlin et al. (2006b):

$$241 \quad \delta^{18}\text{O}_p = \frac{(\delta^{18}\text{O}_{pl} \cdot \delta^{18}\text{O}_{wnl} - \delta^{18}\text{O}_{pnl} \cdot \delta^{18}\text{O}_{wl})}{(\delta^{18}\text{O}_{pl} - \delta^{18}\text{O}_{pnl} - \delta^{18}\text{O}_{wl} + \delta^{18}\text{O}_{wnl})} \quad (1)$$

242 where $\delta^{18}\text{O}_{pl}$ and $\delta^{18}\text{O}_{pnl}$ are the isotopic composition of P- PO_4 for the labeled and unlabeled
 243 samples, respectively, and $\delta^{18}\text{O}_{wl}$ and $\delta^{18}\text{O}_{wnl}$ are the isotopic composition of water for the
 244 labeled and unlabeled reagents, respectively.

245 The efficiency and reliability of the modified Tb10 protocol used here were evaluated by
 246 considering the repeatability of the extraction/purification process, the final Ag_3PO_4 yield and
 247 its purity. The purity was assessed by i) dissolving the obtained Ag_3PO_4 and comparing the

248 actual concentration of phosphate in the solution with the theoretical concentration, i.e. the
249 amount expected if all of the solid mass was Ag_3PO_4 , and ii) determining the C and N
250 concentration of the obtained Ag_3PO_4 on the elemental analyzer (Flash 2000, Termo
251 Scientific); these two concentrations are used as indicators of possible contamination by
252 organic matter and/or nitrate. The purity of the obtained Ag_3PO_4 crystals was also assessed
253 through quantification of the oxygen yield during mass spectrometry analyses. This was done
254 by graphically comparing the weight of the Ag_3PO_4 standards and samples introduced into the
255 machine against the peak area of the produced CO (“oxygen yield”), which is proportional to
256 the oxygen content in the sample. If other sources of oxygen or contaminants were present in
257 the sample, the oxygen yield should be different from the standard Ag_3PO_4 . P- PO_4 losses at
258 each step of the purification were also quantified (see Pistocchi et al. 2014 for more details).

259 *2.3.2 Isotopic analyses and calculation of the theoretical equilibrium values*

260 The oxygen isotope composition of the purified Ag_3PO_4 was measured using a thermal
261 conversion elemental analyzer (Variopyro Cube Elementar), coupled to an IsoPrime100
262 isotopic ratio mass spectrometer (IRMS) at the Plant Nutrition Group laboratory (ETH Zurich).
263 Two to three replicates of each sample were weighed (300 to 600 μg) in silver capsules with a
264 small added amount of glassy carbon powder. The temperature of the TC/EA furnace was kept
265 at 1450°C for pyrolysis.
266 During each run, repeated measurements of an internal standard (Acros Organics Ag_3PO_4 >
267 97.5%, $\delta^{18}\text{O}_p = +14.2\text{‰}$) and two international benzoic acid standards (IAEA 601 and IAEA
268 602, $\delta^{18}\text{O} = +23.3\text{‰}$ and $+71.4\text{‰}$, respectively, IAEA, Vienna, Austria) were included and
269 used to calibrate and correct the instrumental drift. The $\delta^{18}\text{O}_p$ values presented in this paper are
270 reported in the standard δ notation with respect to VSMOW (Vienna Standard Mean Oceanic
271 Water), i.e. as ‰ deviation values. Analytical uncertainties, as determined from the replicate
272 analysis of the standards, were estimated to be below 0.3‰.

273 In order to assess the extent to which the sedimentary P-PO₄ pools investigated in this study
274 could be secondary recycled P-PO₄ (i.e. P-PO₄ that has been cycled by the river biota), a
275 theoretical equilibrium $\delta^{18}\text{O}_p$ value – i.e. the oxygen isotopic composition the phosphate should
276 have if it is in equilibrium with the ambient water – was calculated for each sample using the
277 following equation (Chang and Blake, 2015):

$$278 \quad \delta^{18}\text{O}_p = (\delta^{18}\text{O}_w + 10^3) \cdot e^{\frac{\left[\left(14.43 \cdot \frac{10^3}{T}\right) - 26.54\right]}{10^3}} - 10^3 \quad (2)$$

279
280 where $\delta^{18}\text{O}_p$ is the calculated oxygen isotopic composition of the P-PO₄ sample at equilibrium
281 in ‰, $\delta^{18}\text{O}_w$ is the measured oxygen isotopic composition of the water in ‰, and T is the
282 temperature in degrees Kelvin.

283 Considering the stability of the $\delta^{18}\text{O}_w$ values during low-flow periods in the Redon River due to
284 the dominance of the ground water contribution to the discharge during these periods (Al-
285 Charideh, 1988), we used the $\delta^{18}\text{O}_w$ of the river water measured at each sampling date as the
286 reference isotopic value for water in equation 2. We considered both the minimum and
287 maximum temperatures measured over the period running between the sampling date and the
288 closest big storm flow event (around 10-20 days; Table 1 in the Supplementary Material), since
289 the sediments are (partially) renewed at each storm flow. The temperatures measured at the
290 time of sampling fell within these ranges. For samples 9 to 12 (sources), we used the $\delta^{18}\text{O}_w$
291 values measured in the local water sources, including rain, groundwater and soil water data,
292 and the extremes for the soil temperature measured within the previous 10 days at an agro-
293 meteorological station close to the studied sites. This allowed us to have a conservative and
294 time-integrated estimate of the equilibrium interval. For these samples, we also calculated the
295 deviation from equilibrium using the average stream water temperature and isotopic
296 composition, to verify whether or not their $\delta^{18}\text{O}_p$ values would fall within the equilibrium range
297 once transferred into the river. For each sample, the deviation from equilibrium was calculated

298 as the interval (in ‰) between the measured $\delta^{18}\text{O}_p$ values and the closest values for the
299 equilibrium range calculated from equation 2

300

301 **3. Results**

302

303 **3.1 Method validations**

304 Repeated extractions and purifications ($n = 3$ or 4) on the same sediment sample yielded $\delta^{18}\text{O}_p$
305 values with a standard deviation < 0.4 ‰. This repeatability can be considered as acceptable,
306 since it is within the range of the precision obtained on the repeated standards measurements on
307 the TC/EA-IRMS.

308 The purity of Ag_3PO_4 was 97%, as determined by analyzing the P- PO_4 in the redissolved
309 Ag_3PO_4 (see section 2.3.1, Table 2 in the Supplementary Material). The carbon and nitrogen
310 concentrations were generally lower than 1% and 0.1%, respectively (Table 2 in the
311 Supplementary Material). The C content was comparable to that measured in the internal
312 standard Ag_3PO_4 (0.5%) and much lower than the 23-29% or 0.2% reported by Weiner et al.
313 (2011) for C and N, respectively. The efficiency of the purification procedure was confirmed
314 by the oxygen yield given by the mass spectrometer (Fig. 1 in the Supplementary Material). All
315 of the samples were close to the regression line defined by the Ag_3PO_4 standards, indicating
316 that the amount of oxygen in the samples was close to the stoichiometry of the pure compound.
317 Most of the labeled/unlabeled pairs did not show significantly different $\delta^{18}\text{O}_p$ values. Only four
318 samples (2, 5, 8 and 9) showed different labeled and unlabeled values, meaning that the organic
319 P or condensed phosphate was hydrolyzed during the HCl extraction. These values have been
320 corrected with equation 1, as all of the other values are the average of the labeled and unlabeled
321 results.

322

323 **3.2 P speciation**

324 Table 3. P speciation and isotopic data

325

326 The P content and speciation varied strongly from one sample to another, with the Pt value
327 ranging from 459 to 1256 mgP/kg, Pi from 179 to 631 mgP/kg and Po from 36 to 450 mgP/kg.
328 P_{Olsen} accounted for 0.7 to 10.6 % of Pt depending on the samples. The highest P_{Olsen}
329 concentrations (65-67 mgP/kg, representing 8.4 to 10.6% of Pt) were found in the river
330 sediments sampled immediately downstream from the WWTP discharge. These variations are
331 the consequence of differences in the source, erosion and deposition conditions of the
332 sediments as well as differences in the P inputs that accumulate along the river (Table 1). The
333 downstream sediments (samples 5 to 8) and streamlet sediments (samples 1 and 2) were
334 enriched in Pt, Pi and P_{Olsen} compared to the sediments coming from the head of the catchment
335 (samples 3) or to the riverbank material (sample 12). The same relative enrichment is observed
336 for the cultivated topsoil (sample 10) compared to its lower horizon (sample 11).
337 Pt and P_{Olsen} were weakly and positively correlated ($r^2 = 0.49$, $p < 0.01$) as well as the P_{Olsen}/P_i
338 ratio with Pt concentrations of $r^2 = 0.76$ ($p < 0.01$), providing evidence that the P enrichment
339 process recorded by the downstream sediments is accompanied by an increase in the most
340 reactive fraction of the total P- PO_4 pool. The highest Po concentrations were found in the
341 topsoil samples (9 and 10).

342

343 **3.3. Measured isotopic signatures and comparison with the equilibrium values**

344 The measured $\delta^{18}\text{O}_p$ values for the HCl extracts varied between +11.8‰ (deep soil horizon,
345 sample 11) and +18.3‰ (wetland soil, sample 9; Fig. 2a). No significant statistical
346 relationships were found between the $\delta^{18}\text{O}_p$ values for the samples and their Pt, P_{Olsen} or Pi
347 content or ratios, nor with the water characteristics, notably the SRP concentrations. The

348 riverbank sample collected in the head of the catchment (sample 12) corresponds to a weakly
349 weathered moraine, as confirmed by field observations, i.e. no visible weathering and no
350 vegetation or biofilm on the banks, since they are eroded at each storm flow, and by the P
351 speciation data (very low $P_{O_{isen}}$ and P_o fractions; see Table 3). This sample is representative of
352 both the principal source of the upstream, unpolluted sediments and the parent material of most
353 of the soils in this catchment (CIPEL, 1988). Its $\delta^{18}O_p$ value (+15.9‰) indicates that it has a
354 mixed igneous/sedimentary origin, as this value is intermediate between the $\delta^{18}O_p$ values
355 reported for igneous or metamorphic apatite (approximately +7‰; Tamburini et al., 2012) and
356 for sedimentary apatite (between +17 and +23‰; Tamburini et al., 2014).
357 Interestingly, the variability in the $\delta^{18}O_p$ values for the allochthonous sources that may possibly
358 contribute toward the river particulate-P load was higher than the variability in the riverbed
359 sediments. Whereas while the soil, riverbank and riverbed sediments collected in the head of
360 the catchment (samples 1 to 3 and 9 to 12) are representatives of these sources (Dorioz and
361 Ferhi, 1994) and exhibit $\delta^{18}O_p$ values varying by > 6 ‰ (from +11.8 to +18.3‰), this variation
362 is reduced to approximately 2‰ for the sediments immediately upstream/downstream the
363 WWTP and to 1‰ for the outlet sediments (+12.8 to +13.8‰; Fig. 2a), the $\delta^{18}O_p$ signatures of
364 which were comparable to those recorded by the lake sediments (samples 13 and 14; Table 3).

365
366 [Figure 2]

367
368 We compared the measured $\delta^{18}O_p$ values with the theoretical equilibrium calculated with
369 Equation 2 (see Table 2). The deviation between the calculated and measured $\delta^{18}O_p$ values is
370 presented in Fig. 2b. The $\delta^{18}O_p$ value for the riverbank moraine sample (sample 12) was above
371 the calculated equilibrium by ca. 2‰, as was the wetland soil sample (9). This was not the case
372 for the cultivated soil samples that showed $\delta^{18}O_p$ values that were either very close to (-0.1‰,

373 topsoil, sample 10) or slightly lower than (-0.7‰, deep horizon, sample 11) the calculated
374 equilibrium values.

375 The riverbed sediment samples collected in the streamlets of the agricultural sub-catchment
376 (samples 1, 2) exhibited $\delta^{18}\text{O}_p$ values that were lower than the calculated equilibrium values by
377 ca. 2‰. For the sediments located within the main hydrogeological network, the differences
378 between the measured and calculated equilibrium values generally tended to be lower (see
379 Table 3 and Fig. 2b): ca. $\pm 1.5\%$ in the middle part, upstream and downstream from the
380 WWTP discharge point (samples 4 to 6) and $< 0.5\%$ for the sediments collected at the outlet
381 (samples 7 and 8) and the lacustrine sediments (samples 13 and 14).

382

383 **4. Discussion**

384

385 **4.1 Significance and variability of the measured and equilibrium $\delta^{18}\text{O}_p$ values**

386 The present study shows that the method originally established and optimized by Tamburini et
387 al. (2010) for soils is suitable for CaCO_3 -rich riverbed sediment samples, following some
388 modifications to the methodology. However, as for soils, it should be kept in mind that the
389 extraction is not entirely specific with regards to the nature of the extracted P-PO_4 . First, 1 M
390 HCl can be used to extract the fraction that is primarily composed of Ca-bound P, which can
391 contain a broad spectrum of P compounds (more or less crystallized apatite forms, CaCO_3 -
392 bound P, etc.). This reagent can also extract some P-PO_4 sorbed onto Fe and Al oxides, if used
393 as a single extractant. Thus, the $\delta^{18}\text{O}_p$ signatures obtained with this method correspond to a
394 pooled P-PO_4 isotope signature of a mixture of different P fractions with a different degree of
395 stability, exchangeability and bioavailability (Jarvie et al., 2002) rather than to the signature of
396 a geochemically specific P-PO_4 compartment. Despite this limitation, the oxygen isotope
397 composition of P-PO_4 extracted from sediment using HCl can provide information that is not

398 available with other methods, including the extent to which a P pool has been biologically
399 processed or derived from primary inputs or secondary anthropogenic sources (Tamburini et al.,
400 2014; Davis et al., 2014).

401 The calculation of the equilibrium $\delta^{18}\text{O}_p$ values is another critical point for the interpretation of
402 the $\delta^{18}\text{O}_p$ isotope data. Information regarding the kinetics of the equilibrium is only derived
403 from cell-free or soil incubation experiments (Blake et al., 2005; Melby et al., 2013), but not
404 from conditions that are close to the natural state. Our strategy was to take samples during low-
405 flow periods because of the stability of the $\delta^{18}\text{O}_w$ value and the temperature, but there is still
406 the question of how to choose appropriate time points when calculating the theoretical
407 equilibrium values with equation 2. This is also relevant for soils where sharp variations in the
408 $\delta^{18}\text{O}_w$ value in the soil surface layers due to the progressive evaporation of the soil water
409 (Hsieh et al., 1998) might be a major source of uncertainty. Amelung et al. (2015) suggest
410 calculating the equilibrium $\delta^{18}\text{O}_p$ value with the annual average $\delta^{18}\text{O}_w$ for rainwater, arguing
411 that the biological recycling of Ca-bound P in soil is a long-term process. Using an annual
412 rainwater $\delta^{18}\text{O}_w$ value and temperature values, the calculated equilibrium value for the
413 agricultural soil samples (10 and 11) would increase by approximately +2‰, resulting in
414 measured signatures below the equilibrium.

415 Except for a small number of studies (Elsbury et al., 2009; Gooddy et al., 2016; Markel et al.,
416 1994; McLaughlin et al., 2006a), freshwater ecosystems have been poorly explored in terms of
417 $\delta^{18}\text{O}_p$. Our results expand the global $\delta^{18}\text{O}_p$ library for these systems, comparing, for the first
418 time, the oxygen isotope signatures of HCl-P extracted from river sediment samples from
419 upstream to downstream with those of the main potential allochthonous sources in the
420 catchment. The isotope signatures of the analyzed river sediments, ranging from +12.2 to
421 +15.8‰, lie in the interval reported in previous studies, i.e. +8.6 to 22.5 (Markel et al., 1994),

422 +12.4 to 23.5‰ (Davies et al., 2014) and +9.7 to +20.0‰ (Tamburini et al., 2014). The same is
423 true for our soil samples (Amelung et al., 2015; Tamburini et al., 2014; Zohar et al., 2010).

424

425 **4.2 Biological recycling versus tracing the sources using $\delta^{18}\text{O}_p$ values**

426 *4.2.1 Soils*

427 Tamburini et al. (2012, 2014) have shown that the HCl-P fraction in the upper and most
428 biologically active part of the oldest soil of a chronosequence was formed by secondary
429 minerals precipitated from P_i that were previously cycled by biota. The close-to-equilibrium
430 $\delta^{18}\text{O}_p$ values measured in the agricultural soil samples (samples 10 and 11) can be viewed as
431 evidence that biological cycling had a large impact on the HCl-P fraction (which represents
432 roughly half of the P_t) in these samples. The soil P is supplied by the parent material and
433 fertilizations and is redistributed throughout the soil profile via biological activities. As the
434 parent material is low in P_t , P_{Olsen} and P_o , the high concentrations observed in the topsoil
435 sample (see Table 3), as in other cultivated soils, is the long-term result of the accumulation
436 and transformation of P favoring a high P bioavailability, which supports the assumption of an
437 intense biological activity.

438 The most common fertilizers used in the area have $\delta^{18}\text{O}_p$ values that are higher (16.2-22.5‰) or
439 lower (9.5‰; Guyard 2010) than the values for the studied agricultural soils. Even if we cannot
440 exclude that the observed equilibrium values were only the result of a mixing between the
441 parent material (close to +15.9‰) and a lighter fertilizer input, we assume that the fertilizer has
442 little influence because the fertilizer inputs are almost totally cycled over the long-term (> 1
443 year) and the amount applied is too small (2-3% of the HCl-P soil stock) over the short-term
444 (one season).

445 The high and above-equilibrium $\delta^{18}\text{O}_p$ value in the wetland surface soil (sample 9) can also
446 reflect the impact of biota, e.g. plants. The $\delta^{18}\text{O}_p$ value for this sample (18.3‰) cannot be

447 explained by the mixing of any of the potential input materials present in the area (i.e. bank
448 material or cultivated soil), as +18.3‰ is too far outside the range of the values (11.8 and
449 15.8‰) recorded by these input materials. The only source with a sufficiently high $\delta^{18}\text{O}_p$ value
450 would be plant-derived phosphate. Phosphate in plants is enriched in ^{18}O , with the $\delta^{18}\text{O}_p$ values
451 ranging from 28 to 31‰ (Pfahler et al., 2013; Tamburini et al., 2014). This suggests that the
452 extracted fraction might partially originate from the release of P from plant litter, either directly
453 or via decomposition. A significant fraction of this enriched P pool that is not taken up by
454 microbes (Liang and Blake, 2006; Oberson and Joner, 2005) might have precipitated or been
455 sorbed into the soil mineral pool. Interestingly, this sample had a particularly high Po
456 concentration. Preferential sorption/desorption reactions with soil minerals could also have
457 helped increase the $\delta^{18}\text{O}_p$ value for this sample. However, previous studies have reported both
458 negative and positive fractionation associated with sorption (Jaisi et al., 2010; Melby et al.,
459 2013b). In any case, this effect is probably too small (<1 ‰) to justify the observed value by
460 itself.

461 When compared to the equilibrium values calculated using the average river conditions (the
462 values in brackets in Table 3, Fig. 2b), the subsoil/embankment sample and the wetland topsoil
463 are no longer found within the equilibrium range. The situation is different for the agricultural
464 topsoil sample, the actual $\delta^{18}\text{O}_p$ value of which falls within the equilibrium range, meaning that
465 under these conditions the oxygen isotopic composition alone would not help to distinguish
466 between a sedimentary P- PO_4 pool inherited from this soil and in-stream recycled P- PO_4 .

467
468 *4.2.2 River sediments upstream and downstream from the wastewater treatment plant*
469 *discharge*

470 The September sampling showed that the input of P from wastewater was marked by a
471 threefold increase in the SRP concentration in the river water sampled upstream and

472 downstream from the discharge point (Table 1), accompanied by a drastic change in the Pt, Pi
473 and P_{Olsen} concentrations in the corresponding sediments (Table 3). No changes were detected
474 in the Po content, meaning that the impact of the point discharge on this pool in the bed
475 sediments was negligible (Dorioz et al., 1998a). Interestingly, despite the small distance (< 100
476 m) separating the upstream and downstream sites, the $\delta^{18}\text{O}_p$ values were also distinct, with a
477 difference of 3‰. Upstream from the WWTP discharge point, the P pools in the sediment are
478 mainly impacted by sources comprising a mixture of riverbanks and agricultural soil erosion
479 (materials similar to samples 12 and 1, 2). The $\delta^{18}\text{O}_p$ value (12.2‰) and P speciation for
480 sample 4 were intermediate between those of these source materials.

481 The two September samples (4 and 5, immediately upstream and downstream from the WWTP)
482 deviate from the isotopic equilibrium values (by -1.5 and +1.4‰, respectively; Fig. 2b),
483 suggesting the existence of conditions that prevent the complete biological turnover of P at both
484 sites (also see Goody et al., 2016). Under the assumption that biological equilibration is the
485 dominant process during sewage treatment (Gruau et al., 2005), the $\delta^{18}\text{O}_p$ value for dissolved
486 Pi from the WWTP would vary between 13.0 and 14.8‰ using equation 2 (confirmed by the
487 value of 14.0‰ measured by Guyard, 2010), meaning that the higher $\delta^{18}\text{O}_p$ value downstream
488 (15.8‰) cannot be interpreted as the result of a mixing between wastewater derived P and P
489 from the upstream sediments. One possible explanation for this could be that part of the Pi
490 injected into the river by the WWTP is rapidly (within a few meters from the injection point)
491 and continuously sorbed onto the sediment particles, as also hypothesized by Dorioz et al.
492 (1998a), and that the heavier isotopologues are preferentially sorbed. However, there is no
493 consensus regarding the associated fractionation (Jaisi et al., 2010; Melby et al., 2013b). An
494 alternative explanation would be that the upstream sample 4 represents a local anomaly, and the
495 signature of the downstream sediments, sample 5, derives from local riverbank erosion. Sample
496 12 can be considered as representative of riverbank material and its $\delta^{18}\text{O}_p$ value is comparable

497 to that of sample 5. However, this hypothesis is unlikely as it does not explain the drastic
498 change in P speciation observed in the sediments downstream from the WWTP (see Table 3).
499 Only the locality situated immediately downstream from the WWTP discharge point could be
500 sampled in March 2012 (sample 6). The SRP, Pt, Pi and P_{Olsen} concentrations were as high as
501 those found in sample 5, which was collected in September, and much higher than the
502 concentrations measured further upstream, at location 3, once again providing evidence for the
503 strong impact of the WWTP discharge on the river and sediment P pool. The $\delta^{18}\text{O}_p$ value for
504 sample 6 was similar to the September sample (sample 5), although it was close to the
505 calculated isotopic equilibrium. The lack of a reference value equivalent to sample 4 in March
506 means that it is not possible to conclude whether the observed value resulted from the complete
507 equilibration of the sedimentary P-PO₄ pool or if it was instead produced by selective sorption
508 processes as we suggest for sample 5. A repeated seasonal sampling of the sediments
509 immediately upstream and downstream from the WWTP could possibly clarify this point.

510

511 *4.2.3 Head and outlet sediments*

512 The possible mixing of isotopically different sediment source materials coupled with
513 differences in the sediment retention time complicates the interpretation of the $\delta^{18}\text{O}_p$ values for
514 the riverbed sediments. Under conditions of low biological activity and a short residence time,
515 the isotopic signature of the sediments is expected to represent primary source signatures. In
516 this context, an isotopic disequilibrium would indicate the existence of factors that limit the in-
517 stream P cycling, intended as uptake by the biota, intracellular equilibration and subsequent
518 release. As judged from the present data, a limiting situation for in-stream P recycling could
519 have been prevalent in the head of the Redon River catchment, where strong out-of-equilibrium
520 $\delta^{18}\text{O}_p$ signatures have been recorded (see Table 3 and Fig. 2b). This could be first related to the
521 physical conditions in this part of the catchment - high-energy stream (total renewal of the

522 sediments with each storm flow) and a short distance from the sources, i.e. riverbanks - which
523 both determined the short residence times. The oligotrophic state of the running water (SRP <
524 10 µg P/l, NO₃ < 0.5 mg N/l) and the low P bioavailability of the riverbed sediments (as
525 indicated by low P_{Olsen} concentrations) could contribute to limit the development of benthic
526 biota and thus P uptake and subsequent release. For the same reasons the contribution of
527 extracellular mineralization of organic P is expected to be negligible. The lack of an isotopic
528 difference and the small differences in the Pt, Pi and Po content between the riverbank sample
529 (12) and the sediment sample (3) collected immediately downstream further confirms the direct
530 filiation between the source and the sediment.

531 The hypothesis that a rapid transfer does not significantly modify the δ¹⁸O_p signature is further
532 supported by the disequilibrium signature recorded in the streamlet samples 1 and 2 and their Pi,
533 Po and P_{Olsen} concentrations, as all of these parameters are intermediate between the values
534 found in the analyzed deep and topsoil samples. This suggests i) a mixed origin for these two
535 sediment samples derived from both superficial soils and ditch banks or field embankments;
536 and ii) that the biological activity was not intense or not long enough to equilibrate the original
537 P oxygen isotope signatures, consistently with the expected very short residence time of the
538 sediments in the streamlets (stream order of 1).

539 The situation was different for the sediments that were collected closer to the river outlet or in
540 the lake (stream order higher than 3). Close-to-equilibrium values were found in these samples.
541 A corresponding increase in the SRP concentration in running water (a factor of 8 between
542 upstream and the outlet; see Table 1), as the result of the cumulative anthropogenic inputs in
543 the river, was also observed.

544 The equilibrium values observed in the lowest part of the Redon River suggest the occurrence
545 of processes that are able to efficiently recycle the sedimentary P-PO₄. This hypothesis would
546 be consistent with the decrease in the δ¹⁸O_p values observed during the September campaign

547 downstream from the WWTP discharge point, i.e. between samples 5 and 8. There was a 3‰
548 decline in the $\delta^{18}\text{O}_\text{P}$ value between sample 5, which was collected immediately downstream
549 from the WWTP discharge point, and sample 8, which was collected approximately 2 km
550 further downstream. This decrease was accompanied by a strong narrowing of the gap between
551 the measured and calculated equilibrium $\delta^{18}\text{O}_\text{P}$ values. Whereas this gap was +1.4‰ for
552 sample 5, it was reduced to < 0.4‰ for sample 8 (see Table 3). The isotopic equilibration
553 inferred by these data would likely occur in the sediments, given that in this kind of torrential
554 rivers most of the biological activity essentially takes place in the benthic system. Equilibration
555 could be further promoted by the longer residence time of the sediments in the downstream part
556 of the river (Doriz et al., 1989). It may have been also driven by the enhanced bio-availability
557 of P in the lower part of the river, notably downstream from the WWTP, as evidenced by the
558 sharp increase in the P_{Olsen} concentration starting from this point (see Table 3, Figs. 2a and 2b).
559 Most likely, the overall process should be viewed more as the integration of a succession of
560 partial equilibrium occurring over different storage stages under similar equilibrium conditions,
561 as it is unlikely that the bulk of the P- PO_4 could be completely cycled by the benthic biota
562 during a single low-flow period.

563 However, we lack information about the real residence time of the sediments and the P
564 recycling rates in the Redon River. Considering that the measured $\delta^{18}\text{O}_\text{P}$ values for the
565 sediments collected near the river outlet (samples 7 and 8) are identical within error to those
566 measured in the upstream streamlet sediments (samples 1 and 2) and are within the range of the
567 values found in the possible sediment sources (see Fig. 2a), we cannot entirely exclude that the
568 $\delta^{18}\text{O}_\text{P}$ values in these downstream sediments are simply source-inherited isotopic values. More
569 direct evidence for the real residence time of the sediments and the ability of aquatic biota to
570 efficiently recycle P- PO_4 in the downstream part of the river is needed to decide between these
571 two hypotheses.

572

573 4.3 Potential and limits of the oxygen isotopes tool

574 We hypothesize that is possible to trace the potential sources of sediment particulate-P-PO₄
575 using oxygen isotopes, assuming a limited number of potential sources and under conditions of
576 low biological activity and a short residence time in the river network. In the studied torrential
577 system, it appeared possible to trace these sources in the head of the catchment (only two
578 sources, rapid and high-energy transfer, limiting conditions for the biological activity). It would
579 be also possible at the catchment scale, during winter high-flow periods, when the sediments
580 that originate from banks and agricultural soil erosion are rapidly transferred as suspended
581 matter and without significant interaction with point source discharge and biological activity. In
582 this case, the $\delta^{18}\text{O}_\text{p}$ value would primarily be the result of a two end-members mixing between
583 the cultivated soil (values close to 11-14‰) and moraine material (values close to 15-16‰).
584 However, the isotopic tracing is expected to be less straightforward during the summer and
585 early fall periods, as the storm flows that occur at that time primarily mobilize riverbed
586 sediments. Therefore, it could become more difficult to perform a quantitative assessment of
587 the original particulate-P sources using $\delta^{18}\text{O}_\text{p}$ due to an overprinting of the sediment isotopic
588 signature induced by the high biological activity. During summer low flows, an even stronger
589 overprinting of the original isotopic signature is expected to occur in the downstream part of
590 the river, due to the input of anthropogenic, bioavailable P, a longer sediment residence time
591 and increased biological activity. This being the case, $\delta^{18}\text{O}_\text{p}$ could then instead be used as a
592 tool to assess the in-stream biological turnover of the P.

593

594

595 5. Conclusion

- 596 • In this study, we successfully adapted Tamburini et al.'s method (2010) for soil for use
597 with river sediments to provide the first assessment of the variability in $\delta^{18}\text{O}_\text{p}$ from HCl-
598 extracted P in river sediments along a gradient of increasing anthropogenic impact.
- 599 • We observed a contrast between the $\delta^{18}\text{O}_\text{p}$ signature and the speciation of the sediments
600 collected in the head and at the outlet of the catchment. The sediments that were collected
601 upstream have isotopic signatures that are far from isotopic equilibrium and within the range of
602 the signatures found in the possible sediment sources (i.e. riverbank material), while those
603 collected at the outlet showed $\delta^{18}\text{O}_\text{p}$ values close to isotopic equilibrium. We related this
604 difference to the contrast between the highly dynamic character (short residence time) of the
605 transfer system in the head of the catchment (preservation of the isotopic signature of the
606 source material in the sediments) and the “quieter” sediment transport system in the lower
607 catchment, which, along with the input of the bioavailable anthropogenic P, likely fostered the
608 recycling of P by benthic biota, leading to an equilibration in the $\delta^{18}\text{O}$ signatures. However,
609 with the available data, we cannot exclude that the signatures at the outlet were the result of the
610 mixing of sediments with higher (upstream/wetland sediments) and lower signatures
611 (agricultural soil erosion).
- 612 • Uncertainties remain about the calculation of the equilibrium window. For aquatic
613 environments, the duration of the in-stream equilibration processes should be taken into
614 account, together with the perturbation caused by changes in the hydrological regime. This
615 would require an experimental approach for a proper estimation.
- 616 • In terms of the quantitative tracing of potential particulate-P sources, future
617 investigations should focus on winter high flows, when rapid sediment transfer maximally
618 reduces the possibility of biological interactions. On the other hand, the sensitivity of $\delta^{18}\text{O}_\text{p}$ to
619 biological processes opens up new promising perspectives for studying in-stream processes. In

620 situ labelling experiments could help develop these studies by providing unique information
621 about the biological uptake, turnover and/or organic P mineralization.

622 • More insights about the way and extent the biological activity controls P speciation in
623 the soil and sediments could be possibly provided by the development of refined extraction
624 methods that are able to selectively and quantitatively extract other P pools like the one
625 associated with Al and Fe hydroxides, P_{Olsen} and P_o or finer fractions (e.g. $< 50 \mu\text{m}$, $< 2 \mu\text{m}$).

626

627

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634

635

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Table and Figure captions

Table 1. Characteristics of the sediment sampling sites.

Table 2. Characteristics of the soil sampling sites.

Table 3. P speciation and oxygen isotope composition.

Figure 1. Redon River catchment and sampling sites; the capital letters refer to the sampling locations, the numbers are the sample code.

Figure 2. Oxygen isotope composition ($\delta^{18}\text{O}_p$) and deviation from isotopic equilibrium. The gray circles represent potential source samples, white circles represent riverbed sediment samples, and $\text{Pols} = \text{P}_{\text{Olsen}}/\text{P}_t$ in % (note that the deviation of the potential sources from equilibrium in this image have been calculated according to the average within-river conditions).

Figure 3. Conceptual model describing the evolution of $\delta^{18}\text{O}_p$ for the riverbed sediments in the studied torrential system.

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Table 1 Characteristics of sediment sampling sites

Site	SO	Sample code	Sampling date	Q	EC	SRP	Altitude	Drainage area	Land use of the drainage basin	Main sources of stream P
				L/s	mS/cm	mg P/L	m a.s.l	ha		
A	1	1	September 3, 2012	<1	543	0.0012	576	4	Cultivated land	NPS (erosion of cultivated soils and tiches)
B	1	2	September 3, 2012	1	557	0.009	577	17	Cultivated land	NPS (erosion of cultivated soils and tiches)
C	2	3	March 26, 2012	35	598	0.008	696	510	Steep forest	NPS (natural: mainly river bank erosion)
D	3	4	September 3, 2012	55	7203	0.027	510	2310	Forest and cultivated land	NPS (natural, agricultural)
E	3	5	March 26, 2012	105	654	0.056	508	2311	Forest and cultivated land	NPS (natural, agricultural) + PS (WWTP)
		6	September 3, 2012	55	753	0.092				
F	4	7	March 26, 2012	340	597	0.062	405	3300	Forest and cultivated land	NPS (natural, agricultural) + PS (WWTP)
		8	September 3, 2012	85	975	0.067				

SO = Strahler stream order (Strahler, 1957)

Q = water discharge

EC = electric conductivity

SRP = soluble reactive phosphorus concentration in water, measured on filtered samples (0.45 μm) by the molybdate-blue method (Murphy and Riley, 1962)

NPS = Non-point source, PS = Point source

WWTP = waste-water treatment plant

Table 2 Characteristics of soil sampling sites

Sample code	Date of sampling	Location	Soil depth	Altitude	Sampling conditions
			cm	m a.s.l	
9	September 3, 2012	Wetland topsoil	-1 to -5	552	Organic soil, saturated
10	September 3, 2012	Cultivated topsoil ^a	-10 to -30	558	Cambisol on moraine deposit, dry
11	September 3, 2012	Cultivated soil (embankment) ^a	-60 to -70	558	Cambisol on moraine deposit, saturated
12	September 3, 2012	Riverbank in a forested upland area	-	980	Würm moraine deposits, saturated

^a samples collected within an agricultural sub-catchment of 33 ha (Figure 1)

Table 3. P speciation and oxygen isotopes composition

Sample code	Sample type	Pt	P _{Olsen}	Pi	Po	P _{Olsen} /Pt ^a	$\delta^{18}\text{O}_p$	St. Dev.	n	Deviation from equilibrium (‰) ^b
		mgP/kg	mgP/kg	mgP/kg	mgP/kg	%	‰	‰	-	‰
1	River bed sediments ^d	791	14	401	180	2.4	12.5	0.3	3	- 2.4
2		751	20	323	208	3.8	13.3 ^s	0.3	2	- 1.6
3		528	8	411	78	1.7	15.5	0.6	4	+ 1.3
4		609	18	398	202	2.9	12.2 ^s	0.4	2	- 1.5
5		1018	65	586	194	8.4	15.8 ^s	0.7	2	+ 1.4
6		962	67	470	166	10.6	15.2	0.3	3	< 0.4
7		734	26	439	138	4.5	13.8	0.4	2	< 0.4
8		736	21	324	90	5.2	12.8	0.1	3	- 0.6
9	Wetland soil	1163	32	179	383	5.7	18.3 ^s	0.2	2	+ 2.1 (+ 3.6) ^c
10	Agric. topsoil	1256	43	631	450	6.2	14.0	0.4	2	< 0.4 (< 0.4) ^c
11	Embankment/ deep soil	755	5	363	76	1.1	11.8	0.4	2	- 0.7 (- 2.0) ^c
12	Riverbank (moraine)	511	3	305	36	0.7	15.9	0.2	2	+1.9 (+1.2) ^c
13	Lake	459	15	365	121	3.5	12.9	0.3	2	< 0.4
14	sediments	539	17	420	70	3.1	13.9	0.5	2	+ 0.8

^a As a sum of Pi and Po fractions

^b See section 2.3.2 and equation 2

^c Values not in brackets are calculated according to the temperature and water isotopes conditions at the sampling site, the values in brackets are calculated according to the “within-river” conditions

^d Sediment samples are ranked according to the corresponding drained area

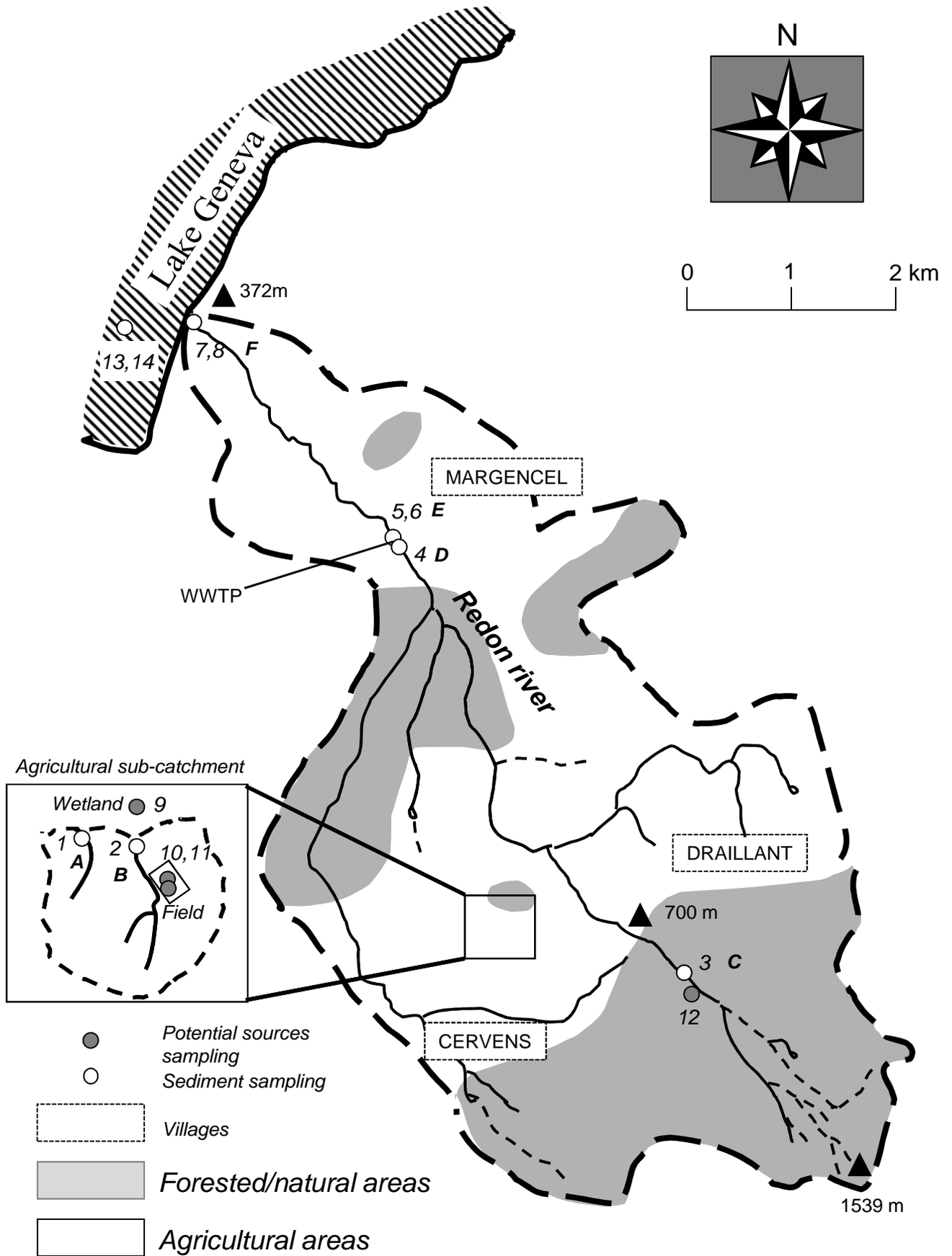


Figure 1. Redon River catchment and sampling sites, capital letters refer to sampling locations, numbers to sample code

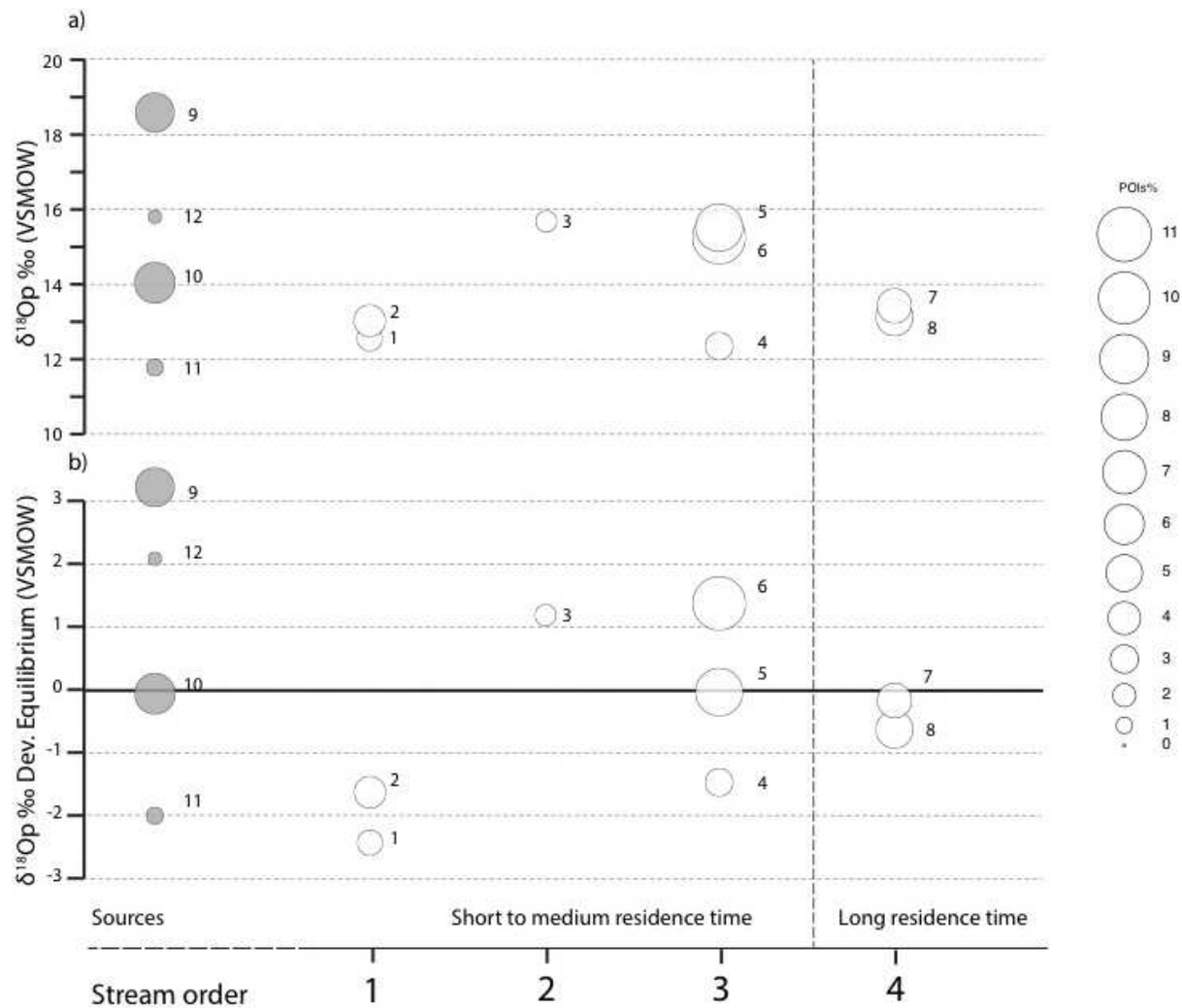


Figure 2. Oxygen isotopes composition $\delta^{18}\text{Op}$ and deviation from isotopic equilibrium. Grey circles represent potential sources samples, white circles represent riverbed sediments samples, $P_{ols} = P_{Olsen} / P_t$ in % (note that deviation from equilibrium of potential sources in this picture have been calculated according to average within-river conditions)

Highlights

- We provided the first assessment of the oxygen isotopic composition of HCl-extracted phosphate in river sediments
- We observe a contrast in the oxygen isotope composition between upstream and downstream sediments
- This contrast can be related to anthropogenic impacts on the river system and to the sediment residence time
- We have shown that in some situations it is possible to trace particulate-P sources of sediments with this isotopic tool