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Relevance of using the non-reactive geochemical signature in sediment core to estimate historical tributary contributions

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Key words: geochemical fingerprinting, sediment core, non-reactive fraction of metals, suspended particulate matter (SPM)

Highlights:

Assessment of historical sediment sources with an original fingerprinting approach

Apportionment modelling using total and residual concentrations of inorganic elements

More tracers available with non-reactive fraction to discern historical SPM sources

Total concentrations of tracers lead to inaccurate historical sources apportionment

More reliable SPM contribution from Upper Rhône tributaries with residual fraction

1 **1. Introduction**

2 Suspended particulate matter (SPM) is important for biogeochemical cycling and aquatic ecosystem
3 functioning, but it can also have negative impacts on water quality, ecosystems, and the economy (i.e.
4 causing dysfunctions in power plants that use river water, filling reservoirs) (Navratil et al., 2012;
5 Torres-Astorga et al., 2018). Identifying SPM sources can help improve river management to reduce
6 the sediment and associated contaminants load in rivers by implementing remediation actions in
7 source areas. In general, sediment source fingerprinting studies focus on estimating spatial sources of
8 SPM in hydrosystems or on methodological aspects (Walling, 2005; Collins et al., 2017).

9 Fingerprinting methods for tracing sources of SPM or sediment, which has evolved considerably since
10 the 1970s, employ tracers (Walling, 2013; Collins et al., 2017). Tracer science has also evolved with
11 analytical techniques developed to greatly increase the number of physical, biological, geochemical
12 and other properties that can serve for source tracing (Walling, 2013; Collins et al., 2020). A majority
13 of fingerprinting studies have used geochemical properties as tracers (e.g. trace metals/metalloids,
14 major elements; Collins et al., 2020), which is explained by the inclusion of these properties in the
15 composition of soils and SPM that are typical of each watershed. Thus, these properties provide an
16 effective tool to discriminate the potential sources identified.

17 As signalled in the review of D'Haen et al. (2012), few studies have focussed on the application of
18 source tracing to sediment cores. However, the historical inputs of SPM from different sources can
19 provide retrospective information on environmental events (floods) or modifications of SPM fluxes,
20 such as dam construction, bank stabilization or revegetation (Collins et al., 1997; Navratil et al., 2012).
21 Dhivert et al. (2016) showed that past anthropogenic inputs were higher for several metals/metalloids
22 (e.g. Bi, Cd, Hg, As, Pb), leading to increased concentrations of some properties in the deepest layers
23 of the core. We therefore deduce that there are differences in concentrations between the sources and
24 the deepest layers of the sediment core, which means that the elements concerned are reactive because
25 the geochemical signature of the deposited SPM has changed over time. Moreover, source
26 fingerprinting in sediment cores does carry some limitations, since total metal concentrations that
27 include anthropogenic inputs and the use of contemporary SPM are commonly used in literature to
28 trace historical sources.

29 In order to account for the conservatism of tracers in the total fraction, fingerprinting studies generally
30 apply a mass conservation test called the range test (Collins et al., 2017). However, Owens et al.
31 (2016) showed that the range test lacks reliability as some tracers could pass the test yet still have non-
32 conservative behaviour. Moreover, a conservative property could be out of the range of the considered
33 sources, as there might be one unidentified source. Given the challenge of establishing tracer
34 conservatism, using the total concentration of tracers may add uncertainties on the model results due
35 to the reactivity of some of the metals (i.e. including metals and metalloids in the following text). To
36 address this issue, Dabrin et al. (2014) tested the use of trace elements analyzed in the non-reactive

37 fraction to trace qualitatively the origin of contemporary SPM from two sources (tributaries) in the
38 highly reactive Gironde estuary. More recently, a similar approach proved useful to trace the SPM
39 contributions of five tributaries on the Upper Rhône River (Dabrin et al., 2021). Furthermore, several
40 studies show that total metal concentrations in sediment have widely varied over the past decades (e.g.
41 Audry et al., 2004; Dhivert et al., 2016), and may be modified as a result of diagenetic processes that
42 occur in the sediment column (D'Haen et al., 2012; Collins et al., 2017). It is therefore essential to
43 ensure the conservatism of properties for tracing SPM sources over a large time-scale (i.e. using
44 sediment cores). Thus, the use of the non-reactive fraction could be advantageous for the application
45 of fingerprinting methods in a multitude of environments (i.e. estuaries, delta, acid mine drainage)
46 and, then, implement management actions aimed at reducing major sediment inputs to rivers.

47 There are still only few studies focusing on the potential reactivity of metals in the total fraction
48 (Ankers et al., 2003; Carter et al., 2006; Pulley et al., 2015). Pulley et al. (2015) applied the range test
49 to investigate tracer conservatism (i.e. magnetic, radionuclide, and geochemical tracers) in layers of
50 six sediment cores in the Nene River watershed (UK). If some properties failed the range test for some
51 core layers, then the corresponding layers were removed from further analysis (Pulley et al., 2015).
52 This method, advocated by Collins et al (2017), results in the loss of a large amount of information on
53 sources over time, information which may be essential for this type of study. Another limitation in the
54 identification of historical inputs of SPM is the difference in grain size between contemporary SPM
55 sources and sediment core layers. As many metals (i.e. As, Cu, Cd, Ni, Co, Zn, Pb) have a strong
56 affinity with fine particles (Horowitz, 1991; Koiter et al., 2015), SPM can have higher concentrations
57 of some metals than the coarser particles deposited in sediment cores. To homogenize the particle size
58 distribution between sediment core layers and source samples (SPM from tributaries or surface soil
59 samples), some studies include a particle size correction of the geochemical concentrations (metals,
60 rare earths and major cations; Manjoro et al., 2017).

61 Given the methodological limitations to ensuring tracer conservatism, here we set out to: (1)
62 investigate the relevance of using the non-reactive fraction of metals in fingerprinting approaches
63 applied on a sediment core, and (2) assess the historical contributions of SPM inputs in a sediment
64 core collected on the Upper Rhône River, downstream of three major tributaries, using the total and
65 non-reactive fractions. For that purpose, we focussed on historical SPM inputs in the Rhône River
66 basin, which is one of the largest watersheds and the main contributor of SPM to the Mediterranean
67 Sea (Launay, 2014; Zebracki et al., 2015; Delile et al., 2020).

68

69 **2. Materials and Methods**

70 **2.1. Study area**

71 The Rhône River is one of Europe's major rivers and the biggest sediment input to the Mediterranean
72 Sea, with an inter-annual mean of $324 \text{ t.km}^{-2}.\text{year}^{-1}$ for the 1994–1995 period (Launay, 2014;
73 UNEP/MAP, 2003). Its watershed covers an area of $95,600 \text{ km}^2$ and a wide diversity of geological and
74 climatic conditions (Zebracki et al., 2015; Delile et al., 2020). Since 2010, concentrations and fluxes
75 of SPM and associated contaminants of the Rhône River and its tributaries are monitored and managed
76 under the Rhône Sediment Observatory (OSR) program. A monitoring network up and running since
77 2009 collects a large set of SPM samples and data, which are freely available from the online database
78 of the OSR (Thollet et al., 2018). Our study focussed on a simplified system on the upper part of the
79 Rhône River catchment ($\approx 7000 \text{ km}^2$). This area includes three main tributaries located upstream of the
80 sediment core location (Fig. 1), i.e. the Arve, the Fier, and the Guiers Rivers, which have catchment
81 areas of 2164 km^2 , 700 km^2 and 614 km^2 , respectively. The Lake Geneva outlet, located upstream of
82 the Arve River, is considered a negligible source of SPM to the Rhône River, in line with Launay
83 (2014) who reported that Lake Geneva acts as a reservoir for deposited SPM and that the quantities of
84 SPM discharged at its outlet into the Rhône River are negligible. In terms of contributions of each
85 tributary in terms of average SPM fluxes, the Arve, Fier and Guiers Rivers brought 0.49 , 0.18 and 0.07
86 Mt.year^{-1} respectively, for the period 2001–2011 (Launay, 2014).

87 2.2. Sampling strategy

88 2.2.1. Sources sampling

89 The SPM samples were collected on the three main tributaries of the Upper Rhône River, the Arve,
90 Fier and Guiers Rivers, which are the potential SPM sources identified in this study (Fig. 1). Each
91 station is located a few kilometres upstream from the confluence with the Rhône River (Thollet et al.,
92 2018). The SPM were collected using particles traps deployed throughout the year and retrieved every
93 month. For the Guiers River, due to repeated vandalism, we had to carry out spot sampling using a
94 continuous centrifuge (Westfalia KA 2-86-76) or by manually sampling large volumes of water. A
95 more detailed description of the sampling methods can be found in Masson et al. (2018). Overall, 22
96 SPM samples ($n = 7$ [flood: 3, base water flow: 4] for the Arve River, $n = 15$ [flood: 8, base water
97 flow: 7] for the Fier River, $n = 6$ in flood condition for the Guiers River) were collected between 2012
98 and 2019, and were used to identify the SPM signature of each tributary.

99 2.2.2. Targeted sediments: sediment core sampling

100 A 101 cm-long sediment core was sampled from a flat-bottom boat with Uwitec gravity corer
101 (including hammer accessory) equipped with a transparent PVC liner (diameter, 90 mm). This core
102 was collected in a secondary channel of the Rhône River (close to the sediment core sampled in 2008
103 by Desmet et al. (2012)) at a point located downstream of the confluence of the Guiers River
104 (45.7034806 N , 5.555225 E), during a low-water period in September 2018 (Fig. 1). This site is
105 always submerged with inflows from the Rhône River downstream and from La Morte River upstream

106 (see Morereau et al. (2020) for a detailed description of the sampling area). An apparent 31%
107 compaction of the sediment core was observed. We identified 25 roughly 4 cm-thick layers and each
108 layer was sub-sampled using a ceramic knife in order to collect from 32.7 to 133.8 g of wet sediment.
109 A subsample of fresh sediment was stored for particle size analysis. Other sediment subsamples were
110 freeze-dried, grinded and stored in plastic bags until further analytical steps.

111 2.3. Analysis of trace and major elements in the non-reactive fraction and sediment core 112 dating

113 The analysis of geochemical properties in SPM and sediment samples involved the quantification of
114 20 trace and major elements in the total and reactive fraction. The total fraction was determined after
115 triacid mineralization (SI.1). The reactive fraction was obtained by a soft extraction using hydrochloric
116 acid (1 M) at room temperature (Dabrin et al., 2014). The difference between the concentrations of the
117 two fractions gives a so-called 'non-reactive fraction'. Metals were analyzed in both fractions by
118 inductively-coupled plasma optical emission spectroscopy or triple quadrupole inductively-coupled
119 plasma mass spectrometry. Furthermore, samples were analyzed for particle size distribution,
120 according to the method described by Masson et al. (2018), in order to correct the metal concentrations
121 when necessary. More details on analytical methods can be found in the Supplementary material SI 1
122 and in Dabrin et al. (2014; and 2021).

123 The sediment core dating using $^{210}\text{Pb}_{\text{xs}}$ (^{210}Pb in excess) and ^{137}Cs concentrations is detailed in the
124 Supplementary material SI 1 (see Morereau et al. (2020) for additional information).

125 2.4. Statistical analysis and mixing model

126 2.4.1. Particle size correction of metal concentration data

127 Statistical tests (Student's *t*-test or Wilcoxon test) were used to compare SPM particle size distribution
128 between tributaries. As sediment core samples have coarser particles than SPM samples (D'Haen et
129 al., 2012; Huang et al., 2019), it is essential to correct the metal concentration data. Here, despite
130 ongoing debate surrounding this methodological aspect, we applied the particle size correction
131 implemented by Gellis and Noe (2013), which is based on examining the relationship between tracer
132 (metal) concentrations and particle size distribution. If this relation was significant (Pearson's test, p
133 < 0.05), a correction was applied according to equation 1:

$$134 \quad C_n = C_{is} - [(D_{50(S)} - D_{50(\text{Sed})}) \times p] \quad \text{Eq. 1}$$

135 where C_n is corrected concentration of tracer i , C_{is} is initial concentration of tracer i in source s , $D_{50(S)}$
136 is median particle size value of source s , $D_{50(\text{Sed})}$ is average D_{50} value for all target sediment samples,
137 and p is slope of the regression line.

138 2.4.2. Selection of tracers to discriminate the SPM sources

139 To discriminate the three potential SPM sources (the Arve, Fier and Guiers Rivers), we selected
 140 tracers via a two-step procedure that included two statistical tests: a Kruskal-Wallis test and a
 141 discriminant factor analysis (DFA) (Collins et al., 1997). The range test was performed to test tracer
 142 conservatism before the tracer selection step. For this test, tracer concentration in the target sediment
 143 (layers of the sediment core) must be within the range of tracer concentrations in the three potential
 144 sources (Manjoro et al., 2017; Pulley et al., 2018). We defined the range test from the minimum and
 145 maximum values of the concentrations of each source (C_{is}) and the standard deviation (SD) (Eq. 2). To
 146 fit the range test, the concentrations in each sediment core layer (C_i) must be within the defined limits
 147 (Sanisaca et al., 2017).

$$148 \quad [\min(C_{is})]_{\text{mean}} - \text{SD} \times [\min(C_{is})]_{\text{mean}} < C_i < [\max(C_{is})]_{\text{mean}} - \text{SD} \times [\max(C_{is})]_{\text{mean}} \quad \text{Eq. 2}$$

149 This test eliminates properties for which the concentration in sediment samples is outside the range of
 150 the SPM source concentrations. To discriminate the three identified potential sources, we first selected
 151 tracers based on the range test using a Kruskal-Wallis test to highlight redundant elements between
 152 sources (Collins et al., 2010). Then, the fingerprint properties that showed a significant difference
 153 between the three sources ($p < 0.05$) were tested by DFA to ensure an optimum combination of tracers.
 154 Finally, DFA was used to determine the percentage of correctly classified SPM source samples
 155 (ideally, 100% for each source) and the discriminatory power of each tracer. The results are
 156 summarized in Table 1.

157 2.4.3. Estimation of the source contributions and associated uncertainties

158 There are several models for estimating source contributions, but the mixing model combined with
 159 Monte Carlo simulation remains the most widely used method in the literature. Here we employed this
 160 model to quantify the SPM source contributions, as detailed in Eq. 3:

$$161 \quad C_i = \sum \sum (P_s \times C_{is}) / 1000 \quad \text{Eq. 3}$$

162 where P_s is percentage contribution from the SPM tributary s , C_{is} is concentration of tracer i in the
 163 tributary s , n is number of tributaries, and C_i is tracer concentration in the target sediment samples
 164 (Hughes et al., 2009; Haddadchi et al., 2013). Note that this model is based on two conditions: the
 165 source contributions should be between 0 and 1, and the sum of the contributions is equal to 1 (Hughes
 166 et al., 2009; Navratil et al., 2012; Collins et al., 2017).

167 To estimate the uncertainties associated with modelled contributions, the mixing model was combined
 168 with a Monte Carlo analysis. For each target sediment sample (i.e. layers of the sediment core), this
 169 analysis provided 1000 model iterations that represent the optimized number of replicates most
 170 widely-used in the literature. These iterations are used to determine the mean of the contributions from
 171 each source and the associated uncertainty in the form of standard deviation (SD) and 95% confidence

172 intervals (CI95). Some studies also use an estimator of model precision, the mean absolute error
173 (MAE), calculated according to Eq. 4 (Huang et al., 2019):

$$174 \text{ MAE} = \left\{ 1 - \frac{\sum |C_i - (\sum P_s \times C_{is})|}{C_i / m} \right\} \quad \text{Eq. 4}$$

175 where m is number of properties. If the MAF is greater than 0.85, the model results are reliable.

176

177 **3. Results and Discussion**

178 **3.1. Geochemical signature of total and non-reactive fractions of tributary SPM**

179 Geochemical composition of SPM in each source (tributaries) collected between 2012 and 2019 are
180 reported in Fig. 2. The description and comparison of concentrations between tributaries are presented
181 in Supplementary material 3 (SI.3).

182 Based on the relative proportion of the metals in the non-reactive fraction compared to the total
183 fraction, we determined three mains groups: group 1 (Fig. 2 a) gathers low-reactivity properties (83%–
184 99% of total fraction, Al, Ba, Cr, Li, Ti and V), group 2 (Fig. 2 b) gathers moderate-reactivity
185 properties (58%–78% of total fraction, As, Co, Fe, Mg and Ni) and group 3 (Fig. 2 c) gathers high-
186 reactivity properties (17%–46% of total fraction, Cd, Cu, Zn, Mn, Sr and Pb). Thus, the use of Cd, Cu,
187 Zn, Mn, Sr and Pb, which are highly reactive elements, should clearly be avoided for fingerprinting
188 studies as they can decrease the reliability and accuracy of the estimation of source contributions by
189 the models (Lacey et al., 2017; Collins et al., 2017).

190 First, Al, Ba, Cr, Li, Ti and V (group 1, Fig. 2a) showed only minor differences in concentrations
191 between total and non-reactive fractions. Nonetheless, the reactivity of the properties differs somewhat
192 according to tributary. In the case of Al, Cr and Ti, concentrations did not differ significantly
193 (Student's t -test and Wilcoxon tests; see SI.2) between the total and non-reactive fractions for the
194 Arve, Fier and Guiers Rivers. Indeed, the differences between the total and non-reactive Al
195 concentrations were lower than 5% for the three tributaries. For Cr, the total fraction was characterized
196 by approximately 10% higher concentration values than in the non-reactive fraction for all tributaries.
197 For Ti, calculations showed little difference in concentrations between the two fractions (lower than
198 2% for the three tributaries). For Ba, Li and V, the differences between the two fractions were
199 significant, but total fraction concentrations were less than 20% higher than the non-reactive fraction
200 concentrations for all tributaries. These six metals (Al, Ba, Cr, Li, Ti, V) generally had low reactivity
201 in the Rhône tributaries, which is consistent with the literature. For example, aluminium can be used
202 as a reference for calculating the enrichment factor in sediment samples (Ollivier et al., 2011). The
203 enrichment factor reflects the anthropogenic inputs of a given metal in a river system. Given that these
204 anthropogenic inputs have a high reactivity, the enrichment factor could serve as an effective tool for
205 representing the reactive fraction of the SPM. We thus conclude that total Al concentrations can be

206 used in fingerprinting studies in practically any river system. For Ba and V, Ollivier et al. (2011)
207 calculated an enrichment factor of 1.02 ± 0.16 and 1.15 ± 0.12 , respectively, in Rhône River SPM,
208 which means that Ba and V have low reactivity.

209
210 Second, concentrations of As, Co, Fe, Mg and Ni (group 2, Fig. 2b) were, in general, significantly
211 higher in the total fraction than in the non-reactive fraction, with differences ranging from 20% to 45%
212 depending on focal metal and tributary. This is consistent with results obtained by Ollivier et al.
213 (2011) for Ni analysed in SPM from the Rhône River. Indeed, this study supports its classification as
214 moderate reactivity property (proportion of 70% of the non-reactive fraction) with an enrichment
215 factor determined as 1.47 ± 0.3 .

216 Finally, concentrations of group-3 metals (Cd, Cu, Mn, Pb, Sr and Zn; Fig. 2c) were significantly
217 higher in the total fraction than in the non-reactive fraction. The difference ranged from 50% to 80%
218 depending on metal and tributary considered. Ollivier et al. (2011) also studied the reactivity of Pb,
219 among other properties, in the SPM collected at the Arles station, located at the outlet of the Rhône
220 River watershed. The results obtained from the calculation of Pb enrichment factor (3.89 ± 1.03)
221 showed that Pb is influenced by anthropogenic inputs (Ollivier et al., 2011). We therefore conclude
222 that Pb is not conservative in the total fraction. These studies showed the same trends for Cd and Zn.
223 Ollivier et al. (2011) calculated enrichment factors of 2.24 ± 1.08 for Cd and 3.32 ± 0.92 for Zn,
224 meaning that these two metals are influenced by anthropogenic inputs. Cd and Zn are therefore
225 considered non-conservative properties.

226 3.2. Historical contamination trends and influence on the non-reactive fraction

227 Figure 3 shows the metal concentrations in the two studied fractions as a function of time (from 1984
228 to 2018) in the sediment core sampled in the Upper Rhône River. Overall, similar sedimentary profiles
229 are observed for metals of group 1 and 2, for both fractions, with more differences in concentration
230 levels between the two fractions for group 2 (from 37 % to 50%) compared to group 1 (from 2% to
231 20%). Properties of group 3 differ from group 1 and group 2 by different temporal trends. Indeed,
232 there are peaks that appear at the top or bottom of the core for the total fraction for metals of group 3
233 (e.g. Cd, Cu and Zn). In general, in the total fraction of metals from group 1 and 2, concentrations of
234 Al (mean of $47.4 \pm 1.7 \text{ g kg}^{-1}$), Ba ($249 \pm 10 \text{ mg kg}^{-1}$), Li ($47.1 \pm 2.1 \text{ mg kg}^{-1}$), V ($73.1 \pm 3.1 \text{ mg kg}^{-1}$),
235 As ($10.7 \pm 1.3 \text{ mg kg}^{-1}$), Co ($12.6 \pm 0.6 \text{ mg kg}^{-1}$), Fe ($27.9 \pm 1.5 \text{ g kg}^{-1}$) and Mg ($10.9 \pm 0.8 \text{ g kg}^{-1}$) did
236 not show any general decreasing or increasing temporal trend over the study period (1984–2018). For
237 the non-reactive fraction, concentrations of Al, Ti, Ba, Cr, Li and V (group 1) and As, Co, Fe, Mg and
238 Ni (group 2) followed a similar trend to total fraction concentrations. On the contrary, Cd, Cu, Pb, Sr
239 and Zn (group 3) did not show temporal trends, except for Cd that tended to decrease from the bottom
240 to the top of the core, but not as much as in the total fraction.

241 Another distinctive feature observed on the sedimentary profiles of the total and non-reactive fractions
242 is the presence of a peak of concentration in 1994 for metals of the group-1 and 2 (Fig. 3a and 3b) that
243 is not present for the group-3. These higher concentrations in 1994 may be explained by a generalized
244 flood on the Upper Rhône tributaries in October 1993 and January 1994 (DREAL, 2011). Some of the
245 metals of group 3, such as Cu and Zn, showed a significant increase in total concentrations in the
246 deepest layers (192 mg kg⁻¹ and 258 mg kg⁻¹, respectively, in 1987). Cadmium showed three peaks in
247 1988 (1.13 mg kg⁻¹), 1997 (0.81 mg kg⁻¹) and 2008 (0.64 mg kg⁻¹). These concentrations for Cu, Zn
248 and Cd are higher than the geochemical background values (in the order of 0.51-28.8 mg kg⁻¹, 11.7-
249 116.7 mg kg⁻¹ and 0.05-0.30 mg kg⁻¹, respectively) reported by Dendeviel et al. (2020) for the Rhône
250 River. Concentrations of Mn and Sr showed an increasing trend from 1984 to 2018 with a peak in
251 2014 and 2016 for Mn (1173 mg kg⁻¹ and 1204 mg kg⁻¹, respectively) and in 2011 for Sr
252 (434 mg kg⁻¹). Lead concentrations remained relatively stable, varying by 10% (25.1 ± 2.5 mg kg⁻¹)
253 over time with no marked peaks. These values are similar to the Pb concentrations measured in a
254 sediment core (1965–2007) collected from the same site in 2008 (concentrations between 16 and 28
255 mg kg⁻¹; Dendeviel et al., 2020). In contrast, the non-reactive concentrations of Cd (0.044 ± 0.033 mg
256 kg⁻¹), Cu (8.45 ± 4.28 mg kg⁻¹), Mn (102 ± 27 mg kg⁻¹), Sr (54.7 ± 13.4 mg kg⁻¹) and Zn (38.8 ± 5.1
257 mg kg⁻¹) were low and did not show any peaks (Fig.3c). The non-reactive concentrations were more
258 stable over time compared to total concentrations due to the exclusion of the reactive part of the SPM
259 by HCl extraction. The notion of conservatism is more important when tracing historical inputs than
260 contemporary SPM sources. A higher reactive fraction proportion of As (50%), Cd (93%), Cu (86%),
261 Fe (48%), Pb (86%) and Zn (69%) was found in sediment core samples than in SPM samples (24%,
262 60%, 69%, 28%, 69% and 55% for As, Cd, Cu, Fe, Pb and Zn, respectively). This is consistent with
263 the literature, which highlights that metals in the sediment core are subject to temporal changes driven
264 by diagenesis and the evolution of contaminants inputs, resulting in peaks in metal concentrations over
265 time (D'Haen et al., 2012; Collins et al., 2017).

266 Despite this shift of reactivity between SPM and sediment core samples, metals follow the same
267 pattern and agree with the proposed classification according to their low, moderate or high reactivity.
268 For example, for group 1 (Fig. 3a), taking into account all layers of the core, the Al and Ti
269 concentrations of the non-reactive fraction accounted on average for 92% and 98%, respectively, of
270 the total fraction. Similarly, the average proportions of the non-reactive fraction in relation to the total
271 fraction for Ba, Cr, Li and V range from 80 to 98% according to metal, and are of the same order of
272 magnitude as what we found in SPM samples. This is consistent with the work of Dendeviel et al.
273 (2020), who determined the natural geochemical background concentrations of some metals in the
274 Upper Rhône River by collecting a sediment core from the pre-industrial period. The measured
275 reference value for Cr, for example, was 63.8 mg kg⁻¹ using total extraction (Dendeviel et al., 2020).
276 Here, the Cr values calculated in the non-reactive fraction were relatively close to the reference value,

277 with an average of $63.7 \pm 5.3 \text{ mg kg}^{-1}$. We thus consider that total Cr concentrations do not undergo
278 significant anthropogenic inputs.

279 For As, Co, Fe, Mg and Ni (Group 2), the average proportion of the non-reactive concentration ranged
280 from 50% to 60%. These values are similar to those reported in SPM, except for As and Fe that
281 showed higher values in the sediment core. For Fe, this proportion averaged 48% in the sediment core
282 vs $28 \pm 3\%$ in SPM for all tributaries. For As, the proportion of the non-reactive fraction for the
283 sediment core samples averaged around 50% vs 70%–80% for the SPM samples, which suggests that
284 As is more reactive in the sediment core than in SPM. This is corroborated by studies from Hung et al.
285 (2009) and Tessier (2015) who highlighted the high reactivity of As under diagenetic conditions. In
286 fact, Smedley and Kinniburgh (2002) notably showed that As is associated with Fe and Mn oxides,
287 which are dissolved under reducing conditions releasing As into the interstitial water of the sediment.

288 Finally, the highly reactive metals such as Cd, Cu, Mn, Pb, Sr and Zn (group 3) showed concentration
289 differences of 70%–95% on average between the total fraction and non-reactive fractions
290 concentrations in the sediment core, following a similar pattern to SPM. In contrast to the observations
291 made for both other groups, not all these metals show the same temporal trend. For example, in the
292 total fraction, Cd, Cu and Zn concentrations increased strongly in the deepest layers (from 1985 to
293 1988) whereas Mn and Sr concentrations were higher in the more recent layers of the core.
294 Conversely, in the non-reactive fraction, all these properties showed a relatively stable evolution of
295 concentrations along the sedimentary profile. Thus, in the total fraction, the significant differences in
296 concentrations of Cd, Cu, Mn, Sr and Zn between the bottom and top of the sediment core suggest
297 metal inputs due to the various anthropogenic pressures over time, whereas the non-reactive fraction
298 helps to reduce the impact of metal reactivity on estimations of source contributions.

299 3.3. Relevance of the non-reactive fraction to optimize tracer selection

300 In order to obtain reliable results on SPM source contributions in the Rhône River using the total
301 fraction, it is essential to take into account the conservatism of the tracers and to identify and exclude
302 non-conservative properties from the procedure. This is carried out using the range test (Fig. 4) and
303 can be illustrated by Fig. 2. The results of statistical tests are given in Supplementary Material SI.2,
304 (Table SI.2).

305 In the total fraction, half of the properties (Li, V, Co, Fe, Cd, Cu, Pb and Zn) had significantly higher
306 concentrations in sediment core than in SPM samples. This suggests that these metals are more
307 reactive in the sediment core than in SPM samples. Note too that anthropogenic inputs were not the
308 same 40 years ago as in contemporary SPM, which may explain the difference between sources and
309 sediment core for some of these properties (Cd, Cu, Pb and Zn). In contrast, in the non-reactive
310 fraction, only Co, Li and V still had significantly higher concentrations in the sediment core than in
311 SPM samples. In the non-reactive fraction, As had significantly higher concentrations in the SPM of

312 the Arve River than in the sediment core, whereas there were no differences between the Fier and
313 Guiers Rivers and sediment core samples. Consequently, the non-reactive fraction increased the
314 number of tracers available for the application of the fingerprinting procedure. These observations can
315 be confirmed by the results of the range test.

316 The range test was performed for all studied elements in the total and non-reactive fractions by
317 comparing concentrations in the three main SPM sources with concentrations in each of the sediment
318 core layers (see Eq. 2). According to each sediment core layer (Fig. 4), more elements do not satisfy
319 the range test in the total fraction (from 2 to 7) than in the non-reactive fraction (from 0 to 3),
320 especially in the deepest sediment core layers. All these properties are therefore considered non-
321 conservative and were removed for further analysis. In the total fraction, concentrations of Co, Cd, Li
322 and V were outside the range of the source values for most of the sediment core layers, which is
323 consistent with results illustrated in Figure 2. For Cu, Pb and Zn, only the deepest layers (63-100 cm)
324 did not meet the range test. These results are consistent with the trends illustrated in Fig. 3 showing an
325 increase in Cu and Zn concentrations in the deepest layers. Concentrations of Pb prior to 1997 did not
326 increase as strongly as for Cu and Zn but were nevertheless higher than the range of the source values.
327 For Fe and Mn, the superficial layers (first 20 cm) of the sediment core had total concentrations
328 outside the range test reflecting the concentration increases seen in recent years (2011–2018 for Fe and
329 2014–2016 for Mn) (Fig. 3). Finally, as shown in Fig. 3, the 1994 layer features a concentration peak
330 for almost all the metals studied, which can explain why Al, Ba, Cd, Co, Cu, Fe, Li, Mg, Pb, Ti and V
331 did not pass the range test for the total fraction. In the non-reactive fraction, the number of metals that
332 do not satisfy the range test was significantly lower (Fig. 4). For the deepest layers of the sediment
333 core, there are less metals that are out of the range test for the non-reactive fraction than for the total
334 fraction (Fig. 4). Moreover, for the 1994 layer, there were fewer metals outside the range test with the
335 non-reactive fraction (7) than the total fraction (11). Nonetheless, as suggested in the literature
336 (Collins et al., 2017), we decided to remove the 1994 layer from further methodological steps in order
337 to keep enough tracers for the mixing model. The higher metal concentrations (Al, Ti, Ba, Cr, Li, V,
338 As, Co, Fe, Mg and Ni) in this 1994 layer of the sediment core may be explained by the widespread
339 flooding of all three tributaries on the Upper Rhône River. Owens et al. (1999) and Manjoro et al.
340 (2017) signalled that using recent SPM to trace source histories likely introduced uncertainties. During
341 such major flood events, the concentrations of some properties can be much higher than those
342 measured in recent SPM (Fig. 2), so the SPM sources in 1994 are not readily associated with the
343 2012–2019 SPM used (Owens et al., 1999; Manjoro et al., 2017). Using the non-reactive fraction
344 helped to overcome the issue of the source sampling period in general, but was not sufficient to
345 compensate for such extreme past events.

346 Hence, considering the total fraction, many properties were removed for the analysis of sediment
347 sources, and some of the selected properties are potentially reactive. There were twice as many

348 properties out of the range test for the total fraction compared to the non-reactive fraction (Fig.4). For
349 As, Cu, Pb, Mn and Zn, the non-reactive fraction was respectively about 49%, 14%, 14%, 16% and
350 28%, respectively, of the total fraction in the sediment core. These metals thus have high reactivity in
351 the total fraction but can still be integrated into the model because they pass the range test. Owens et
352 al. (2016) also highlighted that a non-conservative property could still pass the range test. Based on
353 our range test results, the non-reactive fraction presents a great advantage by increasing the number of
354 tracers available for the study and, thus, reducing the potential reactivity of the tracers.

355 The total fraction includes geological signature and anthropogenic inputs of the tributaries, while the
356 non-reactive fraction reflects only their geological signature and geochemical background. According
357 to the number of metals available following the range test, using the non-reactive concentrations of
358 Cu, Fe, Mn, Pb, V and Zn is more relevant for source fingerprinting in sediment core than total metal
359 concentrations.

360 3.4. Historical SPM source inputs on the Upper Rhône River

361 The mixing model results are presented in Fig. 5 showing the tributary source contributions as a
362 function of time along the sediment core (from 1984 to 2018) derived from concentrations of metals in
363 the total and non-reactive fractions. For the layers from 2012 to 2018, the source contributions of each
364 tributary did not differ between the total and non-reactive fractions. For the layers from 1984 to 2004,
365 source contributions were significantly different between total and non-reactive fractions used in the
366 mixing model, which is explained by the use of more reactive metals for total fraction due to higher
367 anthropogenic inputs (Fig. 4) (Dendeviel et al., 2020).

368 Regarding the uncertainties on the source contributions, the 95% confidence intervals (CI95; SI.3)
369 were low for both fractions and all tributaries. Another way of presenting the uncertainty associated
370 with these results is to calculate the mean absolute error (MAE), as used by Huang et al. (2019). The
371 MAE values associated to source contributions ranged from 0.88 to 0.95 for total fraction and from
372 0.90 to 0.99 for the non-reactive fraction for the 1984-2018 period. A Student's *t*-test concluded that
373 the MAE values were significantly higher, meaning that estimations are more reliable, for the non-
374 reactive fraction than for the total fraction.

375 Our results show that the Arve River was the main SPM source contributor to the Rhône River over
376 the past decade for total fraction for the 2001–2018 period (mean of $52.9 \pm 7.2\%$) and from 1984 to
377 2018 for the non-reactive fraction (mean of $61.6 \pm 8.5\%$). Results using the two fractions showed
378 significant differences, i.e. the Fier River contributed significantly more and the Guiers River
379 significantly less for the total fraction compared to the non-reactive fraction. The Fier River had a
380 mean contribution of $45.3 \pm 10.3\%$ for the total fraction against $22.6 \pm 6.1\%$ for the non-reactive
381 fraction for the 1984–2018 period, while the Guiers River had a mean contribution of $7.3 \pm 4.8\%$ for
382 the total fraction against $15.8 \pm 7.4\%$ for the non-reactive fraction for the same period. The higher

383 contributions of the Fier River compared to the Guiers River for the 2016–2020 and 2012 periods
384 reflects the much higher SPM fluxes of the Fier River than the Guiers River (average annual water
385 flow of $0.18 \text{ Mt}\cdot\text{year}^{-1}$ and $0.07 \text{ Mt}\cdot\text{year}^{-1}$, respectively, for the 2001–2011 period; Launay, 2014). Due
386 to these differences in model results between the two metal fractions, we further investigated the
387 relevance of our mixing model by comparing the estimated contributions of tributary sources using
388 water flow, flood events (2013–2019) and SPM flux data (2012–2019) depending on the data available
389 in the OSR network database (Thollet et al., 2018) (Fig.5).

390 Several major hydrological events occurred during the study period that may explain the observed
391 variations in source contributions. For the 1987 period, the estimations via our mixing model using the
392 non-reactive fraction clearly show a main contribution of the Arve River ($60 \pm 16 \%$) and a lower
393 contribution from the Fier River ($25.5 \pm 17.9\%$) for this period. In contrast, the total fraction equally
394 distributed the SPM inputs between the Arve and Fier Rivers ($46 \pm 21\%$ and $51 \pm 20\%$, respectively).
395 Knowing that a major 100-year flood that occurred in 1987 on the Arve watershed (Grand Bornan),
396 with a water flow of $150 \text{ m}^3 \text{ s}^{-1}$ causing intense erosion of the banks and thus a significant sediment
397 load to the river (DREAL, 2011), this suggests that the contributions of the Arve River were more
398 relevant with the non-reactive fraction.

399 For the 1990 period, as illustrated in Fig. 5, in the total fraction, the Arve and Fier Rivers displayed
400 similar SPM inputs ($49 \pm 23\%$ and $47 \pm 21\%$, respectively) and the Guiers River had a smaller
401 contribution ($3.3 \pm 12.4\%$). In the non-reactive fraction, the Arve River contributed to $75 \pm 18\%$ of
402 SPM inputs, and the Fier and Guiers Rivers contributed far less ($16 \pm 17\%$ and $7.9 \pm 14.3\%$,
403 respectively). This 1990 period recorded a ten-year flood on the three tributaries, with water flows of
404 $643 \text{ m}^3 \text{ s}^{-1}$ for the Arve River, $700 \text{ m}^3 \text{ s}^{-1}$ for the Fier River, and $403 \text{ m}^3 \text{ s}^{-1}$ for the Guiers River
405 (DREAL, 2011; Thollet et al., 2018). Using the available SPM flux data (2015–2019), it is assumed
406 that the average interannual contributions in terms of SPM flux are higher for the Arve River (relative
407 contribution of 65%) than the Fier River (26%), which corroborates that the estimates of contributions
408 using the non-reactive fraction is more relevant than the total fraction.

409 For the Arve River, the contribution peaks observed in the non-reactive fraction in 2003, 2005, 2011
410 and 2012 can be explained by dam flushing operations that occurred downstream in 2003 and 2012
411 (Lepage et al., 2020). For 2005 and 2011, represented by major SPM inputs from the Arve River, a
412 major flood in 2005 with a water flow of $361 \text{ m}^3 \text{ s}^{-1}$ and two floods in 2011 with discharges of 435 and
413 $395 \text{ m}^3 \text{ s}^{-1}$ (Thollet et al., 2018), explain the larger contribution of the Arve River.

414 For the 2010 layer, the main source of SPM was the Arve River, which contributed $45 \pm 22 \%$ and
415 $46 \pm 21\%$ respectively for the total and non-reactive fractions. This higher contribution from the Arve
416 River, compared with the Fier (39% and 18% respectively for the total and non-reactive fraction) and
417 Guiers (17% and 36% respectively for the total and non-reactive fraction) contributions was consistent

418 with the water flow data (with a maximum of $504 \text{ m}^3 \text{ s}^{-1}$) and SPM fluxes. The annual SPM fluxes
419 represent relative contributions of 65%, 26% and 9% of SPM respectively for the Arve, Fier and
420 Guiers Rivers for the 2001–2011 period. In 2010, for the non-reactive fraction, there was also a higher
421 contribution of the Guiers River ($36 \pm 21\%$) than the Fier River ($18 \pm 16\%$). For this period, a 10-year
422 flood occurred on the Guiers River with a maximum water flow of $157 \text{ m}^3 \text{ s}^{-1}$. The Fier River was also
423 subject to a 100-year flood ($505 \text{ m}^3 \text{ s}^{-1}$). Given that coarser particles deposit faster, we suppose that
424 following the Guiers flood, a majority of these coarse particles would have settled and been deposited
425 downstream. Furthermore, the Guiers River has a particle size distribution substantially closer to the
426 grain size of the sediment core, unlike the Fier River, which could further explain the greater
427 contribution of the Guiers River. For the non-reactive fraction, during the period from 2014 to 2016,
428 there was another peak in the contributions of the Guiers River that could be explained by numerous
429 large floods (about 14 floods with a water flow exceeding $100 \text{ m}^3 \text{ s}^{-1}$) that occurred during this period.

430 Using the total fraction, our mixing model results show lower contributions for the Arve River than for
431 the non-reactive fraction. According to the hydrological data, our results suggest that the use of the
432 non-reactive fraction is more relevant to estimate source contributions compared to the total fraction.
433 Thus, the use of this non-reactive fraction holds promising potential for the application of sediment
434 fingerprinting. Indeed, using tracers of the total fraction, the tracer selection step can lead to a set of
435 properties that are reactive or characterized by different concentrations between the deepest layers of
436 the core and the contemporary SPM. Arsenic is the most illustrative example, since non-reactive
437 concentrations in the sediment core represent only 49% of total concentrations, compared to 76% for
438 contemporary SPM.

439 **4. Conclusion**

440 In this study, we used the non-reactive concentrations of metal implemented in a mixing model with a
441 Monte Carlo simulation to estimate historical source contributions in the Upper Rhône River. The
442 results showed that the non-reactive fraction of metals has clear advantages over the total fraction in
443 tracing historical SPM sources through our model. First, for the range test, the non-reactive fraction
444 allows to obtain a greater number of available metals before Kruskal-Wallis test and DFA, and ensures
445 that concentrations of the most reactive metals (i.e. Cd, Cu, Mn, Pb, Sr and Zn) are conservative (i.e.
446 relatively stable and homogeneous concentrations along the sediment profile). Historical SPM
447 contributions modelling showed that using tracers in the non-reactive fraction was more consistent
448 with historical floods and flow data, and with higher MAE (mean absolute error) values than the total
449 fraction, meaning that uncertainties associated with source contributions for the non-reactive fraction
450 were lower and therefore the results were more reliable. In addition, the use of total concentrations of
451 reactive metals could induce unreliable results. Indeed, some of the selected tracers according to the
452 range test were showed to be highly reactive. According to the results of the non-reactive fraction, the

453 SPM from the Arve River mainly constituted the SPM deposited in the Upper Rhône River from 1984
454 to 2018, which was relevant to hydrological data. Thus, our study demonstrated that the non-reactive
455 fraction of metal tracers used in fingerprinting studies can be used in a variety of environmental
456 conditions and at various spatial and temporal scales to provide a robust quantification of sediment
457 sources. Moreover, we believe that the non-reactive fraction might also be more effective than the total
458 fraction in tracing the historical SPM sources in a sediment core collected at the outlet of the Rhône
459 River watershed as undertaken in further studies. Since deposited sediments were subject to changes in
460 physicochemical conditions caused by the salinity gradient and to diagenetic processes, which may
461 alter total metal concentrations, the non-reactive fraction may be a more effective tool in fingerprinting
462 studies than the total fraction.

463

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476

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Figures

Figure 1: Location of the Rhône River watershed in Europe and France (43.3302-4.8455). Map of the suspended particulate matter (SPM) stations of the Rhône Sediment Observatory (OSR) network (squares) and zoom on the study area of the Upper Rhône River and locations of SPM and sediment core sampling sites.

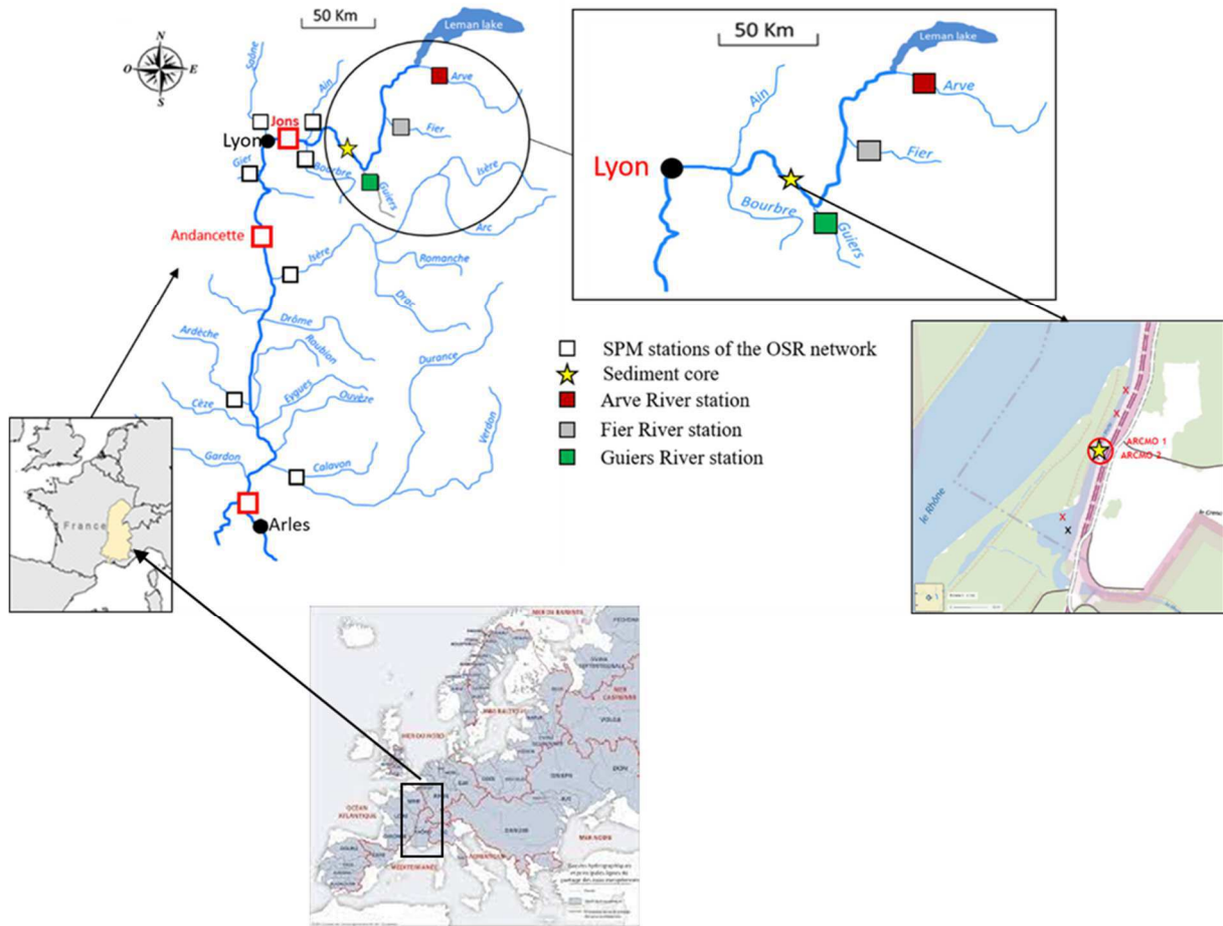
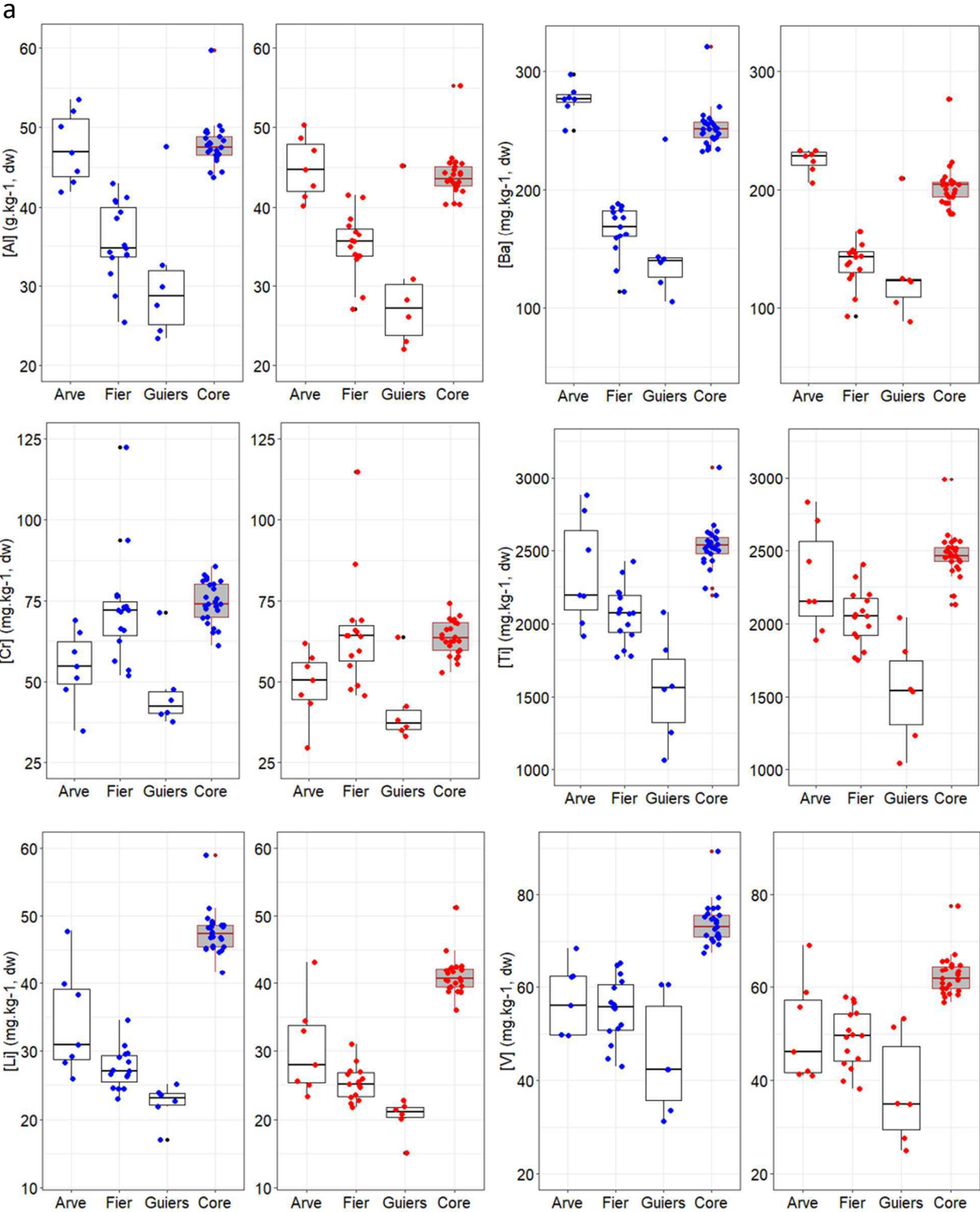
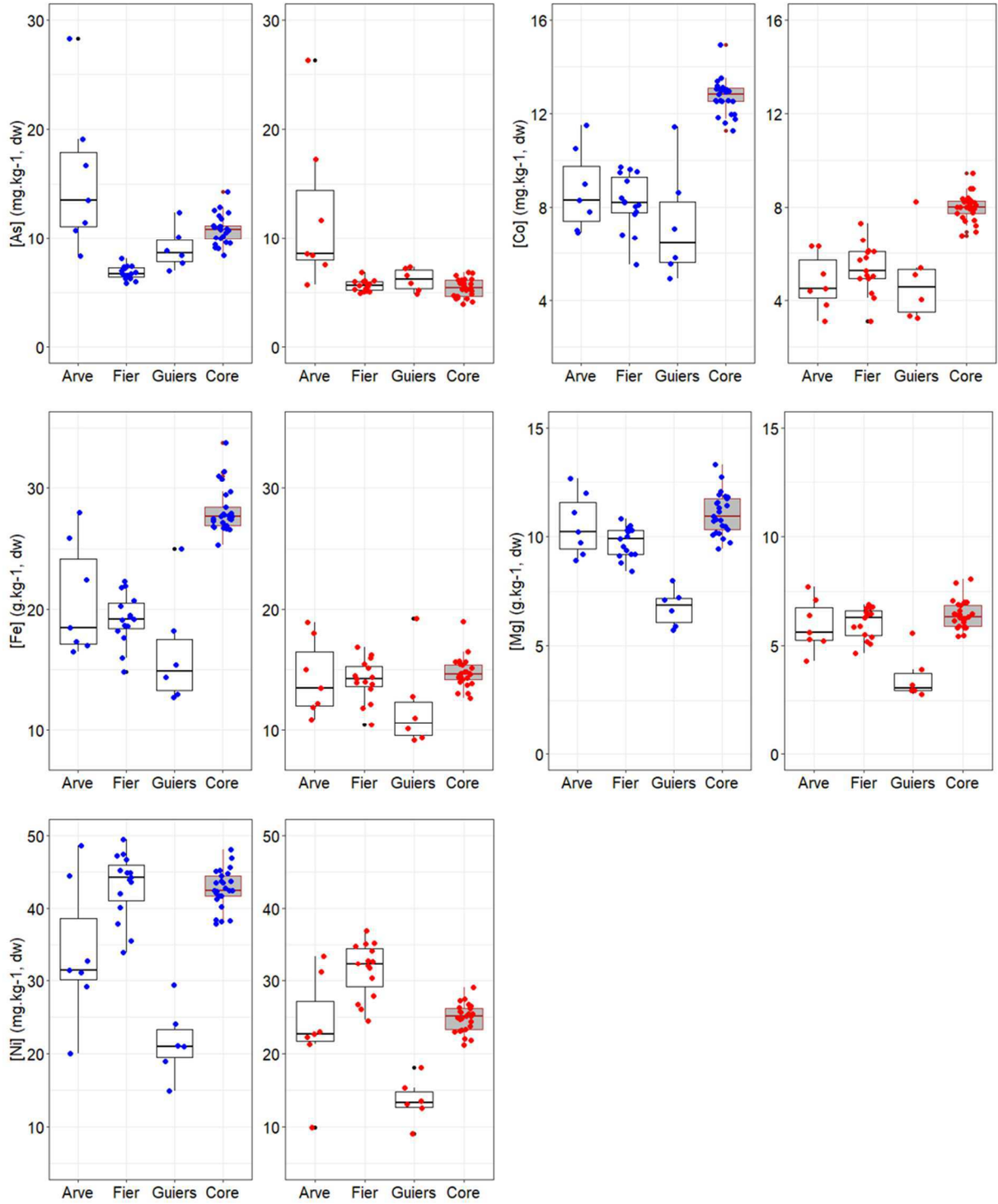


Figure 2: Metal concentrations (particle-size corrected data) in suspended particulate matter of the three tributaries of the Rhône River (Arve, Fier and Guiers Rivers) and sediment core sampled in the Rhône at La Morte site. The three identified reactivity groups represent properties that are not reactive (a), moderately reactive (b), and highly reactive (c). Concentrations in the total (blue dots) and residual (red dots) fractions.



b



C

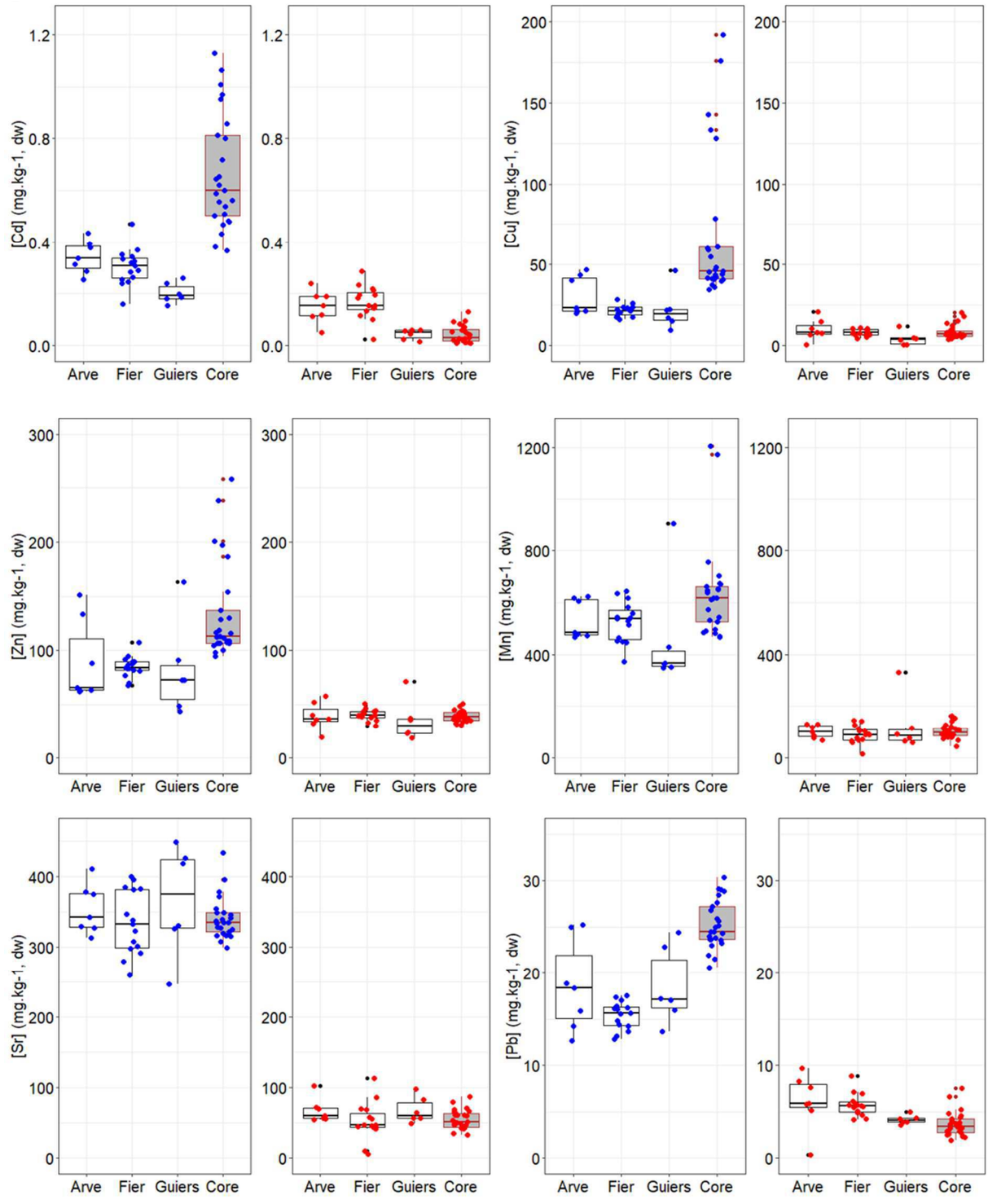
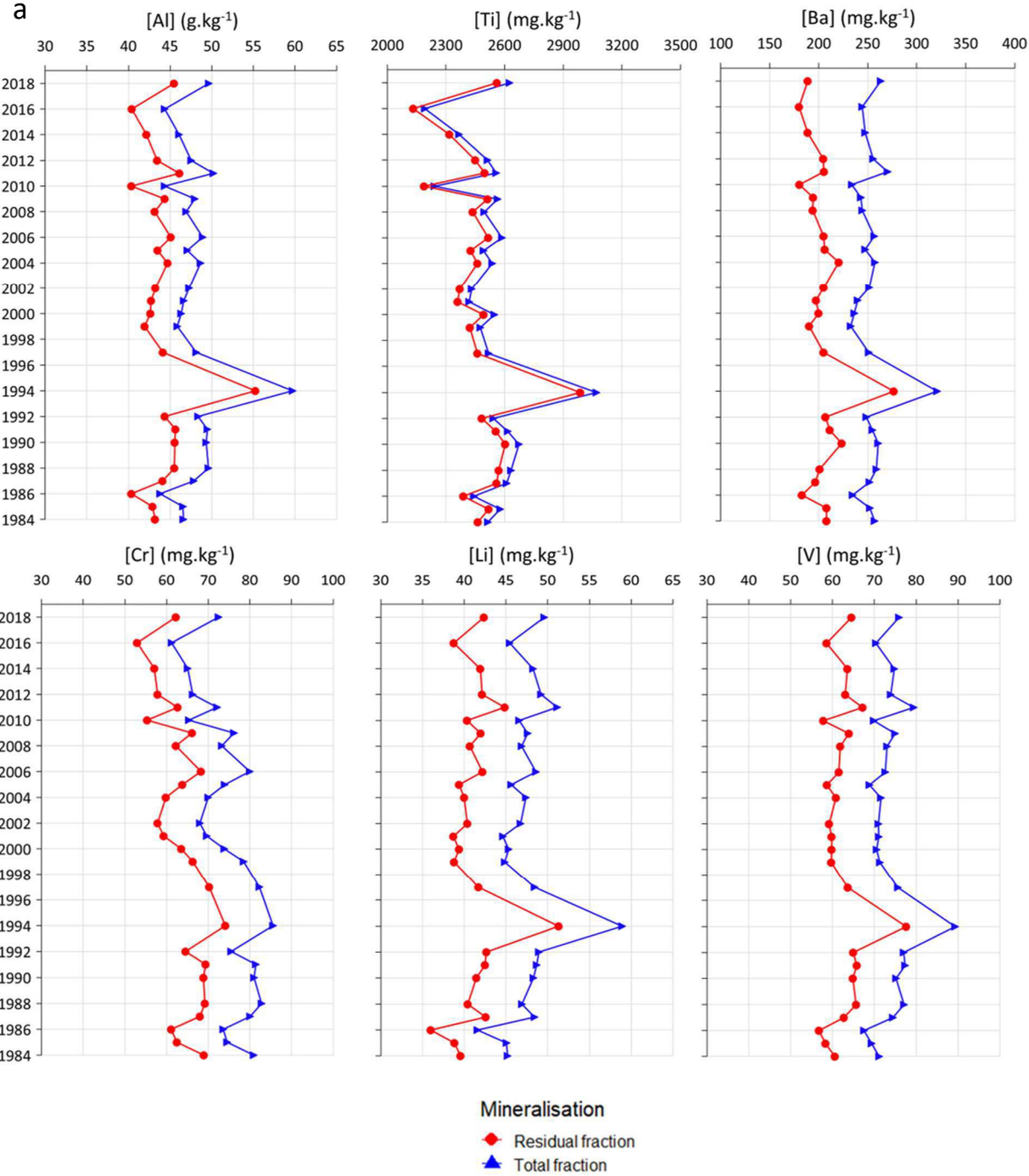
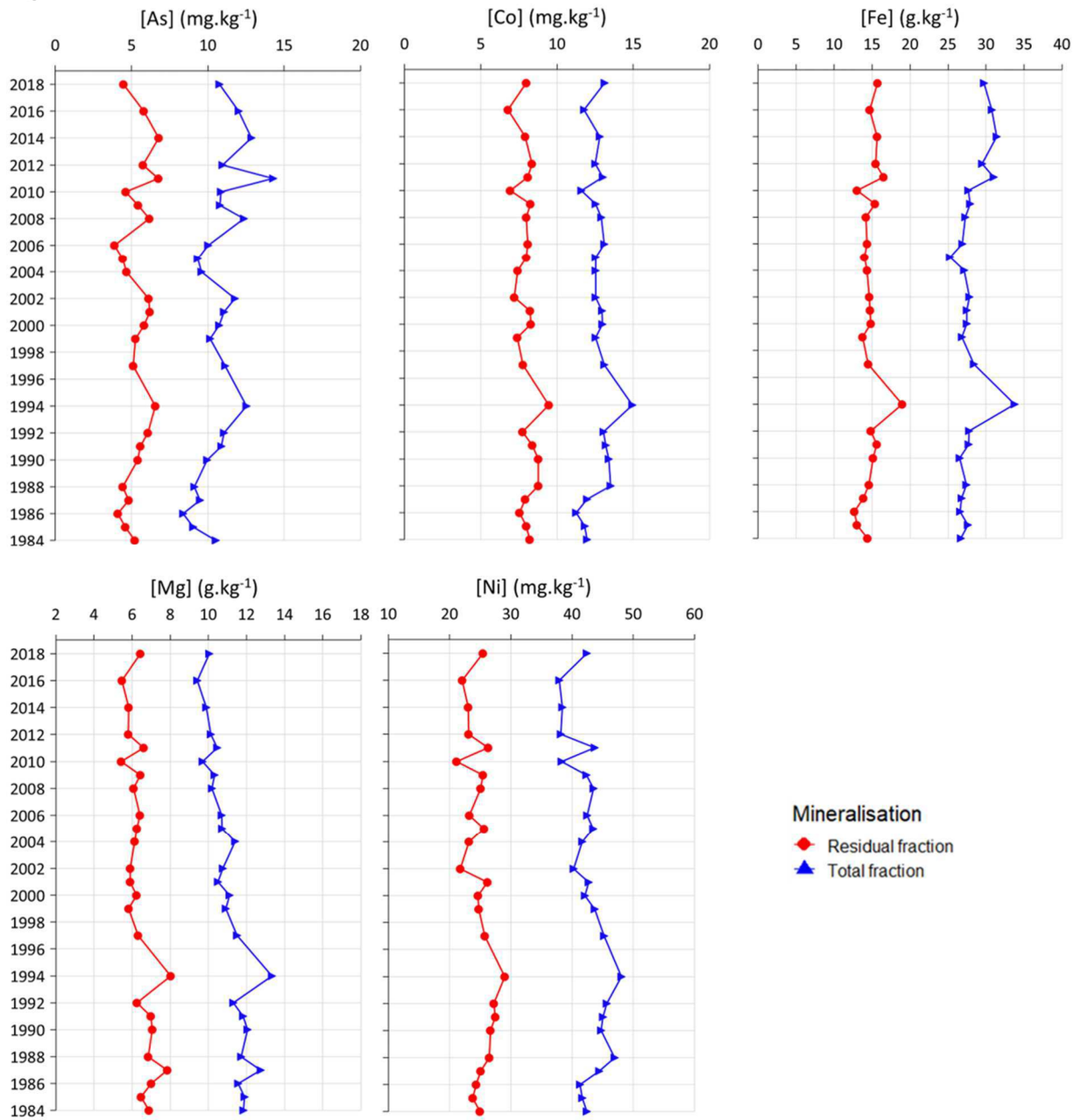


Figure 3: Metal concentrations in the sediment core sampled in the Rhône at La Morte site. The three identified reactivity groups represent properties that are not reactive (a), moderately reactive (b) and highly reactive (c). Example of three behaviours of metals along the sediment core by comparing concentrations in the total (blue line) and residual (red line) fractions.



b



C

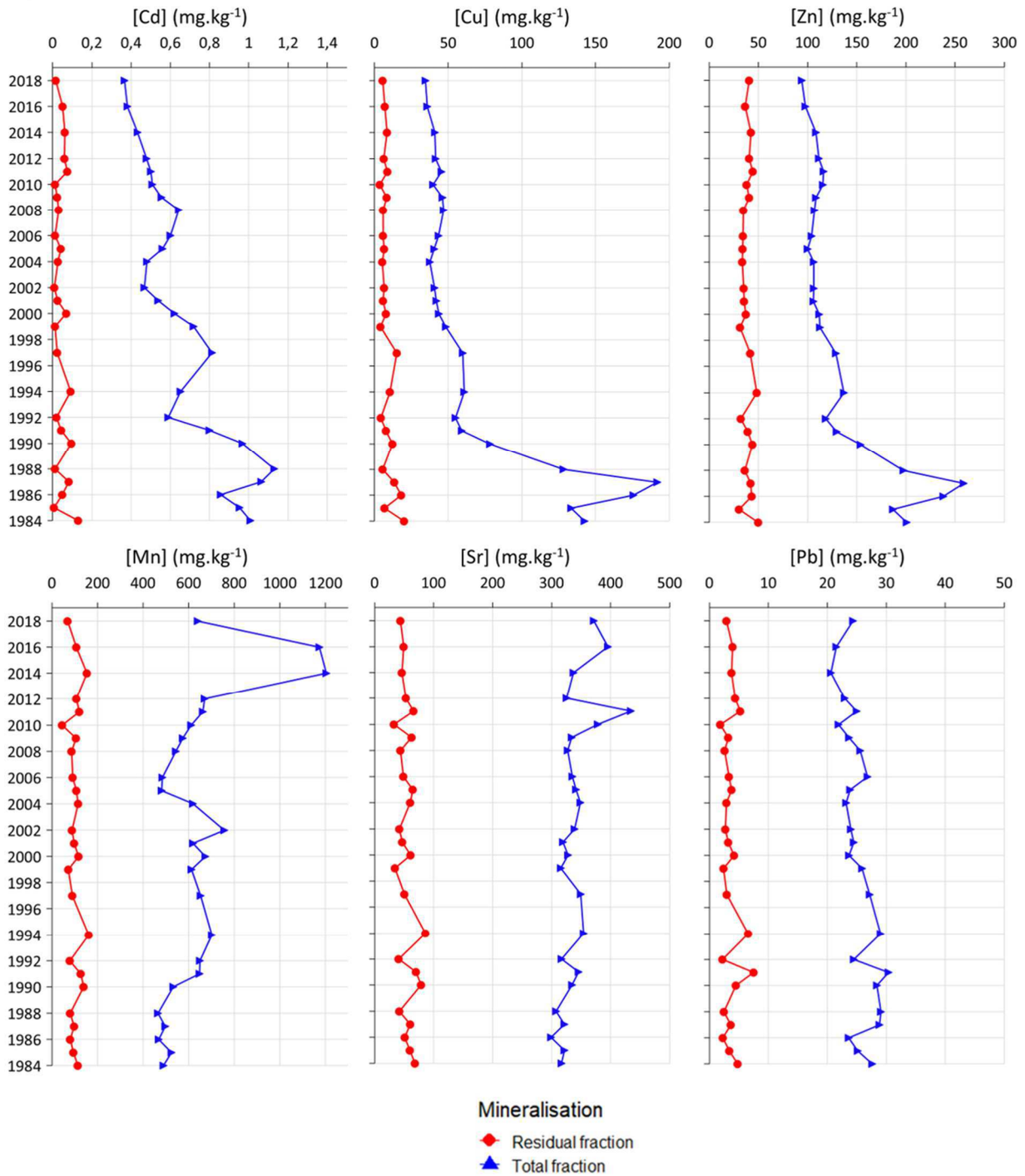
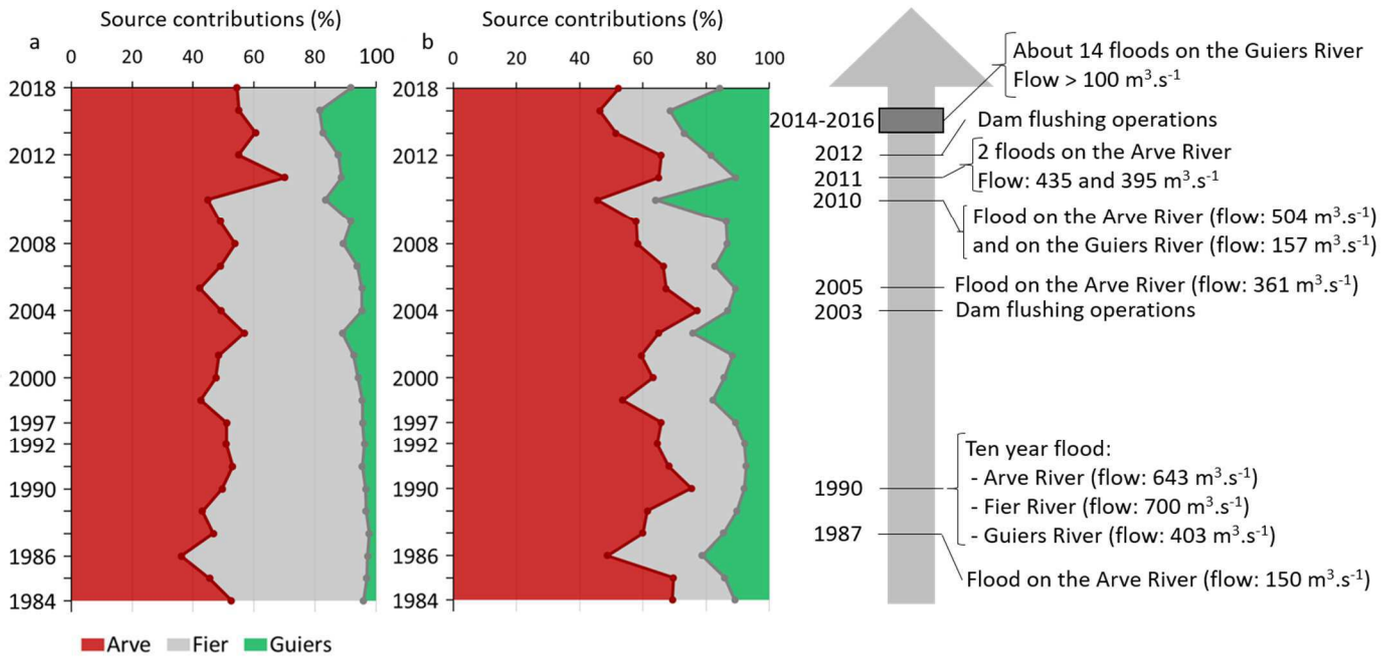


Figure 4: Geochemical properties excluded after the range test for the concentrations of metals in the total (a) and the residual (b) fractions of each layer of the sediment core according to the age of each sediment strata.



Figure 5: Historical relative SPM contributions (in %) of the Arve, Guiers and Fier Rivers as inferred from the selected geochemical tracers in the total (a) and residual (b) fractions of the sediment core.



Tables

Table 1: Statistical results (Range test, Kruskal Wallis test and Discriminant Factor Analysis) for the tracer selection procedure for total and residual fractions of SPM sources. The green “V” to show the geochemical tracers that are retained at each step and red “X” for those that fail. For DFA, the red “x” shows the elements removed from the procedure while the elements retained are characterised by a value representing the discriminatory power of elements.

Properties (corrected concentrations)	Total fraction			Residual fraction		
	Range test	KW test	DFA	Range test	KW test	DFA
Al	V	V	100	V	V	X
As	V	V	100	X	X	X
Ba	V	V	78.6	V	V	78.6
Cd	X	X	X	X	X	X
Co	X	X	X	X	X	X
Cr	V	V	X	V	V	X
Cu	X	X	X	V	X	X
Fe	X	X	X	V	X	X
Li	X	X	X	X	X	X
Mg	V	V	100	V	V	X
Mn	X	X	X	V	X	X
Ni	V	V	96.4	V	V	96.4
Pb	X	X	X	V	V	X
Sr	V	X	X	V	X	X
Ti	V	V	X	V	V	X
V	X	X	X	V	X	X
Zn	X	X	X	V	X	X

Supplementary Material 1: Analytical procedures for suspended particulate matter (SPM) and sediment samples

Particle size distribution was analyzed on fresh SPM samples using a Cilas 1190 particle size analyzer, according to ISO standard 13320 (ISO, 2009) under ultrasound and sample agitation. For the sediment core samples, grain size analysis was performed on a Mastersizer 2000[®] (Malvern Panalytical, Instruments Ltd., Malvern, UK) with a hydro SM small volume wet dispersion unit.

The analysis of geochemical properties involved the quantification of 20 trace and major elements in two fractions: total fraction and reactive fraction. The total fraction was determined after triacid mineralization (nitric acid, hydrochloric acid and hydrofluoric acid, proportioned 1.5 mL HCl 12M, 0.5 mL HNO₃ 14M, 2 mL HF 22M on a heating plate. The reactive fraction was obtained by a soft extraction using hydrochloric acid (1 M) at room temperature (Dabrin et al., 2014). Metals were analyzed in both fractions by inductively-coupled plasma optical emission spectroscopy (ICP-OES, Agilent 720-ES, Agilent Technologies) or triple quadrupole inductively-coupled plasma mass spectrometry (TQ-ICP-MS, Thermo iCAP-TQ, Thermo Fisher Scientific) according to their limit of quantification (Table S1) and the concentration in the samples. Aluminium (Al), barium (Ba), cobalt (Co), chromium (Cr), copper (Cu), iron (Fe), magnesium (Mg), manganese (Mn), strontium (Sr), titanium (Ti), vanadium (V) and zinc (Zn) were analyzed by ICP-OES. Arsenic (As), cadmium (Cd), lithium (Li), nickel (Ni) and lead (Pb) concentrations were determined by TQ-ICP-MS. To control the accuracy of the results, certified reference materials (IAEA-158, marine sediment for total extraction, and LGC-6187, river sediment) were analyzed in triplicate for each analytical series. Coefficient of variation was lower than 11% for each element analyzed (Table S2). In addition, blanks were systematically included in mineralization and analytical series to confirm that the samples were not contaminated during the analytical process.

A Student's *t*-test or Wilcoxon test was used to compare tracer concentrations between total and non-reactive fractions of SPM, concentrations of metals between tributaries, and concentrations of metals between the top and the bottom of the sediment core. Level of significance was 0.05.

Table S1: Limits of quantification of trace (expressed in mg kg⁻¹) and major (expressed in g kg⁻¹) metals according to the analytical technique used in this study (ICP-OES and ICP-MS).

Metals	ICP-OES				ICP-MS	
	LQ_total extraction		LQ_HCl extraction		LQ_total extraction	LQ_HCl extraction
	g kg ⁻¹	mg kg ⁻¹	g kg ⁻¹	mg kg ⁻¹	mg kg ⁻¹	mg kg ⁻¹
Ag					0.025	0.0063
Al	2.5		0.63			
As					0.025	0.0063
Ba		2.5		0.63		
Cd					0.025	0.0063
Co		2.5		0.63		
Cr		5.0		1.25		
Cu		5.0		0.63		
Fe	2.5		0.63			
Li					0.025	0.0063
Mg	2.5		0.63			
Mn		2.5		0.63		
Ni					0.05	0.0125
Pb					0.025	0.0125
Sr		2.5		0.63		
Ti		2.5		0.63		
V		2.5		0.63		
Zn		2.5		0.63		

Table S2: Coefficient of variation (%) associated with metals and analytical techniques (ICP-OES and TQ-ICP-MS).

Metals	Coefficient of variation ICP-OES (%)		Coefficient of variation TQ-ICP-MS (%)	
	Total extraction	HCl extraction	Total extraction	HCl extraction
Al	8.3	8.9	-	-
As	-		9.3	4.2
Ba	7.7	4.1	-	-
Cd	-		9.3	7.9
Co	8.2	7.9	-	-
Cr	9.4	10.0	-	-
Cu	9.1	10.9	-	-
Fe	7.0	8.8	-	-
Li	-	-	6.3	9.5
Mg	8.1	6.7	-	-
Mn	9.4	5.4	-	-
Ni	-	-	7.3	4.9
Pb	-	-	4.5	5.1
Sr	5.4	4.0	-	-
Ti	9.0	8.6	-	-
V	6.6	5.8	-	-
Zn	9.2	8.0	-	-

The sediment core was dated using $^{210}\text{Pb}_{\text{xs}}$ (^{210}Pb in excess) and ^{137}Cs concentrations, which are widely used to date contemporaneous (<100 years) sediment cores (Appleby, 1998). Dry samples of sediment were conditioned in 17 or 60 mL boxing depending on the quantity of sediment available. Each box was placed in vacuum-sealed packages and stored for at least one month before measurement with a germanium detector to ensure the secular equilibrium of the ^{210}Pb necessary to determine the concentration of $^{210}\text{Pb}_{\text{xs}}$ (Morereau et al., 2020).

A peak of ^{137}Cs , related to the atmospheric fallout from Chernobyl in 1986, was observed at a depth of 90 cm and used as a proxy to date the sediment core. According to $^{210}\text{Pb}_{\text{xs}}$, sedimentation rate was estimated to be 2.9 cm year^{-1} factoring in the 31% compaction. Dating was confirmed using additional information input such as flood events (Morereau et al., 2020).

Supplementary Material 2: Results of statistical tests (t test or Wilcoxon test) comparing the concentrations of metals (1) for each tributary (Arve, Fier and Guiers Rivers) and for the sediment core between both analytical fractions and (2) between each tributary and the sediment core for both fractions.

Table SI.1: The green “V” show the significant differences between the concentrations of metal measured in the total and residual fractions ($p < 0.05$) and the red “X” for those that show no significant differences.

Metals	Arve River	Fier River	Guiers River	Sediment core
Al	X	X	X	V
As	X	V	V	V
Ba	V	V	X	V
Cd	V	V	V	V
Co	V	V	X	V
Cr	X	X	X	V
Cu	V	V	V	V
Fe	V	V	X	V
Li	X	V	X	V
Mg	V	V	V	V
Mn	V	V	V	V
Ni	V	V	V	V
Pb	V	V	V	V
Sr	V	V	V	V
Ti	X	X	X	V
V	X	V	X	V
Zn	V	V	V	V

Table SI.2: Results of statistical tests (t test or Wilcoxon test) comparing D50 and concentrations (particle size corrected) of properties in the three sources (Arve, Fier and Guiers Rivers) and the sediment core. The green “V” to show the significant differences between samples ($p < 0.05$) and red “X” for those that show no significant differences.

Properties	Total fraction			Residual fraction		
	Arve vs Core	Fier vs Core	Guiers vs Core	Arve vs Core	Fier vs Core	Guiers vs Core
D50	X	V	X	X	V	X
Al	X	V	V	X	V	V
As	X	V	V	V	X	X
Ba	V	V	V	V	V	V
Cd	V	V	V	V	V	X
Co	V	V	V	V	V	V
Cr	V	X	V	V	X	V
Cu	V	V	V	X	X	V
Fe	V	V	V	X	X	V
Li	V	V	V	V	V	V
Mg	X	V	V	V	X	V
Mn	X	V	V	X	X	X
Ni	X	X	V	X	V	V
Pb	V	V	V	V	V	X
Sr	X	X	X	X	X	X
Ti	X	V	V	X	V	V
V	V	V	V	V	V	V
Zn	V	V	V	X	V	X

Supplementary Material 3: Results of statistical tests comparing the concentrations of metals between tributaries.

The green “V” show the significant differences between the concentrations of metal measured in the total and residual fractions ($p < 0.05$) and the red “X” for those that show no significant differences.

Elements	Total fraction			Non-reactive fraction		
	Arve vs Fier River	Arve vs Guiers River	Fier vs Guiers River	Arve vs Fier River	Arve vs Guiers River	Fier vs Guiers River
Al	V	V	X	V	V	X
As	V	V	V	X	X	X
Ba	V	V	X	V	V	X
Cd	X	V	V	X	V	V
Co	X	X	X	X	X	X
Cr	V	X	V	V	X	V
Cu	X	X	X	X	X	X
Fe	X	X	X	X	X	V
Li	X	V	V	X	V	V
Mg	X	V	V	X	V	V
Mn	X	X	V	X	X	X
Ni	V	V	V	V	V	V
Pb	X	X	X	X	V	V
Sr	X	X	X	X	X	X
Ti	X	V	V	X	V	V
V	X	X	V	X	X	V
Zn	X	X	X	X	X	X

Generally, among the three sources, the lowest concentrations in the total fraction for most of the metals, such as Al, Ba, Cd, Cr, Li, Mg, Ni and Ti, were measured in Guiers samples. The highest mean concentrations were found in the Arve River for Al ($47.5 \pm 4.5 \text{ g kg}^{-1}$), As ($15.4 \pm 6.7 \text{ mg kg}^{-1}$) and Ba ($276 \pm 14 \text{ mg kg}^{-1}$), and in the Fier River for Cr and Ni concentrations ($72.5 \pm 17.2 \text{ mg kg}^{-1}$ and $43.1 \pm 4.5 \text{ mg kg}^{-1}$, respectively). Total concentrations for Cd, Li, Mg and Ti are significantly different between the Guiers and Arve and between the Guiers and Fier but not significantly different between the Arve and Fier Rivers. Concerning the other metals (Co, Cu, Fe, Mn, Pb, Sr, V and Zn), the total concentrations do not display significant differences between the three tributaries.

In the non-reactive fraction, the Guiers River had the lowest concentrations for the same metals (Al, Ba, Cd, Cr, Li, Mg, Ni and Ti) as for the total fraction, except for Pb. Concentrations of Pb in the Guiers River were significantly lower from the concentrations in the other two tributaries. The highest mean concentrations in the non-reactive fraction were found in the Arve River for Al ($45.0 \pm 3.8 \text{ g kg}^{-1}$), As ($12.2 \pm 7.2 \text{ mg kg}^{-1}$) and Ba ($224 \pm 10 \text{ mg kg}^{-1}$) and in the Fier River for Cr ($65.2 \pm 17.1 \text{ mg kg}^{-1}$) and Ni ($31.5 \pm 3.7 \text{ mg kg}^{-1}$), this following the same pattern as for the total fraction. For Cd, Cu, Fe, Li, Mg, Pb, Ti and V, the concentrations in the non-reactive fraction were significantly different between the Guiers and Arve Rivers and between the Guiers and Fier Rivers but not

significantly different between the Arve and Fier Rivers. There were no significant differences in concentrations of Co, Mn, Sr and Zn between the three tributaries.

Supplementary Material 4: The enrichment factor (EF) values and standard deviation (SD) associated obtained for selected trace and major elements in SPM of the Rhône River.

Matrix	Elements	EF (study period)	SD	River systems	References
SPM	Ba	1.02	0.16	Rhône River	Ollivier et al (2011)
	V	1.15	0.12		
	Ni	1.47	0.3		
	Pb	3.89	1.03		
	Cd	2.24	1.08		
	Zn	3.32	0.92		

The EF was calculated according to following equation (according to Ollivier et al. (2011)):

$$EF = (C_{ie}/[Al]_e)/(C_{ir}/[Al]_r)$$

where C_{ie} is the concentration of element i in SPM/sediment samples, $[Al]_e$ is the aluminium (Al) concentration in SPM samples, C_{ir} is the concentration of element i in a reference sample and $[Al]_r$ is the Al concentration in a reference sample. Concentrations of Al are used as reference since they are not influenced by anthropogenic inputs (except for sites with bauxite extraction). In addition, aluminium is used as a particle size proxy, which allows the values to be compared without introducing the bias of grain size variation.

Supplementary Material 5: Comparison of uncertainties (expressed as 95% confidence interval: CI95%) associated with the mixing model output for the geochemical tracers selected in the total fraction with uncertainties calculated for the geochemical tracers selected in the residual fraction for SPM from the three tributaries (Arve, Fier and Guiers Rivers).

