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Reconstruction of historical suspended particulate matter contributions of Rhône River tributaries to the Mediterranean Sea

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Keywords: sediment fingerprinting, trace metals, non-reactive fraction, sediment core, floods

Highlights:

Reconstruction of historical SPM inputs of the Rhône River to the Mediterranean Sea Influence of past anthropogenic inputs on source contribution modelling

Non-reactive fraction successfully estimated past SPM inputs

Geochemical (mixing) model results are consistent with documented past flood events

1 **1. Introduction**

2 An excess of suspended particulate matter (SPM) in rivers can cause environmental and economic 3 damages (e.g., clogging spawning beds, degrading water quality or filling reservoirs; Navratil et al., 4 2012; Torres-Astorga et al., 2018). Since the 1970s, sediment fingerprinting approaches have been 5 widely used to identify the SPM sources. Most fingerprinting studies in the literature use geochemical 6 tracers, such as trace and major elements, to assess contemporary SPM inputs in rivers. However, few 7 studies have focused on historical sediment inputs over a larger time-scale using the fingerprinting approach on sediment cores, which can provide retrospective information on past hydrological events 8 9 (floods) or modifications of SPM fluxes, such as dam construction, bank stabilization or revegetation

(Collins et al., 1997; Navratil et al., 2012). Manjoro et al. (2017) studied several methodological factors 10 11 related to the work on sediment cores, such as the spatial representativeness of a sediment core, the 12 number of tracers selected and the optimal number of model iterations, on model estimations of source contributions. Other studies investigated the application of geochemical (Collins et al., 1997; Pulley et 13 al., 2015), colorimetric (Pulley et al., 2018), magnetic (Pulley et al., 2015) and radiometric (Pulley et 14 15 al., 2015) tracers for estimating source contributions of contemporary SPM in a sediment core; this was 16 done for periods ranging from a few decades to 250 yr. The main objectives of fingerprinting studies 17 applied on sediment cores are to investigate changes in SPM sources over time (Collins et al., 1997; 18 Manjoro et al., 2017). The results were interpreted by comparing estimates of the historical source 19 contributions against information related to land-use changes (Huang et al., 2019), reservoir construction (Gateuille et al., 2019), implementation of actions to reduce soil erosion (Wang et al., 2018), or available 20 hydrological data (Navratil et al., 2012). 21

22 The reconstruction of historical SPM inputs into a river generally implies measurements of tracers in 23 contemporary SPM used as source samples. However, Pulley et al. (2015) is one of the few studies that reported the risk of using non-conservative tracers (e.g., changes in organic matter content, particle size 24 25 selectivity, geochemical and magnetic reactions along the sedimentary profile) to assess historical 26 source contributions in sediment cores. Begorre et al. (2021) recently showed that the main issue with 27 using fingerprinting approaches on sediment cores is the non-conservative behaviour of the geochemical tracers (i.e., total trace and major elements) because (i) tracers undergo diagenetic processes that modify 28 29 their partitioning and concentrations in sediment layers, and (ii) tracers have been largely modified since the mid-1970s, notably because of changes in anthropogenic inputs (Audry et al., 2004; Dhivert et al., 30 31 2016). The proposed alternative was to use the non-reactive fraction to limit the effects of nonconservative behaviour of the selected tracers. One advantage of this novel approach is that it increases 32 33 the number of conservative tracers available after passing the range test (Begorre et al., 2021).

Focusing on the Rhône River, historical sediment contamination by polychlorobiphenyls and metals
(Mourier et al., 2014; Dendeviel et al., 2020) and contributions from contemporary sources of SPM
across the river catchment (Zebracki et al., 2015; Dabrin et al., 2021) have previously been documented.

At the scale of the Bléone River, which is a sub-catchment of the Durance River (a tributary of the 37 38 Rhône River), Navratil et al. (2012) reconstructed the geological sources of SPM over time via the 39 analysis of a sediment core study using radionuclide and geochemical tracers. However, to date, no study has documented historical SPM contributions of the main Rhône River tributaries to the river 40 outlet. Zebracki et al. (2015) quantified the contributions of three main SPM sources, which included 41 42 tributaries grouped according to hydrological characteristics (Cévenol, Oceanic and Mediterranean rivers), and used radionuclide data to show that the Durance River is the main relative contributor of 43 44 SPM at the outlet of the Rhône River. Interest in the study of the sources of sediments on the Rhône River is based on the major sedimentary contributions of the Rhône River to the Mediterranean Sea 45 (Delile et al., 2020). Given that fine sediments affect water quality and aquatic biodiversity (Koiter et 46 al., 2013), it is essential to identify sediment sources to reduce sediment inputs to the Mediterranean 47 Sea. Dendeviel et al. (2020) have worked on historical metal contamination along the Rhône River 48 continuum; but, to the best of our knowledge, no studies have been published on historical sediment 49 50 sources to the Mediterranean Sea.

In this context, the main objective of the present study was to highlight the relevance of using tracers in 51 52 the non-reactive fraction to retrace the historical SPM contributions of the main tributaries of the Rhône 53 River basin to its global SPM discharge to the Mediterranean Sea over the last 40 yr. This study is based 54 on the collection and analysis of a sediment core located close to the outlet of the Rhône River basin, which is only connected to the main channel during major flood events, and on SPM samples collected 55 56 within the Rhône Sediment Observatory (OSR) monitoring network, which has been operational since 2012. To overcome the methodological biases resulting from changes in anthropogenic inputs and 57 58 diagenetic processes potentially affecting concentrations of trace and major elements in sediment cores, we applied an original fingerprinting method based on the analysis of the non-reactive fraction of 59 60 elements. This fingerprinting approach was previously used by Begorre et al. (2021) on a sediment core sampled on the Upper Rhône River, but the depth resolution of the sediment core was too limited to 61 investigate sediment inputs at the flood-event scale, in contrast to the sediment core collected 62 63 downstream of the Rhône River basin for the present study.

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65 **2.** Material and Methods

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2.1. Study area: the Rhône River basin

67 The Rhône River is one of Europe's major rivers and the largest supplier of sediments to the Mediterranean Sea, delivering an inter-annual mean of 5.5 Mt yr⁻¹ over the 2008–2018 period (Delile et 68 al., 2020). The basin covers an area of 95,600 km² spanning a broad diversity of geological and climatic 69 70 conditions (Zebracki et al., 2015; Delile et al., 2020). Over the last 40 yr, eight major flooding events 71 were reported at the Beaucaire station close to the outlet of the Rhône River basin: in 1982 (maximum water discharge, $Q_{max} = 8025 \text{ m}^3 \text{ s}^{-1}$, 1990 ($Q_{max} = 5300 \text{ m}^3 \text{ s}^{-1}$), 1993 ($Q_{max} = 9800 \text{ m}^3 \text{ s}^{-1}$), 1994 72 $(Q_{max} = 11,006 \text{ m}^3 \text{ s}^{-1}), 1996 (Q_{max} = 8981 \text{ m}^3 \text{ s}^{-1}), 1997 (Q_{max} = 8020 \text{ m}^3 \text{ s}^{-1}), 2002 (Q_{max} = 9700 \text{ m}^3 \text{ s}^{-1})$ 73 and 2003 ($Q_{max} = 11,500 \text{ m}^3 \text{ s}^{-1}$) (DREAL, 2011). Concentrations and fluxes of SPM and associated 74 75 contaminants are monitored in the Rhône River and its main tributaries under the Rhône Sediment 76 Observatory ("OSR") program (Fig. 1). A monitoring network that has been set up and running since 77 2009 collects a large set of SPM samples and data via stations located across the entire basin (Thollet et 78 al., 2021). The study area includes eleven tributaries located upstream of the sediment core location near 79 the Rhone River outlet (Fig. 1), i.e., the Arve, Fier, Guiers, Ain, Bourbre, Saône, Gier, Isère, Ardèche, 80 Durance and Gardon rivers.

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2.2. Sampling strategy

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2.2.1. Suspended particulate matter source sampling

The SPM samples were collected at five river stations representing the potential SPM sources to characterize geochemical signatures of sediment sources: the Middle Rhône River at the Andancette station (including inputs from seven tributaries, i.e., the Arve, Fier, Guiers, Ain, Bourbre, Saône and Gier rivers), the Isère, Ardèche, Durance and Gardon river stations (Fig. 1). The Andancette station is located on the Rhône River, upstream of its confluence with the Isère River. The other stations are located downstream of each tributary, at a few kilometres upstream from their confluence with the

Rhône River (Thollet et al., 2021). The SPM samples were collected using integrative particle traps 90 91 deployed throughout the year and retrieved every month. In order to collect SPM samples during specific 92 events, such as the Cévenol floods, event-based samplings were carried out using a continuous flow centrifuge (Westfalia KA 2-86-76) or by manually sampling large volumes of water. A more detailed 93 description of the sampling methods can be found in Masson et al. (2018). Overall, 42 SPM samples 94 were collected between 2011 and 2019 from five sources: the Middle Rhône River (n=11), the Isère 95 River (n=14), the Ardèche River (n=6), the Gardon River (n=2) and the Durance River (n=9) (see Table 96 97 1).

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2.2.2. Targeted sediment core sampling

99 To assess the historical SPM inputs from Rhône River tributaries, a 300-cm-long sediment master core 100 was reconstructed by combining seven individual sediment cores collected from two holes (16 cm apart) 101 at different depths at the Mas des Tours site, which is located 54 km upstream of the Mediterranean Sea 102 (43.740000 N; 4.625194 E; Fig. 1), in May 2018 (Morereau et al., 2020). This sampling site was selected 103 because it is only connected to the Rhône River during major floods events. Core drilling was carried 104 out using a Cobra TT percussion driller equipped with a transparent 90-mm diameter PVC liner. Using a XRF core scanner (ITRAX, Cox Analytical Systems, Sweden), 43 layers ranging in thickness from 2-105 106 15 cm were identified. Each layer was subsampled using a ceramic knife to collect several grams of 107 fresh sediment. A subsample was stored for particle size analysis. Other sediment subsamples were freeze-dried, ground, and stored in plastic bags until further analysis. All information about core dating 108 109 are fully reported and discussed by Morereau et al. (2020).

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- 111

2.3. Physicochemical analysis of SPM and sediment samples

The particle size distribution was determined on fresh SPM samples using a Cilas 1190 particle size analyzer under ultrasound and sample agitation, according to ISO standard 13320 (ISO, 2009). For the sediment core samples, the particle size distribution was determined on a Mastersizer 2000© instrument (Malvern Panalytical, Instruments Ltd., Malvern, UK) with a small-volume wet dispersion unit. Tests carried out to compare results obtained by these two instruments showed that there are no significantdifferences between the results.

The analysis of geochemical properties involved the quantification of 20 trace and major elements 118 (metals) in the total and reactive fractions (see supplementary material "Database Rhône River"). As 119 120 highlighted by Dabrin et al. (2021), soft extraction using HCl has been used for many years to identify metals adsorbed on the reactive fraction of SPM, which is mainly present in anthropized systems. The 121 total fraction was determined after triacid mineralization (12 M hydrochloric acid, 14 M nitric acid and 122 123 22 M hydrofluoric acid, respectively proportioned at 1.5 mL, 0.5 mL and 2 mL) on a heating plate. The reactive fraction was obtained by soft extraction using hydrochloric acid (1 M) at room temperature 124 (Dabrin et al., 2014). The difference between the concentrations of the two fractions correspond to what 125 we call a 'non-reactive fraction'. Further details can be found in Begorre et al. (2021). Major and trace 126 127 elements were analyzed in both fractions by inductively-coupled plasma optical emission spectroscopy (ICP-OES, Agilent 720-ES) or triple-quadrupole inductively-coupled plasma mass spectrometry (TQ-128 ICP-MS, Thermo iCAP-TQ) according to their limit of quantification and concentration in the samples. 129 Certified reference materials (IAEA-158, marine sediment for total extraction, and LGC-6187, river 130 131 sediment) were analyzed in triplicates for each analytical series to control the accuracy of results. Precision was lower than 11% for each element analyzed. When considering sediment samples, 132 precision of the analysis by ICP-OES was similar between the total and HCl fractions with values in the 133 range of 0.3-9.7% depending on the metal. For analysis by ICP-MS, analytical precision on triplicates 134 135 was lower than 7% for the total fraction against lower than 9% for the HCl fraction. In addition, blanks were systematically included in the mineralization and analytical series to confirm that the samples were 136 137 not contaminated during the analytical process.

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139 2.4. Sediment core dating

Sediment core dating was performed according to the procedure detailed in Morereau et al. (2020).
Briefly, the sediment core could not be dated with the traditional markers such as ¹³⁷Cs and ²¹⁰Pbxs

(Appleby, 1998; Foucher et al., 2021). In fact, the Marcoule nuclear facility located along the Rhône 142 River also releases those radionuclides into the waters in proportion that diluted the contributions of the 143 144 watershed and masked the traditional markers (Provansal et al., 2010). For this reason, the dating of this archive was carried out by modelling using the data on the releases from this facility (Morereau et al., 145 2020). Dry samples of sediment were conditioned in 17-mL or 60-mL boxes depending on the quantity 146 of sediment available and were placed in vacuum-sealed packages and stored for at least one month 147 before analysis to ensure the secular equilibrium of the ²¹⁰Pb necessary to determine the concentration 148 of ²¹⁰Pbxs (Morereau et al., 2020). Measurements of gamma emitters were performed with a germanium 149 detector. Dating was confirmed using additional information input, such as the chronology of past 150 151 flooding events. Morereau et al. (2020) showed that, because of the location of the coring site, the deposited sediments mainly correspond to flood deposits. Dating of the sediment core showed that the 152 43 layers included sediments deposited from 1981 to 2017. The mean apparent sedimentation rate was 153 estimated at 7.8 cm yr⁻¹. 154

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2.5. Statistical analysis and fingerprinting procedure

157 2.5.1. Data treatment before implementation in the geochemical (mixing) model

Trace and major element concentrations in the total or residual (non-reactive) fraction were corrected for differences in particle size between the SPM sources and the sediment core. The correction method applied was the method described by Gellis and Noe (2013) and implied a particle size difference between SPM from tributaries (source samples) and the sediment core (target samples). As illustrated in Supplementary Information SI.1, the SPM of the Ardèche and Durance rivers had significantly different D₅₀ values (median value of the particle size distribution) than the sediment core samples. For metals that correlate positively with D₅₀ values, concentrations were corrected according to Eq. (1):

165 $C_f = C_i - [D50_{(S)} - D50_{m(Sed)}] \times p$ (1)

where C_f is corrected concentration of tracer i, C_i is initial concentration of tracer i in source s, $D50_{(S)}$ is median particle size value of source s, $D50_{m(Sed)}$ is average D_{50} value for all target sediment samples, and p is slope of the regression line.

169 Trace and major elements integrated in the geochemical model were selected by a three-step

170 procedure: a range test to keep conservative metals (Eq. (2)), a Kruskal-Wallis test to remove

171 redundant elements, and a discriminant factor analysis (DFA) to determine the signature that ensures

172 optimal source discrimination. Results of this tracer selection process are reported in Table 2.

173
$$[\min(C_{is})]_{mean} - 0.10 \times [\min(C_{is})]_{mean} < C_i < [\max(C_{is})]_{mean} - 0.10 \times [\max(C_{is})]_{mean}$$
 (2)

174 where C_{is} is concentration of tracer i in source s, and C_i is concentration of tracer i in sediment core

175 layers. The concentration of tracer i in the sediment core must lie within the source range represented

by the minimum and maximum concentrations of tracer i in sources for which a 10% error is accepted.

177 The distribution-mixing model coupled to Monte Carlo simulation resolved Eq. (3) to estimate the178 source contributions with their associated uncertainties from the selected tracers for each layer of the

sediment core. Uncertainties associated with the source contributions were calculated based on 95%confidence interval and mean absolute error (MAE; Eq. 4).

181
$$C_i = \sum_{l=1}^{1000} \sum_{s=1}^{n} ((P_s \times C_{is})/1000)$$
 (3)

where Ps is percentage contribution from SPM tributary s, Cis is concentration of tracer i in tributary s,
n is number of tributaries, and Ci is tracer concentration in the target sediment samples (Hughes et al.,
2009; Haddadchi et al., 2013). Note that this model is based on two conditions: the source
contributions should be between 0 and 100%, and the sum of the contributions is equal to 100%
(Hughes et al., 2009; Navratil et al., 2012; Collins et al., 2017).

187
$$MAE = 1 - (\sum_{i=1}^{m} |[C_i - (\sum_{s=1}^{n} P_s \times C_{si})]/C_i|/m)$$
 (4)

188 where m is number of properties. If the MAE is greater than 0.85, then the model results are reliable.

189

2.5.2. Statistical tests to investigate metal reactivity in the sediment cores

To investigate metal reactivity in the sediment profile, a Student's *t*-test or Wilcoxon test was used to
compare tracer concentrations between total and non-reactive fractions of SPM, concentrations
between tributaries, and concentrations of metals between top (1991–2017 period) and bottom (1981–
1930 period) layers of the sediment core. The level of significance used in statistical tests was set at a
p-value < 0.05.

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3. Results and Discussion

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3.1. Reactivity of tracers in SPM and sediment core from the Rhône River basin

Total and non-reactive metal concentrations in SPM for each studied tributary are reported in Fig. 2. 197 The two layers corresponding to sediment deposited in 2014 and 2017 were removed from further 198 analysis because the non-reactive concentrations of 7 out of the 12 metals analyzed were below the 199 200 limit of quantification, which could narrow the applicability of the fingerprinting method by decreasing the number of available tracers. The study reported here therefore considers the 1981–2013 201 period. To apply a robust fingerprinting method, it is necessary to select metals with low or moderate 202 reactivity in SPM for all tributaries and in the sediment core. To investigate the spatial (contemporary 203 204 SPM) and temporal (sediment core) variability in metal reactivity, we assessed metal reactivity according to percentage of reactive fraction to percentage of total fraction (see Table 2). Based on 205 206 these percentages, we categorized the trace and major elements into three groups: low reactivity (< 20%), moderate reactivity (20%–50%) and high reactivity (>50%). The percentages listed in Table 207 208 2 were determined from the mean percentage values of all samples for each tributary. Only two 209 samples were available for the Gardon River, and so we were unable to reach a definitive conclusion 210 on metal reactivity for this tributary.

Begorre et al. (2021) found that all groups of metals in the Upper Rhône River presented the same
reactivity in SPM across all the studied tributaries: Al, Cr, Ti and V displayed low reactivity, Co and
Ni had moderate reactivity, and Cu, Mn and Sr showed strong reactivity. Here we found a globally
similar pattern of results, although the SPM from the Ardèche and Durance rivers had the highest
reactive fractions of most metals compared to other tributaries. In addition, the reactivity group of Ba,

Fe and Zn varied according to the source considered, whereas Al, Co, Cr, Cu, Mn, Ni, Sr, Ti and V 216 were assigned to the same reactivity group whatever the tributary considered. For example, Ba showed 217 218 low reactivity in SPM at the Andancette (Middle Rhône River) and Isère river stations (18% and 19%, respectively), whereas it was moderately reactive in the Ardèche and Durance rivers (32% and 40%, 219 220 respectively). Iron also showed spatial variability in its reactivity, with low reactivity in SPM from the 221 Isère River (19%), but moderate reactivity for SPM collected at the Andancette, Ardèche and Durance 222 rivers (30%–31%). Finally, Zn showed moderate reactivity in SPM from the Isère and Durance rivers 223 (33% and 43%, respectively), whereas it was highly reactive in SPM from the Andancette (56%) and 224 Ardèche (65%) rivers. This higher reactivity of Zn at the Andancette and Ardèche river stations could 225 be explained by Zn pollution caused by numerous anthropogenic activities, i.e., by vineyards and industries along the Middle Rhône River, and by farming and old mining tailings on the Ardèche 226 River (Ollivier et al., 2011; Dendeviel et al., 2020). 227

Metal concentrations in the sediment core displayed different temporal patterns (Fig. 3). Based on 228 229 linear regression (i.e., data not shown), total concentrations of Al, Ti, V, Mn, Ni, Co, Sr and Fe did not show a significant trend from the deepest layers through to the top of the sediment core. In contrast, 230 231 Cr, Cu, Zn and Ba concentrations in the total sediment fraction decreased from 1981 to 2013 (statistically significant linear regression with R²_{Cr}=0.67 and p<0.05, R²_{Cu}=0.65 and p<0.05, R²_{Zn}=0.67 232 and p<0.05, R²_{Ba}=0.78 and p<0.05). In detail, total Cu, Zn, Cr and Ba concentrations before 1990 233 234 differed significantly from concentrations measured after 1990, and they all followed a decreasing 235 trend in the total fraction from 1981 to 1990 (Fig. 3). For Cr and Ba, this temporal trend is also highlighted for non-reactive concentrations, which could mean that Cr and Ba were not influenced by 236 237 past anthropogenic inputs. Conversely, non-reactive concentrations of Cu and Zn did not vary over time and were significantly different from their total concentrations, suggesting that only the reactive 238 239 fraction of Cu and Zn decreased over time (from 1981 to 1990), which could be explained by historical anthropogenic inputs of reactive metals. This decrease in total Zn concentrations is 240 241 consistent with Zn concentrations in the sediment core (Ferrand et al., 2012) and surface sediments from the National Basin Network of the French Water Agency, which highlighted a decreasing trend 242

from 1986 to 1990 (Ferrand et al., 2012). Furthermore, Dendeviel et al. (2020) highlighted (i) that Cu 243 244 and Zn were delivered all along the Rhône River by multiple anthropogenic activities (i.e., vineyards, 245 mining, cable production, a nuclear power plant, the Marcoule reprocessing spent fuel facility) in the 1980s, and (ii) that Cu and Zn concentrations decreased from 1960 to 1990. Moreover, Morereau et al. 246 247 (2020) reported that the Marcoule facility (on the Rhône River, just north of the Durance tributary) 248 released liquid effluents highly contaminated with Cu and Zn until 1990. Based on these observations, 249 we supposed that some metals are more reactive in the deepest sediment core layers (1981–1990) 250 compared to the more recent layers (1991–2013). We therefore scrutinized metal concentrations in the sediment core separately for these two periods, i.e., 1981–1990 and 1991–2013. 251 252 Concentrations of Al, Ti, V, Mn, Ni, Co, Sr, Fe, Cr and Ba measured in the non-reactive fraction coevolved with the total fraction over time, whereas concentrations of Cu and Zn in the non-reactive 253 254 fraction remained stable along the sediment core. This means that the total concentrations of these two elements may be influenced by variable anthropogenic inputs or variable reactivity, which makes them 255 256 unreliable for tracing historical sediment sources. This is supported by our metal reactivity study based 257 on proportions of the reactive fraction, which classified all metals except Zn into the same reactivity 258 groups whatever the sediment core layer considered. Indeed, Zn showed moderate reactivity in the 259 upper part of the core (50%) but was highly reactive in the deepest layers (1981–1990) of the sediment

River, the reactive fraction of Zn was higher before the 1990 layer (80%) compared to the more recent
layers (66% for the 1991–2013 period).

core (63%). Begorre et al. (2021) also showed that, in a sediment core sampled in the Upper Rhône

Metal reactivity could also differ between SPM tributaries and the sediment core. Al, Co, Cr, Cu, Mn, Ni, Sr, Ti and V showed the same degree of reactivity in SPM tributaries and the sediment core, which is consistent with results obtained by Begorre et al. (2021) for the sediment core sampled in the Upper Rhône River. However, Sr showed significant differences between the highest reactive fraction in SPM from the Durance River (89%) and all other stations (~65%). The reactive fraction of Sr determined in the sediment core layers was similar between the top and bottom of the core at values of around 74–76%. These results showed that SPM from the Durance River were characterized by a

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higher Sr reactivity than sediment core layers and SPM from the other tributaries. This could be 270 explained by the high carbonate contents in SPM from the Durance River because of its sedimentary 271 272 basin (Ollivier et al., 2011). Ollivier et al. (2011) showed that Sr is mainly associated with carbonates, 273 which represent the most reactive fraction of particles, meaning that Sr can easily be removed from 274 particles to the dissolved phase under varying physicochemical conditions. They also highlighted that the Rhône River drains mainly carbonate bedrocks, which are significant sources of SPM during major 275 276 flood events, as in the case of the 2001 and 2002 floods (Ollivier et al., 2011). These events 277 correspond to the increase in total concentrations of Sr found, in the present study, in the recent layers 278 of the core (2001–2013) collected at the outlet of the Rhône River. Liu et al. (2013) studied changes in 279 concentrations of metals associated with reducible and carbonate phases in two coastal sediment cores 280 (Taiwan) and found that metal concentrations were correlated to the carbonate content and that Sr concentrations in the carbonate fraction were higher in the recent layers compared to the deepest 281 layers (Liu et al., 2013). Given that Sr has a high affinity with the carbonate fraction in the Rhône 282 River, it is consistent that total Sr concentrations were higher in the recent layers (2001–2013) of the 283 284 sediment core studied here.

285 The reactive fraction of Ba, Fe and Zn also differed between the sediment core and SPM samples at 286 some stations (Table 2). For example, in case of Ba, the reactive fraction measured in SPM from the 287 Isère (19%) and Andancette (18%) stations was significantly different (p<0.05) than the reactive fraction in the sediment core (39% and 43% for the 1981–1990 and 1991–2013 periods, respectively). 288 289 Conversely, the percentages of the reactive fraction of Ba in SPM from the Ardèche (32%) and 290 Durance (40%) rivers were similar to those found in the sediment core. As reported by Kresse et al. 291 (2007), the erosion of sedimentary rocks constituted by Ba-enriched carbonates could explain the higher reactivity of Ba in SPM from the Ardèche and Durance rivers, in comparison to other 292 293 tributaries. Furthermore, comparison of our results against those obtained for the sediment core 294 sampled on the Upper Rhône (Begorre et al. 2021) shows that Ba reactivity increased downstream of the watershed, i.e., from 20% for the Upper Rhône to 39%–43% at downstream sites. Ba-based 295 296 minerals are generally found in the non-reactive fraction of the particles, but under anoxic conditions,

these minerals are dissolved and then precipitate again in the sedimentary deposits (Henkel et al., 297 2012). This may explain, depending on the presence of such minerals, the increase in the total and 298 299 non-reactive concentrations of Ba with depth. The spatial variation in total and non-reactive Ba concentrations from the Upper Rhône (mean of 253 mg kg⁻¹ and 203 mg kg⁻¹, respectively) to the 300 Rhône River outlet (mean of 577 mg kg⁻¹ and 336 mg kg⁻¹, respectively) may be explained by a 301 missing source of dissolved Ba that precipitates under oxic conditions. Values for the reactive fraction 302 303 of particulate Fe were similar between the Middle Rhône (Andancette station), Ardèche and Durance 304 rivers (30–31%) and also similar to those observed in the sediment core layers deposited between 305 1991 and 2013 (32%). The SPM from the Isère River was the only exception, with a lower percent 306 reactive fraction of Fe (19%). Liu et al. (2013) highlighted that Fe is mainly associated with the oxide 307 fraction of particles. However, to validate this assumption in the case of the Rhône River, it would be 308 necessary to have information about the oxide fraction in SPM samples from each tributary. For Zn, a 309 high reactivity was found in SPM from the Ardèche River (65%), which is almost two-fold higher 310 than the lowest value measured for SPM from the Isère River (33%). This Zn reactivity in SPM of the 311 Ardèche River (65%) was similar to the Zn reactivity in the deepest layers of the sediment core (63% for the 1981–1990 period). We assume that because Zn is statistically selected and used in the 312 geochemical model, the high reactivity of Zn would introduce a bias in the contribution estimates for 313 314 the model based on the total fraction.

Note that the metal reactivity investigation can serve to identify the most appropriate tracers for 315 316 reliably estimating source contributions. Owens et al. (2016) highlighted that the range test, which is commonly used to remove non-conservative elements, is not fully reliable. This was explained by the 317 318 fact that (1) even if total concentrations of a tracer remain within the range values of the sources, they can still evolve because of tracer reactivity and move away from initial concentrations (e.g., metal 319 320 precipitation in the mixing zone may lead to higher tracer concentrations), and that (2) some elements were removed from the procedure because of higher concentrations from an unidentified source 321 322 (Owens et al., 2016). In the present study, based on the range test results, metals included in the low and moderate reactivity groups (i.e., Al, Ba, Co, Cr, Fe, Ni, Ti and V) could be used to reliably trace 323

324	sediment sources. However, total concentrations of metals in the third (high reactivity) group (i.e., Cu,
325	Mn, Sr) are not recommended for estimating source contributions. Zn is a distinctive element as it is
326	moderately reactive at the top of the sediment core (1990-2013 period) while it is highly reactive in
327	the deeper layers (1981–1990; Table 2).

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3.2. Historical reconstruction of relative contributions of SPM sources to the Rhône River

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3.2.1. Selection of tracers in the geochemical model

331 The optimal composite signature integrated in the geochemical model was selected using the range test, as detailed in Section 2.5.1. The range test results (Table 3) show that four elements (i.e., Ba, Cr, 332 333 Cu and Ti) were excluded from this procedure for the total fraction, and that Mn, Sr and Zn, which are highly reactive, were kept for further statistical tests (Kruskal-Wallis and DFA). For the non-reactive 334 335 fraction, only two elements (i.e., Ba and Ti) were excluded from the fingerprinting procedure. 336 Following the range test, the tracers selected by the combination of the Kruskal-Wallis test and DFA 337 were Co, Fe, V and Zn for the total fraction, and Al, Cr, Fe, Mn and Zn for the non-reactive fraction. 338 For the total fraction, the procedure statistically selected Zn to estimate the source contributions, 339 which could influence the reliability of the results. For the non-reactive fraction, two reactive metals 340 (i.e., Mn and Zn) were integrated into the geochemical model, but the estimation of source 341 contributions based on their non-reactive concentrations made it possible to overcome problems associated with metal reactivity or past anthropogenic inputs. These results are in agreement with the 342 343 work of Begorre et al. (2021), showing that tracer selection varied between both fractions leading to a 344 larger number of available tracers for the non-reactive fraction.

345

3.2.2. Global trends in historical SPM inputs to the Mediterranean Sea

Fig. 4 shows the contributions of SPM sources, expressed as percentages, estimated from tracer

- 347 concentrations in the total and non-reactive fractions. Overall, along the whole sediment core (1981–
- 2013), the contributions modeled from the total fraction differed significantly (p<0.05) from the

contributions estimated using the non-reactive fraction for the Durance and Isère rivers. For example, the SPM contributions of the Durance River modeled using the total and non-reactive fractions were $22 \pm 13\%$ and $52 \pm 18\%$, respectively. Given that the sediment core was collected in the Rhône River downstream of the confluence with the Durance, the sediment core is probably strongly influenced by SPM inputs from the Durance River (Vauclin et al., 2021).

Source contributions modeled using the total fraction showed that the Isère and Ardèche rivers were 354 the main contributors to the SPM at the outlet of the Rhône watershed from 2013 to 1991 (27–175 cm 355 356 depth) and from 1990 to 1981 (184-300 cm depth), respectively. Contributions estimated using non-357 reactive concentrations showed that the Durance River was the main source of deposited sediments over time, except in layers from 1989, 1996, 1998 and 2001 for which the main SPM inputs came 358 from other tributaries (Fig. 4-b). Globally, the Middle Rhône River inputs were low and relatively 359 360 stable $(5.6 \pm 3.1\%)$ over time, except in the layers from 1989 (37%), 2001 (19%) and 2003 (16%). In the absence of historical SPM flux data, we compared our results against results from the literature 361 (Zebracki et al., 2015; Poulier et al., 2019). Zebracki et al. (2015) used radionuclide analyses to 362 investigate contemporary sources of SPM transported to the Rhône River outlet between 2001 and 363 364 2011, and identified three groups of tributaries: upstream (Ain, Fier, Isère, and Saône rivers); pre-365 alpine (Durance, Drôme rivers) and Cévenol (Ardèche, Gardon rivers). The Durance River was identified as the main contributor of SPM (53%), followed by the "Andancette + Isère" (35%) and 366 367 Cévenol rivers (11%) (Zebracki et al., 2015). The results obtained here using the non-reactive fraction 368 $(52 \pm 18\%, 22 \pm 13\%, \text{ and } 26 \pm 16\%, \text{ respectively})$ are therefore more consistent with the findings of Zebracki et al. (2015) than the results obtained using the total fraction. We also compared our 369 370 estimates of source contributions against the relative SPM flux contributions calculated over the 2000–2016 period by Poulier et al. (2019) for the Middle Rhône (16%; Upper Rhône River + Saône), 371 372 Isère (25%), and Durance (24%) rivers. The major SPM contributors to the Mediterranean Sea were the Isère and Durance rivers. According to these results, our estimates of SPM contributions based on 373 the total fraction are likely more accurate than those based on the non-reactive fraction for the 2000-374 375 2016 period. However, these results should be interpreted with caution because Poulier et al. (2019)

did not include the Cévenol tributaries. The differences in conclusion between both comparisons 376 377 might be associated with the method applied to evaluate the contribution of SPM sources. Poulier et 378 al. (2019) used the SPM flux data calculated at each sub-watershed outlet without calculating relative SPM proportions delivered to the Rhône River, whereas Zebracki et al. (2015) and the present study 379 380 compared geochemical signatures between SPM sources and sediment core layers. Zebracki et al. 381 (2015) noted the presence of a large number of dams along the Rhône River continuum, which may 382 explain the underestimated contributions from upstream tributaries. Estimates of source contributions 383 using geochemical models were therefore more representative of SPM inputs to the watershed outlet. Regarding result reliability, the MAE calculated for each layer of the sediment core were higher for 384 the non-reactive fraction (average = 99.8%) than for the total fraction (average = 95.4%). Considering 385 each layer individually, it can be observed that all MAE were higher than 85%, which means that the 386 results are reliable except for the 2003 layer for the total fraction (MAE = 81%). 387

388

3.2.3. Temporal variations in historical SPM contributions

The contributions of the Andancette station estimated from total and non-reactive fractions were very low and not significantly different in top $(7 \pm 3\%)$ and $8 \pm 5\%$ for the total and non-reactive fractions, respectively; p=0.88) and bottom $(4 \pm 3\%)$ and $7 \pm 8\%$ for the total and non-reactive fractions, respectively; p=0.07) layers of the sediment core. In contrast, the source contributions of the Isère,

393 Ardèche, Gardon and Durance rivers varied over time (Fig. 4).

394 Differences in source contributions between both fractions depended on the sediment core layers. For 395 the total concentrations, the contributions of SPM sources differed significantly between the top of 396 the core (from 1991 to 2013) and the deepest layers (from 1981 to 1990), whereas for the non-reactive 397 fraction, the contributions were not significantly different between both parts of the sediment core. 398 For example, in the top of the sediment core, the contributions of the Ardèche and Gardon rivers were 399 similar for those estimated based on the total fraction $(13 \pm 6\%)$ and $14 \pm 13\%$, respectively) and the 400 non-reactive fraction $(15 \pm 5\%)$ and $13 \pm 13\%$ respectively), whereas in the deepest layers, the estimated source contributions were significantly different between fractions (total fraction: $32 \pm 16\%$ 401

402 and $22 \pm 14\%$ for the Ardèche and Gardon rivers, respectively; non-reactive fraction: $12 \pm 3\%$ and

 $13 \pm 9\%$, respectively). To explain these differences, a Pearson correlation test and a principal 403 404 component analysis (PCA) were carried out based on the tracers selected by DFA for the total (Co, 405 Fe, V, Zn) and non-reactive (Al, Cr, Fe, Mn, Zn) fractions, and results were interpreted according to 406 the geochemical model results (Fig. 5). These statistical tests showed that the contributions of the 407 Ardèche River were significantly correlated to Zn concentrations in the total fraction (p<0.05; Fig. 5-408 a). In addition, as previously demonstrated (see Section 3.1), Zn was characterized as highly reactive 409 in the 1981–1990 layers, meaning that it was influenced by past anthropogenic inputs (Dendeviel et 410 al., 2020). Consequently, the estimated increasing contributions from the Ardèche River from 1990 to 1981 did not reflect an increase in Ardèche River SPM inputs but instead revealed substantial historic 411 412 anthropogenic inputs of Zn at the global Rhône River scale, as discussed above. 413 3.2.4. Cross-analysis of relative historical SPM contributions and past flood events 414 To discuss and confront the results of the two fingerprinting approaches, we compiled the main 415 historical flood events (DREAL, 2011), and reported them in Fig. 4-c according to each identified 416 layer of the sediment core. The discussion is presented in reverse chronological order. 417 In 2010 (42 cm depth), a flood of the Isère River with a ten-year return period, i.e., a flow exceeding 900 m³ s⁻¹, was recorded, which implies higher SPM inputs from the Isère River compared to the other 418 419 tributaries. For this layer, there was no marked peak in contributions for any particular tributary in 420 either total or non-reactive fraction, as there was no sediment deposition from this flood event at the study site. For the total fraction, the Isère River SPM contribution was 39% against only 13% for the 421 422 non-reactive fraction. Combining the contributions of the Middle Rhône with those of the Isère River 423 for the present study, the SPM contributions modeled using the total and non-reactive fractions were 424 45% and 17% of total SPM inputs, respectively. For the Durance River, the SPM contributions 425 modeled from the total and non-reactive fractions were $36 \pm 19\%$ and $59 \pm 19\%$, respectively. Given that the Durance River is much closer to the sediment core site than the Isère River (see Fig. 1), it is 426 427 possible that the flood peak of the Isère River was not observed at this time in the sediment core at the Rhône River outlet because of SPM storage in the dam reservoir located downstream of the 428 429 confluence of the Isère River and the Rhône River (Zebracki et al., 2015).

In 2003 (96 cm depth), the deposited sediments were characterized by SPM transported during an 430 extensive Mediterranean flood implying a flood on the Durance (1100 m³ s⁻¹) and Ardèche 431 432 (2510 m³ s⁻¹) rivers. Total and non-reactive fractions showed a major contribution of the Durance River of 58% and 54%, respectively, which is consistent with the hydrological data illustrated in Fig. 433 4-c (DREAL, 2011) and with Zebracki et al. (2015), who estimated contributions of the Durance River 434 435 at between 38% and 53% during the Mediterranean floods for the 2000–2012 period. Contributions of 436 the Ardèche River were estimated at 21% and 25% for the total and non-reactive fractions, 437 respectively. These contributions could be explained by the flood event that occurred on the Ardèche River and reached a water flow of 2510 m³ s⁻¹, which is around three times higher than flood threshold 438 $(Q = 845 \text{ m}^3 \text{ s}^{-1}).$ 439

The generalized floods of 2001–2002 at the Andancette station (water flow of 4780 m³ s⁻¹) and the 440 Isère (928 m³ s⁻¹) and Gardon (6700 m³ s⁻¹) rivers was identified at 102 cm depth in the sediment core. 441 The geochemical modeling results based on the total and non-reactive fractions were similar for the 442 Ardèche (7% and 8%, respectively), Durance (11% and 8%) and Gardon (37% and 43%) rivers. Based 443 on the occurrence of a generalized flood at Andancette station, the contribution obtained with the non-444 445 reactive concentrations (19%) was more relevant than the contribution estimated using the total 446 fraction (5%). The SPM contribution from the Isère River was estimated at 23% based on the nonreactive fraction. Contributions modeled for the Gardon River could only be validated based on past 447 flooding events because of the absence of hydro-sedimentary data for this period. The SPM 448 contributions of the Gardon River for the total and non-reactive fractions (37% and 43%, respectively) 449 are consistent with the reported water flow of 6700 m³ s⁻¹, which is around six times higher than the 450 flood threshold ($Q = 402 \text{ m}^3 \text{ s}^{-1}$). Consequently, based on the SPM source contributions modeled here 451 for the Middle Rhône and Isère rivers, the non-reactive fraction appears to be more relevant than the 452 453 total fraction for tracing the SPM contributions of this major event.

454 The sediments deposited in 1996 (131 cm depth) reflect the SPM inputs during a Cévenol flood.

455 According to the 10-yr flood of the Ardèche River, which reached a maximum water flow of 1780 m³

456 s^{-1} , the estimated SPM contribution for this tributary was very small (0.5%). Results obtained for the

sediment core showed that, based on the total fraction, the Isère and Gardon rivers were the main SPM 457 458 contributors at 36% and 39%, respectively, while SPM inputs from the Durance, Ardèche and Middle 459 Rhône rivers were much lower (19%, 0.5% and 6%, respectively). In contrast, the modeling performed with the non-reactive concentrations resulted in more relevant estimates than using the total 460 concentrations, with contributions of 9% from the Ardèche River and 33% from the Gardon River. 461 462 However, it is possible that the geochemical signatures (based on the tracers selected) of the Gardon and Ardèche rivers were not clearly distinct, which would suggest that part of the SPM inputs from the 463 464 Ardèche River would be assimilated as inputs from the Gardon River. For this type of hydrological event, Zebracki et al. (2015) estimated that the contributions were similar between the upstream 465 tributaries (31%; Andancette station, Isère), the pre-alpine tributaries (30%; Durance), and the 466 Cévenol tributaries (39%; Ardèche, Gardon). According to these results, the estimated SPM 467 contributions of the Durance River in the 1996 layer are more similar using the non-reactive fraction 468 (23%) than the total fraction (19%). Moreover, by combining the estimated contributions for 469 470 Andancette station and Isère River, the contribution for the non-reactive fraction (35%) was similar to 471 the value estimated by Zebracki et al. (2015) (31%), in contrast to our geochemical modeling with the 472 total concentrations (42%).

473 In 1993 and 1994 (i.e., at 142–151 cm depth), there were three extensive Mediterranean floods that led to moderate floods of the Isère River and the Middle Rhône at Andancette and high water flows for the 474 475 southern tributaries (peaks of 2350 m³ s⁻¹ and 4340 m³ s⁻¹ for the Durance and Ardèche rivers, 476 respectively). Therefore, the main expected SPM contributions are probably those of the Ardèche and the Durance rivers. As the only hydrological data available for the Gardon River was the occurrence of 477 478 a 10-yr flood in 1993 (DREAL, 2011), it was not possible to validate the SPM contribution of this 479 tributary. Geochemical modeling with the total fraction resulted in a major contribution from the 480 Durance River (45%). In contrast to expected results, the Isère River had a larger relative SPM contribution (29%) than the Ardèche River (15%). Geochemical modeling with the non-reactive 481 482 fraction confirmed the Durance River as the main source of SPM (75%), followed by the Ardèche 483 River (13%). Moreover, SPM inputs from the Isère River were lower using the non-reactive fraction

(7%) than for the total fraction (29%), which once again suggests that the results obtained with thenon-reactive fraction are more reliable than results obtained with the total fraction.

486 Finally, in 1990 (184 cm depth), the Rhône was subjected to a very intense oceanic flood that 487 particularly affected the tributaries located upstream of the city of Lyon (Upper Rhône River) with peak water flows of 4310 m³ s⁻¹ and 1040 m³ s⁻¹ for the Middle Rhône and Isère River, respectively. 488 Our estimates modeled using the total fraction showed that the Isère and Gardon rivers were the main 489 contributors to SPM inputs at the outlet of the Rhône River basin and that the other three tributaries 490 491 supplied less than 4% sediment each. In contrast, contribution estimates modeled using the nonreactive fraction were equivalent between the Isère River (37%) and the Middle Rhône (37%), which 492 493 better reflects the SPM inputs of the major event studied. For this flooding event, the results modeled from the non-reactive fraction are therefore more reliable than the results obtained with the total 494 495 fraction.

496

497 **4.** Conclusion

498 To overcome the non-conservative behavior of metals, we used the metal concentrations in the non-499 reactive fraction of SPM/sediments on a sediment core collected at the outlet of the Rhône basin, a site 500 for which no information on the historical SPM contributions was available. This study focused on the 501 estimation of tributary contributions over the last 40 yr using a geochemical modeling approach. We demonstrated that estimations of SPM contributions were significantly influenced by past 502 anthropogenic inputs responsible for an increase in total Zn concentrations in the deepest layers of the 503 sediment core. In fact, the range test selected Zn as a conservative tracer even though it is highly 504 reactive in the deepest layers of the sediment core, which may bias the results of contribution 505 modeling based on the total fraction. Based on a comparison with the results of Zebracki et al. (2015), 506 who used radionuclide data on SPM, it is clear that the results obtained with the non-reactive fraction 507 508 are closer to those obtained by Zebracki et al. (2015) compared to contributions estimated using the total fraction. Indeed, using the non-reactive fraction, we showed that the main SPM contributor over 509

the 1981–2013 period was the Durance River. Moreover, the detailed study of major past flooding 510 events showed that our estimates of tributary SPM contributions were more reliable and consistent 511 512 when the non-reactive metal concentrations were used, especially to trace SPM sources in the deepest layers of the sediment core. Therefore, when concentrations of a geochemical element are influenced 513 by anthropogenic inputs, as for investigations relative to historical SPM inputs, it is more relevant to 514 515 use the non-reactive fraction rather than the total fraction. This study demonstrated that our original 516 fingerprinting method based on the non-reactive fraction of metals in SPM/sediment is a robust tool 517 for estimating source contributions in a sediment core, as it removed the influence of past anthropogenic inputs on tracer concentrations. Furthermore, this fingerprinting approach made it 518 519 possible, for the first time in the Rhône River basin, to reconstruct the historical contributions of the main tributaries during major flooding events. It would now be instructive to apply this method to 520 SPM/sediments in coastal environments or those affected by significant past anthropogenic inputs 521 (e.g., rivers influenced by mining activities). 522 523 Thus, this fingerprinting method using the residual fraction at the Rhône River basin scale allows to: 524 Increase the number of available tracers after the range test of SPM sources, as demonstrated by Begorre et al. (2021), 525 Remove the influence of past anthropogenic inputs of metals such as Zn, 526 -Provide results that are more relevant when comparing with available hydrological and 527 _ sedimentary data, 528 529 Provide information on the major sources of sediments that are exported into the Mediterranean _ Sea that could impact its ecosystems. 530 531 In terms of recommendations for future studies, as presented by Begorre et al. (2021), the use of total metal concentrations, especially when they are highly reactive, is problematic. Indeed, source 532 533 fingerprinting using total concentrations of metals must be applied only for metals with low reactivity

to improve estimation of SPM source contributions. Therefore, before applying source fingerprinting in

535 a river basin, it is necessary to investigate metal reactivity to avoid the use of high-reactive metals.

Finally, we highly recommend using tracers in the non-reactive fraction when sources and target sampleswere not sampled at the same period (e.g., with a difference of more than 5 or 10 yr).

538

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- 551

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Fig. 1: Location of sampling sites for SPM sources (Middle Rhône River at Andancette station, Isère, Ardèche, Gardon, Durance river stations of the Rhône Sediment Observatory - OSR network) and sediment core (Mas des Tours at the outlet of the Rhône River).

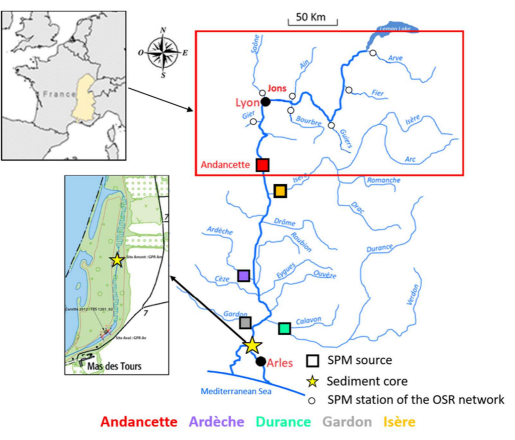
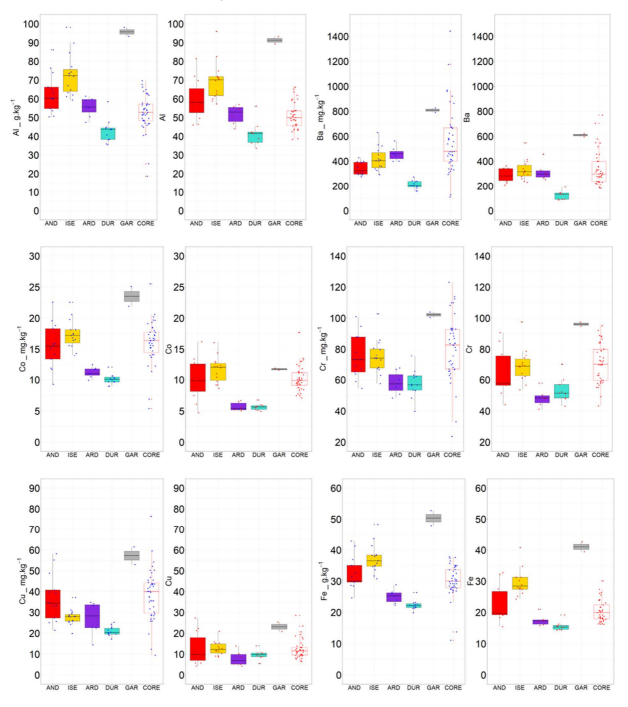
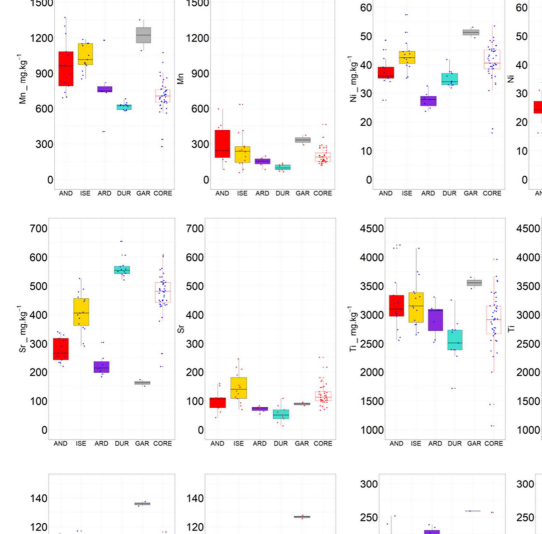


Fig. 2: Metal concentrations (non-corrected data by particle size) measured in the total (blue on the left) and non-reactive (red on the right) fractions of SPM in the Rhône River (Middle Rhône River station at Andancette - AND) and four tributaries (Isère –ISE, Ardèche – ARD, Durance- DUR, Gardon-GAR) and in the sediment core sampled in the Rhône at Mas des Tours site.





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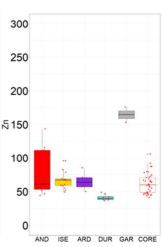
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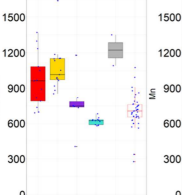
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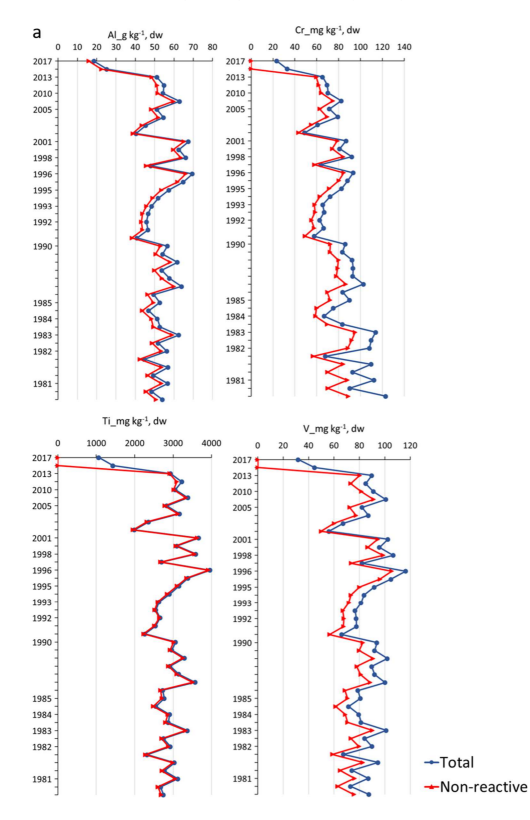
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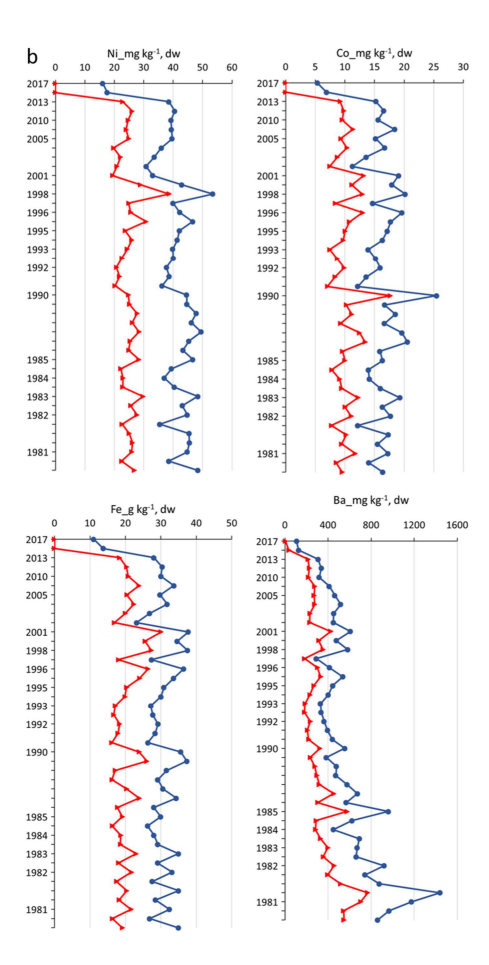
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Fig. 3: Metal concentrations in the sediment core sampled in the Rhône at Mas des Tours site. The three identified groups represent metals that are not reactive (a), moderately reactive (b) and highly reactive (c). Example of three behaviours of metals along the sediment core by comparing concentrations in the total (blue line) and non-reactive (red line) fractions.





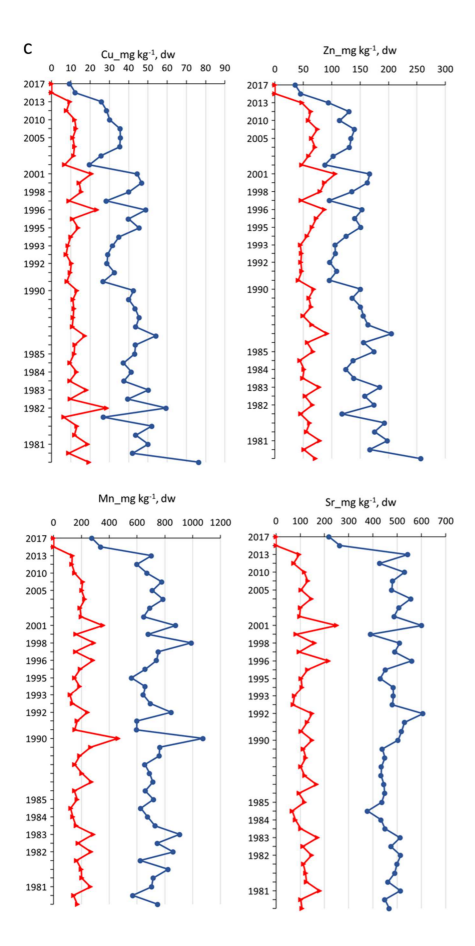


Fig. 4: Profile of source apportionment modelling by using tracers in the total (a) and non-reactive (b) fractions, in sediment core at Mas des Tours site, from 1981 to 2013. The black line represents the 1990 layer. Main historical flooding events (m³ s⁻¹) are reported for the Upper Rhône River and each tributary over the 30-yr period (c)

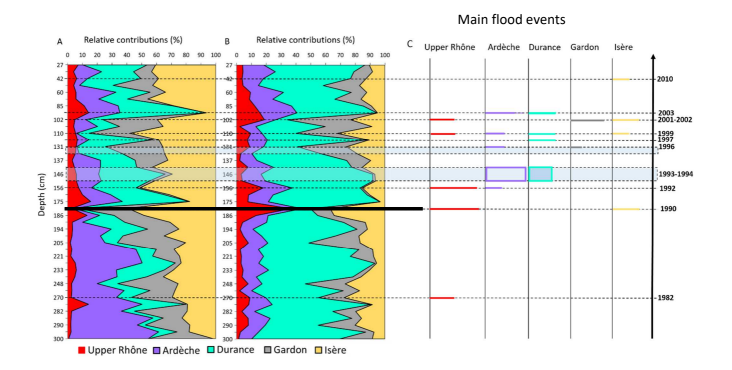
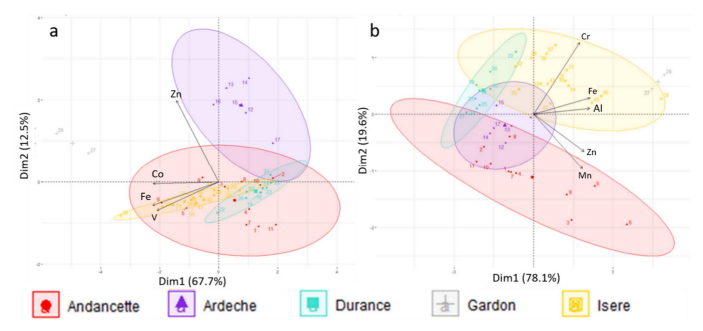


Fig. 5: Principal component analysis (PCA) performed using selected tracer concentrations in the total (a) and non-reactive (b) fractions of SPM at the Rhône River Middle station (Andancette) and tributaries. The concentrations of the tracers used are corrected for particle size. PCA shows correlations (or no correlation) between total (a) or non-reactive (b) concentrations and source contributions.



Tributary	N =42 samples	Water flow	SPM concentration	
		(m³ s⁻¹)	(mg L⁻¹)	
Middle Rhône River	11	263 - 1040	3.5 - 18.8	
(Andancette station)	(flood: 3, base flow: 8)	(2018-2019 period)	(2019)	
lsère	14	175 – 671	6.75 – 1488	
	(flood: 3, base flow: 11)	(2014-2018)	(2014-2018)	
Ardèche	6	9 - 459	1.7 - 13.8	
	(flood: 1, base flow: 5)	(2016)	(2016)	
Gardon	2	-	-	
	(flood: 1, base flow: 1)	(2017)	(2017)	
Durance	9	38 - 1613	6.2 - 545	
	(flood: 2, base flow: 7)	(2011-2016)	(2014-2016)	

Table 1: Summary of sampling periods for each tributary and associated hydro-sedimentary conditions.

Table 2: Proportions (in %) of the reactive fraction compared to the total fraction for each trace and major elements. The results are presented for the SPM tributaries and for the sediment core samples divided in two main periods (1991-2013 and 1981-1990).

Elements	Proportions of the reactive fraction (%)						
	Middle Rhône	lsère	Ardèche	Durance	Gardon	Sediment core (1991- 2013)	Sediment core (1981- 1990)
Al	7±1	3.0±0.5	7.2±0.5	4.5±0.5	4.7±0.5	5.3±0.9	6.3±0.5
Ва	18±4	19±3	32±6	40±8	25±4	39±7	43±6
Со	38±7	32±5	48±6	43±5	50±6	39±4	38±4
Cr	14±3	7±1	16±4	9±1	5.9±0.3	11±2	18±4
Cu	68±9	54±7	68±10	55±8	60±1	66±7	72±6
Fe	31±6	19±2	30±2	30±3	19±1	32±5	37±5
Mn	72±10	76±11	78±7	82±7	72±9	73±6	72±7
Ni	35±5	23±3	33±3	31±2	29.0±0.2	39±5	42±3
Sr	66±9	64±9	66±7	89±7	45.2±0.1	76±7	74±5
Ti	3±1	0.7±0.1	8±2	0.4±0.1	1.8±0.2	1.5±0.7	2.0±0.4
V	13±2	5±1	17±2	10±1	6.7±0.2	11±2	13±1
Zn	56±7	33±6	65±7	43±5	42±1	50±6	63±5

Table 3: Statistical results for tracer selection procedure, which combines the range test, the Kruskal-Wallis test (KW, p<0.05) and the discriminant factor analysis (DFA), for total and non-reactive fractions of SPM sources. The "V" shows the geochemical tracers that are retained at each step and "X" shows those that fail. For the range test, the number of layers, for which metals failed the range test, are indicated in parenthesis. For DFA, the "X" shows the metals removed from the procedure while the metals retained are characterised by a value representing the discriminatory power of the selected tracers.

Elements	Total fraction			Non-reactive fraction		
	Range test	KW test	DFA	Range test	KW test	DFA
Al	V	V	Х	V	V	1.00
Ва	X (5 layers)	Х	Х	X (2 layers)	Х	Х
Со	V	V	0.55	V	V	Х
Cr	X (1 layer)	Х	Х	V	V	0.95
Cu	X (1layer)	Х	Х	V	V	Х
Fe	V	V	0.95	V	V	0.62
Mn	V	V	Х	V	V	0.98
Ni	V	V	Х	V	V	Х
Sr	V	V	Х	V	V	Х
Ti	X (1 layer)	Х	Х	X (1 layer)	Х	Х
V	V	V	1.00	V	V	Х
Zn	V	V	0.88	V	V	0.81

Supplementary material

Fig. SI.1: Box-and-whisker plot of median grain size (D50, values in μ m) in SPM tributaries (Upper Rhône River station at Andancette – AND, Isère –ISE, Ardèche – ARD, Durance- DUR, Gardon- GAR) and in the sediment core sampled in the Rhône at Mas des Tours site.

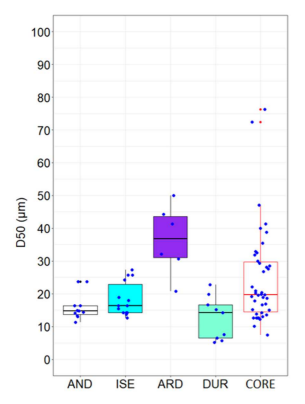


Fig. SI.2: Profile of median particle size (D50; $\mu m)$ in the sediment core.

