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Reactivity of particulate element concentrations: apportionment assessment of suspended particulate matter sources in the Upper Rhône River, France

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Abstract

Purpose Suspended particulate matter (SPM) transport through rivers is a major vector of nutrients and pollutants to continental shelf areas. To develop efficient sediment management strategies, there is a need to obtain quantitative information on SPM sources. For many years, the geochemical properties of SPM have been commonly used as tracers to identify sediment sources. In large watersheds, with numerous sources, the expected alteration of tracers during their transport requires that their reactivity be taken into account.

Materials and methods To overcome this issue, we tested the use of major and trace element signatures in the residual fraction of SPM, using two different extraction methods. This original fingerprinting approach was applied to the Upper Rhône River basin (~20,000 km²) in order to assess the respective SPM contributions of its main five tributaries (Arve, Ain, Fier, Guiers, and Bourbre Rivers) for contrasted hydrological conditions (base flow periods, flood events and dam flushing). By incorporating element concentrations previously corrected from particle size distribution in a mixing model coupled to Monte Carlo simulations, we estimated the associated uncertainties of the SPM contributions from each tributary. The relative SPM contributions obtained using this fingerprinting approach were compared with those calculated with a 1-D hydro-sedimentary model.

Results and discussion The use of element concentrations, such as Zn, P, Cu, Pb, Mn, or Sr, in the total fraction of SPM as conservative fingerprinting properties was not suitable, since they are mainly bound (> 50%) to reactive carrier phases. By using Ba, Ni, Fe, Mg, Cu, Sr, and V concentrations in the SPM residual fraction, the apportionment modeling of SPM sources was successfully assessed. The fingerprinting approach showed that, in base flow conditions, SPM originated mainly from the Arve River. During dam flushing event, the fingerprinting approach consistently estimated that re-suspended sediments came from the Arve River, while the 1-D hydro-sedimentary model estimated a proportion of re-suspended sediment originating within the Rhône River.

Conclusions This original fingerprinting approach highlighted the relevance of using geochemical properties in the non-reactive fraction of SPM in order to obtain reliable information on spatial sources of SPM in a large river basin. This methodology opens up promising perspectives to better track SPM sources in highly reactive environments such as estuaries/delta or in historical sediment archives.

Keywords Fingerprinting · SPM fluxes · Trace metals · Sediment · Hydro-sedimentary model · Uncertainties

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1 Introduction

Natural processes such as soil weathering and erosion strongly control suspended particulate matter (SPM) inputs to rivers (Koiter et al. 2013). These particles are essential for aquatic ecosystems as they control the transport of nutrients, essential elements, and organic matter (House and Warwick 1999; Collins et al. 2005). However, excessive inputs of SPM can

ultimately degrade water quality. Indeed, several contaminants such as trace metals, polychlorinated biphenyls, or polycyclic aromatic hydrocarbons can be bound to particles characterized by large surface specific area and high surface reactivity, which makes SPM a major source of contaminant fluxes in surface freshwaters (Heemken et al. 2000). Gellis and Walling (2011) showed how vital it is to combine source estimation and SPM budget approaches in order to provide resource managers with the necessary information and support for reducing SPM and associated contaminant fluxes. Source estimation approaches can quantify the relative importance of potential sources of SPM, while SPM budget approaches provide information on the magnitude of the SPM fluxes and the links between SPM sources and sinks. Since the estimation of SPM sources through monitoring networks is expensive and requires ongoing maintenance, several fingerprinting methods have been developed in the past 40 years to obtain quantitative information on spatial distribution (e.g. individual tributaries, geological zones) and type (e.g. surface soils and channel banks) of SPM sources (e.g. Walling 2013). These methods are based on the fact that SPM from each sub-catchment are characterized by distinct geophysical and geochemical and biochemical properties, which could be used to determine the relative proportion of SPM from each potential upstream source (Walling et al. 1999). As no single property can reliably distinguish various sources of SPM (Collins et al. 1997), various physico-chemical characteristics of SPM have been used to assess their origin, such as major elements and trace metals (Collins et al. 2012), rare earth elements (Kimoto et al. 2006), mineralogy and magnetism (Rotman et al. 2008), radionuclides (Zebracki et al. 2015), color properties (Martínez-Carreras et al. 2010; Legout et al. 2013), isotopes (Mukundan et al. 2010), pollens (Brown et al. 2008), or biomarkers (Chen et al. 2016).

In order to determine SPM sources at any time and location within a watershed, the properties of SPM used in any fingerprinting method have to be conservative from the source area through the final area of sampling (Walling 2013). In some studies, a simple range test is incorporated into the data screening to confirm that the property values characterizing the target sediment fall within the range of the property values associated with the potential sources (Walling 2013). As mentioned by Walling (2013), enrichment and depletion effects caused by selective mobilization and transport, resulting in different grain size composition and organic matter content between target and source material samples, can be viewed as a form of non-conservative behavior. To overcome this problem, it is possible to remove organic matter from SPM by ashing and analyze only their mineral fraction (e.g. Wilkinson et al. 2013). However, this approach only corrects element concentrations that are strongly linked to particulate organic matter but fails to account for other reactive phases (e.g. carbonates). Lacey et al. (2017) reported in their review

that to address particle size effect, fractionation of source sediment material (e.g. isolation of particle size fractions) or concentration correction (normalization) are often used in sediment fingerprinting studies. Some approaches simply restrict tracer analysis to specific particle size fractions of the SPM sample (e.g. below 63 μm or 10 μm) (e.g. Fu et al. 2008), but these fine fractions are not necessarily representative of the total SPM load. In order to compare samples with similar particle size distribution or organic matter content, several approaches have been applied, such as using pre-sieved sediment or by applying correction factors (Motha et al. 2002). Gellis and Noé (2013) proposed to determine for each source if significant relationships exist between median values of the particle size distribution (D50) or total organic carbon (TOC) content and tracer concentrations. In case of a significant relationship, a correction factor could be applied to the tracer concentration. However, although particle size correction is widely applied, there are some documented limitations. For example, particle size corrections (i.e. positive linearity between particle size and tracer concentration) do not apply to all tracer properties or not equally to all tracers of SPM samples from different sources (Smith and Blake 2014).

In large-sized watersheds, the notion of conservative behavior of tracer properties is even more important than in small watersheds. Indeed, in addition to logistical difficulties to retrieve SPM over the entire system, tracer properties are subjected to intense modifications of physico-chemical conditions (e.g., ionic strength, pH, and redox potential) during their transport, which will influence their partition between particulate and dissolved phases (Dabrin et al. 2009) by adsorbing, precipitating, or releasing them. Thus, tracer properties associated with highly reactive particulate carrier phases, such as Fe and Mn oxy-hydroxides, particulate organic matter, carbonates, and sulfides, may eventually be released in the dissolved phase, modifying the tracer signature in SPM or sediment. But, in the vast majority of fingerprinting studies, tracer properties, such as major and trace element concentrations, are commonly obtained after a total mineralization of SPM or sediment samples (e.g. triacid mineralization by using HCl, HNO₃ and HF; Le Cloarec et al. 2011; Evrard et al. 2011). As this total extraction reflects the signature of the sum of the different carrier phases, this implies that the source signature inferred from total element concentrations may be modified according to the physico-chemical conditions that SPM will encounter during their transport. To overcome this issue, Maher et al. (2009) successfully used magnetic inclusions (protected within host silicate grains) to assess sediment sources in a tropical and marine context. In a similar way, Dabrin et al. (2014) proposed to use trace metal concentrations in the residual fraction of SPM, a fraction supposedly invariable. This approach was successfully applied to distinguish two main SPM sources on the French Atlantic coast.

Validation of fingerprinting estimation also represents a big challenge. One of the most commonly used approaches is to compare model estimation with SPM and discharge monitoring data (e.g. Evrard et al. 2011). However, SPM sampling techniques represent some limitations in terms of practicalities and costs of deployment both spatially and temporally (Collins and Walling 2004). Moreover, SPM monitoring provides information on SPM fluxes but not directly on the origin of the remobilized sediment (Vale et al. 2016). Another way to address this problem is to prepare experimental mixtures in the laboratory with known proportions of SPM and their subsequent analysis (e.g. Franks and Rowan 2000). This validation by using real or virtual sediment mixtures (e.g. Palazon et al. 2015) is highly promising even if dealing with contrasted particle size distribution in source samples represents a challenge. A promising alternative is the use of a 1-D numerical model to simulate sediment transport (e.g. Launay et al. 2015; Maleki and Khan 2016). At the river network scale, most 1-D numerical models are able to simulate complex, multi-reach hydraulic networks, including artificial structures as well as bifurcations and confluences. Thus, these models are able to estimate the relative contributions of water discharge and SPM fluxes from each tributary at a downstream monitoring station, accounting for SPM propagation, deposition, and re-suspension.

In light of these findings, the objective of the present study was to assess the benefits of using major and trace element

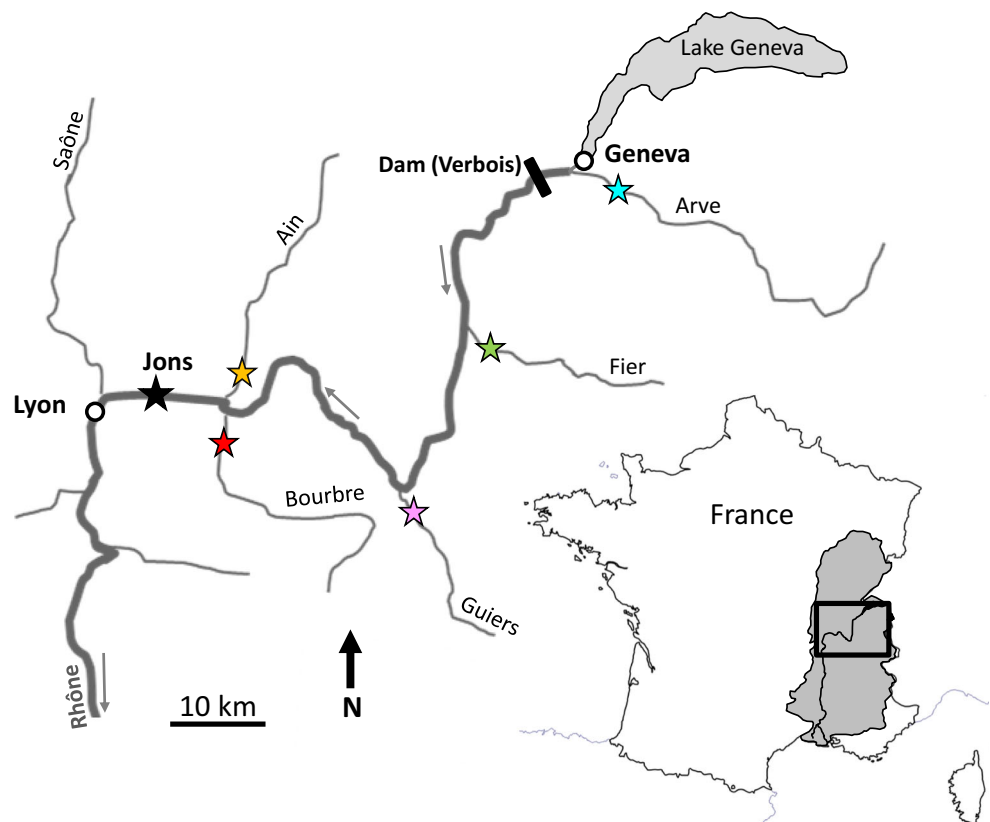
concentrations in the residual fraction of SPM rather than the total element concentrations, in order to fingerprint SPM sources in a large river catchment. The Upper Rhône, a large-sized watershed (~20,000 km²) in France, including five main tributaries (the Ain, Arve, Bourbre, Guiers, and Fier Rivers), was used as a case study. Major and trace elements were analyzed in the total and in the residual fractions of SPM collected during contrasting hydrological conditions at the outlet of the main tributaries and at the outlet of the Upper Rhône River (Jons station) in order to characterize SPM sources. Then, source apportionment modeling was applied to SPM sampled for different hydrological conditions (base flow, floods, or dam flushing) at Jons station. Source apportionments inferred from fingerprinting approaches were compared to simulations obtained on the same sampling periods through 1-D hydro-sedimentary model computations.

2 Material and methods

2.1 Study area and SPM sampling strategy

The Rhône River is one of Europe's major rivers: at its outlet (Beucaire station), its length is 812 km, its mean discharge is 1690 m³ s⁻¹, and its catchment area is 95,590 km² (Fig. 1). It is the main source of water and SPM to the Mediterranean Sea

Fig. 1 Map of the Upper Rhône River (France), with the location of the sampling station at Jons and the 5 sampling stations on the main tributaries (Arve, Fier, Guiers, Ain, and Bourbre Rivers)



since the Aswan Dams were built across the Nile (Antonelli et al. 2008; Ollivier et al. 2011). Data on SPM sources are scarce and limited to the most southern tributaries of the Rhône catchment (Navratil et al. 2012; Zebracki et al. 2015). From Lake Geneva in Switzerland to the Mediterranean Sea, the Rhône River is strongly affected by industrial installations including factories, 21 hydroelectric plants, and four nuclear power plants. The SPM output of Lake Geneva is assumed to be negligible due to the large trapping capacity of the lake (Launay et al. 2019). At Jons station (JON, $\sim 20,000 \text{ km}^2$), upstream of the city of Lyon, the Rhône receives water and SPM inputs from five main tributaries (Fig. 1, Table 1), namely, the Arve (ARV), the Fier (FIE), the Guiers (GUI), the Bourbre (BOU), and the Ain (AIN) Rivers. Monitoring stations (permanent or temporary) have been implemented at the outlet of each tributary since 2011 for the oldest, as part of the Rhône Sediment Observatory (OSR) program (Poulier et al. 2019). For each sampling station, all data of water discharge and SPM concentrations are available on the BDOH website (Thollet et al. 2018).

For each tributary, SPM samples ($9 \leq n \leq 16$, depending on the station) were collected between March 2011 and January 2015 by contrasting hydrological conditions (Table 2), i.e. during base flow periods and flood events (defined as peak water discharges with a return period greater than 2 years). Each tributary drains a large sub-basin that cannot be considered homogeneous in terms of land use, soil types, and geology. Therefore, it is important to collect samples under different hydrological conditions in order to take into account (i) as many different sources of SPM as possible within the sub-basins and (ii) the idea of connectivity, whereas certain parts of the watersheds are only connected during high magnitude and low frequency events (Fryirs 2013). This sampling strategy allowed collecting samples for SPM concentrations ranging from 8.0 to 38 mg L^{-1} (AIN), 10 to 680 mg L^{-1} (ARV), 64 to 210 mg L^{-1} (BOU), 2.3 to 34 mg L^{-1} (FIE), and 8.0 to 269 mg L^{-1} (GUI). Moreover, 16 SPM samples were collected at the mixing station (Jons) during base flow periods, flood events, and dam flushing operations on the Verbois dam (June 2012, Lepage et al. 2020). Indeed, in 2012, the hydroelectric dam at Verbois, located 12 km downstream of the

confluence of the Arve and Rhône Rivers (Fig. 1), was opened to avoid accidental flooding of the city of Geneva by flushing sediments from the Arve River accumulated since 2003. On the Rhône River at Jons station, SPM samples were collected for SPM concentrations ranging from 11 to 280 mg L^{-1} , while during dam flushing operations SPM concentrations ranged from 160 to 660 mg L^{-1} .

Suspended particles were retrieved by two complementary sampling methods (Table 2): time-integrative sediment particle traps (PT) or a continuous flow centrifuge (CFC; Westfalia KA 2-86-075–1978; 9700 rpm) as described by Masson et al. (2018). Comparisons between PT and CFC sampling methods conducted by Masson et al. (2018) have shown that no significant differences were found in PCB and Hg concentrations. We assume that such differences would also be negligible for metals. Two SPM samples were also retrieved using an automatic water sampler (Bühler 4011) at Jons station during the dam flushing event. The PTs were high-quality stainless steel boxes with three holes through the front and back faces allowing water to circulate inside. The PTs were typically installed at 50–100 cm below the water surface. Sampling with CFC was done by pumping water from the river into the centrifuge at a flow rate of $\sim 700\text{--}800 \text{ L h}^{-1}$. The PTs and the CFC provided the amount of SPM (several grams of dry weight) necessary for the analysis of several parameters. Particles retrieved from the PT or obtained after centrifugation were homogenized in a glass bottle with a silicon spatula and were conditioned in an amber glass bottle for chemical analysis and in a polypropylene (PP) tube for grain size determination. Then, SPM collected in glass bottles were freeze-dried, ground, and stored at ambient temperature until chemical analysis. Suspended particulate matters collected in PP tubes were stored at 4 °C in the dark until particle size distribution analysis.

2.2 Particle size distribution

Volumetric particle size distribution of SPM was assessed by laser diffraction using a Cilas 1190 particle size analyzer according to the ISO 13320 standard method (AFNOR 2009b). A representative wet subsample was introduced into the

Table 1 Location of the sampling stations on the Upper Rhône River and characteristics of the 6 studied rivers and watersheds

River	Mean annual discharge ($\text{m}^3 \text{ s}^{-1}$)	Mean annual SPM concentrations (mg L^{-1})	Sampling station name	Surface area (km^2)	Distance from Jons station (km)	Sampling station location Lat./Long (WGS84)	
ARV Arve	73	154	Geneva, Bout du Monde	2060	175	46° 10' 49.19" N	6° 09' 33.74" E
FIE Fier	41	35	Motz	1380	113	45° 56' 00.13" N	5° 50' 28.29" E
GUI Guiers	16	27	Belmont-Tramonet	609	77	45° 34' 23.28" N	5° 39' 04.09" E
BOU Bourbre	7.8	71	Tignieux	728	13	45° 42' 55.20" N	5° 09' 33.05" E
AIN Ain	123	5.4	Pont de Chazay	3600	27	45° 49' 55.11" N	5° 13' 44.46" E
JON Rhône	600	28	Jons	20,300	–	45° 48' 42.62" N	5° 05' 08.86" E

Table 2 List of collected SPM samples, sampling methods, hydrological conditions, and SPM concentrations and particle size during base flow periods, flood events, and dam flushing operations. (ND: not determined)

River	Sample	SPM sampling method	Date/period of sampling	Hydrological condition	Mean discharge ($\text{m}^3 \text{s}^{-1}$)	Mean SPM concentration (mg L^{-1})	Particle size D50 (μm)
Ain	AIN_1	Mobile centrifuge	28/08/2012–25/09/2012	Base flow	368	ND	13
	AIN_2	Mobile centrifuge	25/09/2012–23/10/2012	Flood	688	ND	11
	AIN_3	Mobile centrifuge	23/10/2012–21/11/2012	Flood	511	ND	7
	AIN_4	Mobile centrifuge	21/11/2012–28/01/2013	Flood	679	ND	6
	AIN_5	Mobile centrifuge	08/12/2011	Flood	721	83	5
	AIN_6	Particle trap	02/01/2012	Base flow	72.9	8.4	20
	AIN_7	Particle trap	05/01/2012	Base flow	158	23	18
	AIN_8	Particle trap	06/01/2012	Flood	178	38	19
	AIN_9	Particle trap	11/11/2012	Flood	269	34	27
Arve	ARV_1	Mobile centrifuge	14/03/2014–22/04/2014	Base flow	17.4	ND	ND
	ARV_2	Mobile centrifuge	11/12/2012–22/01/2013	Base flow	41.6	ND	12
	ARV_3	Mobile centrifuge	22/01/2013–19/02/2013	Base flow	101	70	7
	ARV_4	Mobile centrifuge	27/03/2013–16/04/2013	Base flow	40.8	10	12
	ARV_5	Particle trap	16/05/2013–05/06/2013	Base flow	71.1	90	ND
	ARV_6	Particle trap	19/06/2013–30/07/2013	Base flow	76.4	169	28
	ARV_7	Particle trap	30/07/2013–26/08/2013	Flood	62.7	343	50
	ARV_8	Particle trap	26/08/2013–24/09/2013	Flood	92.2	296	28
	ARV_9	Particle trap	09/03/2011	Base flow	122	101	ND
	ARV_10	Particle trap	20/01/2012	Flood	114	680	ND
Bourbre	BOU_1	Mobile centrifuge	05/03/2012–30/03/2012	Flood	22.3	205	21
	BOU_2	Mobile centrifuge	05/03/2012–30/03/2012	Flood	16.9	ND	15
	BOU_3	Mobile centrifuge	30/03/2012–07/05/2012	Flood	30.8	ND	9
	BOU_4	Particle trap	30/03/2012–07/05/2012	Base flow	5.96	164	ND
	BOU_5	Particle trap	15/03/2013–11/04/2013	Base flow	5.96	164	ND
	BOU_6	Particle trap	11/04/2013–06/05/2013	Flood	9.65	161	ND
	BOU_7	Particle trap	01/07/2013–25/07/2013	Flood	9.65	161	ND
	BOU_8	Particle trap	25/07/2013–28/08/2013	Flood	13.7	104	20
	BOU_9	Particle trap	11/11/2012	Flood	15.7	159	24
	BOU_10	Particle trap	02/01/2012	Base flow	5.7	64	19
	BOU_11	Particle trap	27/11/2012	Flood	5.55	134	3
Fier	FIE_1	Particle trap	05/03/2014–15/04/2014	Base flow	48.1	2.0	26
	FIE_2	Particle trap	15/04/2014–20/05/2014	Base flow	40.8	13	17
	FIE_3	Particle trap	20/05/2014–25/06/2014	Base flow	25.5	21	20
	FIE_4	Particle trap	25/06/2014–29/07/2014	Flood	52.4	28	22
	FIE_5	Particle trap	29/07/2014–25/08/2014	Flood	49.2	7.0	24
	FIE_6	Particle trap	25/08/2014–18/09/2014	Base flow	17.5	50	12
	FIE_7	Particle trap	18/09/2014–20/10/2014	Base flow	10.3	20	10
	FIE_8	Particle trap	20/10/2014–25/11/2014	Flood	31.8	34	37
	FIE_9	Particle trap	25/11/2014–20/01/2015	Base flow	18.9	31	26
	FIE_10	Particle trap	18/02/2015–18/03/2015	Flood	53.9	351	43
	FIE_11	Particle trap	18/03/2015–20/04/2015	Flood	66.5	510	65
	FIE_12	Particle trap	01/09/2015–28/09/2015	Flood	26.7	232	32
	FIE_13	Particle trap	28/10/2015–23/11/2015	Flood	17.1	167	37
	FIE_14	Particle trap	21/12/2015–19/01/2016	Flood	61.9	154	43
	FIE_15	Particle trap	19/01/2016–18/02/2016	Flood	72	136	47
Guiers	GUI_1	Mobile centrifuge	05/04/2012–27/04/2012	Base flow	20	8.4	ND

Table 2 (continued)

River	Sample	SPM sampling method	Date/period of sampling	Hydrological condition	Mean discharge (m ³ s ⁻¹)	Mean SPM concentration (mg L ⁻¹)	Particle size D50 (μm)
	GUI_2	Mobile centrifuge	05/04/2012–27/04/2012	Base flow	42.7	ND	ND
	GUI_3	Particle trap	11/05/2012–25/05/2012	Flood	32	48	ND
	GUI_4	Particle trap	11/05/2012–25/05/2012	Flood	32	48	ND
	GUI_5	Particle trap	19/06/2012–19/07/2012	Flood	30.4	57	ND
	GUI_6	Particle trap	05/04/2012	Flood	30.4	57	ND
	GUI_7	Particle trap	27/04/2012	Base flow	2.83	9.0	ND
	GUI_8	Mobile centrifuge	10/12/2018	Flood	109	119	17
	GUI_9	Mobile centrifuge	11/12/2018	Flood	35	19	20
	GUI_10	Manual sampling	15/03/2019	Base flow	15	264	18
	GUI_11	Mobile centrifuge	04/04/2019	Flood	32	18	18
	GUI_12	Mobile centrifuge	20/05/2019	Flood	216	259	9
	GUI_13	Manual sampling	13/06/2019	Flood	32	269	17
Rhône at Jons	JON_1	Mobile centrifuge	08/06/2012–10/06/2012	Flood	1825	130	9
	JON_6	Mobile centrifuge	14/06/2012–15/06/2012	Dam flushing	1330	662	11
	JON_8	Mobile centrifuge	22/05/2012–05/06/2012	Dam flushing	1085	336	13
	JON_15	Mobile centrifuge	11/06/2012–14/06/2012	Flood	1975	283	7
	JON_16	Automatic sampler	20/06/2012–03/07/2012	Dam flushing	804	163	9
	JON_17	Automatic sampler	03/07/2012–17/07/2012	Dam flushing	1290	507	12
	JON_18	Particle trap	17/07/2012–01/08/2012	Flood	641	30	60
	JON_21	Particle trap	01/08/2012–14/08/2012	Dam flushing	1125	510	45
	JON_23	Particle trap	14/08/2012–28/08/2012	Flood	740	62	17
	JON_24	Particle trap	28/08/2012–11/09/2012	Base flow	649	39	18
	JON_25	Particle trap	11/09/2012–25/09/2012	Base flow	427	17	14
	JON_26	Particle trap	04/12/2012–17/12/2012	Base flow	434	13	13
	JON_27	Particle trap	06/01/2012	Base flow	397	12	12
	JON_28	Particle trap	13/06/2012	Base flow	440	11	13
	JON_29	Particle trap	15/06/2012	Base flow	404	9.2	13
	JON_30	Particle trap	16/12/2012	Flood	1078	82	27

analyzer respecting good obscuration rate (typically 15%). An ultrasonic pretreatment was applied before (30 s) and during the analysis to prevent particle aggregation. The volumetric particle size distribution of the sample was computed using the Fraunhofer optical model (AFNOR 2009b). The median diameters of particle size distribution (D50; in μm), D10 and D90, were computed from the volumetric particle size distributions with the manufacturer's data processing software. An internal standard sample of sediment (Azergues River) was systematically used to determine the analytical error (relative standard deviation, $n = 30$) for the D50, D10, and D90 values that were below 3%, 4%, and 2%, respectively.

2.3 Extraction and analysis of major and trace elements

Total trace and major element concentrations in SPM were analyzed from representative sub-samples (30 mg d.w.)

digested in closed Teflon reactors using 1.5 mL HCl (12 M, Suprapur), 0.5 mL HNO₃ (14 M, Suprapur), and 2 mL HF (22 M, Suprapur). Reactors were kept at 110 °C in an automatic heating block (SC154 HotBlock, Environmental Express) for 2 h. After complete cooling, the digested solutions were evaporated to dryness. The dry residues were dissolved with 250 μL HNO₃ (14 M, Suprapur) and 5 mL of ultrapure water (Elga, Veolia) then heated for 30 min at 100 °C. Exactly 3.5 mL of the solution were made up to 10 mL with 6.5 mL of ultrapure water (Elga, Veolia) and stored at 4 °C in the dark until further analysis.

Extraction of acid-soluble elements (HCl 1 M) is commonly applied to quantify the element fraction bound to the most reactive phases of SPM or sediment (e.g. Morse and Luther III 1999). The HCl-available fraction was extracted by HCl (1 M, Suprapur; 12.5 mL) from 200 mg d.w. of SPM continuously agitated for 24 h in PP tubes (50 mL, Starstedt). Samples were then centrifuged (Heraeus Multifuge, X1R) for 20 min at

3000 rpm. Aliquots (10 mL) of the supernatant were transferred into PP tubes (15 mL, Starstedt) and stored at 4 °C in the dark until further analysis.

Depending on the instrumental limits of quantification (LQ) and SPM concentrations, 18 major and trace elements (Al, As, Ba, Co, Cr, Cu, Fe, Li, Mg, Mn, Na, Ni, P, Pb, Sr, Ti, V, and Zn) were analyzed by either ICP-OES (Agilent, 720 Series) or ICP-MS (Thermo X7, Series II). The LQ for each element/extraction

method were determined according to the standard method NF T 90-210(AFNOR 2009a) and are indicated in Table 3. The determination of LQ consists in the measurement of the target LQ concentration in a matrix similar to the samples (HNO₃ or HCl), in duplicate over five different days. The target LQ is validated if the mean value and associated standard deviation are within ±60% of the target LQ value. Blanks and certified reference materials (IAEA-158, marine sediment; LGC-6187, River

Table 3 Element concentrations (means and relative standard deviation) in total and HCl-extracted SPM fractions for the Ain, Arve, Bourbre, Fier, and Guiers Rivers and the Rhône River at Jons station (LQ: limit of quantification; *n*: number of SPM samples)

		Ain (<i>n</i> = 9)			Arve (<i>n</i> = 12)			Bourbre (<i>n</i> = 11)			Fier (<i>n</i> = 9)			Guiers (<i>n</i> = 13)			Jons (<i>n</i> = 16)		
	LQ	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)		
[Al] (g kg ⁻¹)	Total	0.01	29.2	<i>24</i>	47.4	<i>20</i>	31.8	<i>15</i>	35.6	<i>16</i>	28.7	<i>35</i>	55.4	<i>21</i>					
	HCl	0.001	1.55	<i>26</i>	2.56	<i>30</i>	2.08	<i>23</i>	2.01	<i>20</i>	1.29	<i>48</i>	3.72	<i>30</i>					
[As] (mg kg ⁻¹)	Total	0.10	9.30	<i>18</i>	14.1	<i>38</i>	9.21	<i>19</i>	6.50	<i>15</i>	7.93	<i>34</i>	11.8	<i>21</i>					
	HCl	0.01	1.55	<i>17</i>	3.74	<i>51</i>	2.76	<i>21</i>	1.10	<i>13</i>	2.20	<i>53</i>	3.91	<i>40</i>					
[Ba] (mg kg ⁻¹)	Total	2.50	102	<i>19</i>	296	<i>14</i>	203	<i>7</i>	161	<i>15</i>	142	<i>28</i>	292	<i>23</i>					
	HCl	0.62	17.2	<i>24</i>	47.2	<i>28</i>	54.1	<i>21</i>	29.8	<i>19</i>	16.5	<i>49</i>	36.6	<i>21</i>					
[Co] (mg kg ⁻¹)	Total	2.50	6.53	<i>20</i>	8.13	<i>21</i>	8.75	<i>18</i>	8.17	<i>17</i>	6.11	<i>42</i>	10.7	<i>18</i>					
	HCl	1.25	2.49	<i>14</i>	3.69	<i>18</i>	4.91	<i>24</i>	3.30	<i>17</i>	2.38	<i>43</i>	5.92	<i>26</i>					
[Cr] (mg kg ⁻¹)	Total	2.50	41.4	<i>24</i>	49.2	<i>25</i>	53.3	<i>11</i>	68.3	<i>14</i>	42.59	<i>38</i>	57.4	<i>13</i>					
	HCl	0.62	4.03	<i>22</i>	5.13	<i>47</i>	10.2	<i>31</i>	7.17	<i>9</i>	6.67	<i>108</i>	6.52	<i>13</i>					
[Cu] (mg kg ⁻¹)	Total	2.50	16.6	<i>26</i>	26.9	<i>67</i>	40.5	<i>26</i>	20.7	<i>25</i>	16.7	<i>69</i>	23.0	<i>25</i>					
	HCl	1.25	10.7	<i>29</i>	14.2	<i>75</i>	29.8	<i>31</i>	12.6	<i>26</i>	12.7	<i>96</i>	16.6	<i>30</i>					
[Fe] (g kg ⁻¹)	Total	0.002	17.0	<i>23</i>	20.4	<i>21</i>	16.8	<i>18</i>	18.5	<i>15</i>	14.3	<i>37</i>	25.6	<i>20</i>					
	HCl	0.001	3.97	<i>14</i>	6.57	<i>24</i>	5.31	<i>21</i>	4.79	<i>12</i>	3.59	<i>42</i>	9.57	<i>30</i>					
[Li] (mg kg ⁻¹)	Total	0.50	25.9	<i>24</i>	33.2	<i>23</i>	21.8	<i>20</i>	26.0	<i>17</i>	24.5	<i>48</i>	42.7	<i>27</i>					
	HCl	0.06	0.76	<i>14</i>	4.46	<i>42</i>	1.26	<i>16</i>	1.92	<i>17</i>	1.41	<i>47</i>	6.93	<i>38</i>					
[Mg] (g kg ⁻¹)	Total	0.01	5.19	<i>13</i>	9.80	<i>20</i>	4.16	<i>14</i>	9.61	<i>7</i>	6.07	<i>29</i>	11.1	<i>17</i>					
	HCl	0.001	2.22	<i>20</i>	4.13	<i>17</i>	0.99	<i>9</i>	3.65	<i>11</i>	2.52	<i>34</i>	4.41	<i>13</i>					
[Mn] (mg kg ⁻¹)	Total	2.50	415	<i>23</i>	470	<i>22</i>	592	<i>20</i>	504	<i>24</i>	420	<i>49</i>	599	<i>18</i>					
	HCl	6.25	342	<i>21</i>	356	<i>25</i>	507	<i>25</i>	419	<i>27</i>	321	<i>49</i>	472	<i>22</i>					
[Na] (mg kg ⁻¹)	Total	2.50	1850	<i>37</i>	13,300	<i>32</i>	4880	<i>11</i>	6370	<i>25</i>	5380	<i>29</i>	10,000	<i>17</i>					
	HCl	6.25	106	<i>12</i>	78.9	<i>54</i>	89.4	<i>24</i>	93.2	<i>11</i>	68.8	<i>38</i>	82.6	<i>22</i>					
[Ni] (mg kg ⁻¹)	Total	2.50	22.8	<i>24</i>	30.4	<i>30</i>	24.2	<i>22</i>	42.1	<i>16</i>	19.3	<i>43</i>	35.1	<i>17</i>					
	HCl	0.62	5.54	<i>16</i>	9.56	<i>40</i>	8.43	<i>26</i>	11.3	<i>12</i>	6.14	<i>48</i>	12.3	<i>19</i>					
[P] (g kg ⁻¹)	Total	0.50	0.86	<i>31</i>	1.11	<i>76</i>	1.47	<i>18</i>	0.85	<i>20</i>	1.01	<i>41</i>	0.77	<i>12</i>					
	HCl	0.003	0.39	<i>24</i>	0.78	<i>68</i>	0.93	<i>21</i>	0.51	<i>15</i>	0.38	<i>64</i>	0.58	<i>13</i>					
[Pb] (mg kg ⁻¹)	Total	0.10	19.1	<i>21</i>	20.3	<i>40</i>	32.4	<i>22</i>	14.9	<i>17</i>	15.9	<i>36</i>	23.2	<i>21</i>					
	HCl	0.01	12.8	<i>22</i>	11.8	<i>28</i>	25.8	<i>24</i>	9.54	<i>18</i>	11.27	<i>47</i>	17.8	<i>26</i>					
[Sr] (mg kg ⁻¹)	Total	2.50	175	<i>12</i>	335	<i>15</i>	198	<i>5</i>	336	<i>16</i>	297	<i>31</i>	293	<i>7</i>					
	HCl	31.2	150	<i>18</i>	262	<i>21</i>	145	<i>5</i>	288	<i>20</i>	246	<i>32</i>	245	<i>7</i>					
[Ti] (g kg ⁻¹)	Total	2.50	1720	<i>22</i>	2320	<i>21</i>	1600	<i>13</i>	1990	<i>14</i>	1396	<i>41</i>	2580	<i>17</i>					
	HCl	0.001	11.2	<i>20</i>	80.6	<i>80</i>	18.7	<i>16</i>	15.6	<i>25</i>	16.0	<i>48</i>	123	<i>45</i>					
[Vi] (mg kg ⁻¹)	Total	5.00	51.3	<i>26</i>	52.8	<i>23</i>	38.8	<i>21</i>	52.3	<i>19</i>	39.2	<i>46</i>	63.0	<i>21</i>					
	HCl	1.25	6.91	<i>13</i>	5.22	<i>20</i>	5.61	<i>23</i>	5.74	<i>10</i>	5.33	<i>45</i>	7.03	<i>18</i>					
[Zn] (mg kg ⁻¹)	Total	2.50	81.5	<i>30</i>	82.7	<i>37</i>	170	<i>24</i>	81.3	<i>22</i>	66.7	<i>57</i>	97.8	<i>21</i>					
	HCl	12.5	44.8	<i>50</i>	41.7	<i>58</i>	125	<i>31</i>	43.6	<i>26</i>	36.1	<i>65</i>	60.2	<i>25</i>					

Means of element concentrations = bold

Relative Standard Deviation = ital

Sediment) were systematically used to control analytical accuracy and precision. Concentrations of major and trace elements in extraction blanks were systematically below the LQ. According to the element, the accuracy of the total extraction method ranged between 94 and 134% (SI 1) and results were not corrected for extractions yields. Typical expanded uncertainties (i.e. 95% confidence intervals) calculated as international standard NF ISO 11352 (2012) were lower than 20%. Assessment of major and trace element concentrations in the residual fraction of SPM was calculated as the difference between concentrations obtained in the total fraction and HCl-extracted fraction. All trace and major element concentrations inferred from the residual fractions were expressed as a mass of element (μg or mg) related to the total (initial) mass of SPM (d.w.).

2.4 Data treatment

2.4.1 Particle size correction

Correcting major and trace element concentrations from particle size effect may be necessary because the SPM particle size distribution may not be conservative from the sources to the mixing station, due to deposition, erosion, and grain sorting during transport (Lacey et al. 2017). To implement this correction, we applied the methodology proposed by Gellis and Noé (2013). This method is aimed at estimating the element concentrations in the source SPM characterized by the same particle size distribution as the target SPM. It is based on the correlation of the element concentrations in the source samples against their D50, in order to obtain a particle size correction factor, when necessary. In the present study, if the linear correlation of the major or trace element concentrations (total and residual fractions) versus D50 was significant and had a high degree of correlation (Pearson test, $R > 0.7$, $p < 0.05$) for a given tributary, a correction factor was applied to the element concentrations in both the total and residual fractions as follows:

$$C_{i,j} = C_{i,j}^* - m_{i,j} \times (D50_j - D50_{\text{ref}}) \quad (1)$$

where $C_{i,j}^*$ is the initial major or trace element (i) concentration in a tributary (j), $C_{i,j}$ is the major or trace element concentration after particle size correction, $D50_j$ is the median diameter of the tributary sample, $D50_{\text{ref}}$ is the mean D50 of target SPM, and $m_{i,j}$ is the slope of the regression line of the element concentrations i versus D50 in the tributary j .

2.4.2 Range test

To assess major and trace element conservativeness during SPM transport from the five tributaries to the sampling station at the outlet of the Upper Rhône River (Jons), we checked that element concentrations in Jons samples remained within the range of the concentrations obtained in SPM samples of the

five tributaries. We used the method used by Gellis and Sanisaca (2018) consisting for each target sample that each tracer must be bracketed by the source samples' tracer concentrations ($< 10\%$ of the minimum and $> 10\%$ of maximum tracer concentration). Major and trace elements which did not meet this condition were removed from data treatment. This range test was performed on total and residual fractions element concentrations after particle size correction.

2.4.3 Selection of discriminant properties and estimation of the relative contribution of each tributary

Here, we applied the method first formalized by Collins et al. (1997), following a two-stage procedure. The method involved using the Kruskal-Wallis H test to determine suitable properties for discriminating potential sources. Then, a discriminant factor analysis (DFA) was applied to the parameters previously selected by the Kruskal-Wallis H test. A step-by-step algorithm, based on the minimization of the Wilk's lambda, was used to determine the most restrictive combination of parameters that discriminate the maximum number of sources. Then, a linear mixing model was built with the parameters selected by the DA. This model is based on the resolution of a linear equation system, assuming that the SPM flux at the downstream station is a mixture of SPM contributions from the S considered sources as follows:

$$C_i = \sum_{j=1}^S P_j \times C_{i,j} + E \quad (2)$$

where C_i is the concentration of the fingerprint element i in the target sample (e.g. SPM at the downstream station JON); P_j is the proportional SPM contribution of the distinct source j ; $C_{i,j}$ is the concentration of the fingerprint element i in SPM from the distinct source j ; and E is the error accounting for imperfect mixing and ignored sources (e.g. ignored tributaries). This equation assumes that SPM at Jons station are a mix of SPM from the Ain, Arve, Bourbre, Fier, and Guiers tributaries. This hypothesis agrees with the first estimation done by Launay (2014) who showed that for an entire hydrological year, 92% of all SPM fluxes at Jons station corresponded to SPM inputs from these five tributaries. Here, the weighting coefficients have to match the two following conditions:

$$\sum_{j=1}^S P_j = 1 \quad \text{and} \quad P_j \geq 0$$

The least squares method (R software, package nls) was applied in order to minimize the E value:

$$E2 = \left\| \sum_{j=1}^S P_j \times C_{i,j} - C_i \right\|^2 \quad (3)$$

2.4.4 Uncertainties of relative contributions

Finally, a Monte Carlo approach was used in order to take into account the variability of major and trace element

concentrations of each source and to quantify the uncertainties of each tributary SPM contribution obtained with the mixing model. A frequentist approach was preferred to a probabilistic approach since Davies et al. (2018) reported that probabilistic was better at dealing with non-conservative tracers. For each JON sample, a total of 1000 linear equation systems were generated by randomly selecting one $C_{i,j}$ value among all the concentrations measured for each tributary. The proportional SPM contribution of each source was expressed as the mean of the 1000 mixing model iterations. After verifying that the distributions of the model outputs are approximately normal (Shapiro-Wilk test), a confidence interval (C.I.) with a 95% confidence level based on the standard normal distribution obtained from the 1000 iterations of the mixing model was estimated as follows:

$$C.I. = \left[m - k \times \frac{s}{\sqrt{n}}; m + k \times \frac{s}{\sqrt{n}} \right] \quad (4)$$

where m and s are the mean and standard deviation of the n iterations ($n = 1000$) and k is equal to 1.96 for a confidence level of 95%.

2.5 1-D numerical hydro-sedimentary model and simulation

The Rhône River system was modeled using MAGE, a 1-D numerical hydrodynamic code for unsteady flows, coupled with Adis-TS, an advection–dispersion resolution for solute and suspended sediment transport with deposition and erosion (Andriès et al. 2011; Launay et al. 2019). The flow resistance factors of the multi-reach network were calibrated and tested using measured longitudinal water profiles. Cross-sectional profiles of the major tributaries, such as the Arve and Ain Rivers, are included from their confluence with the Rhône to their nearest water and SPM monitoring station. Other tributaries are simply represented as local inputs.

The eight hydropower schemes located upstream Jons station, and their regulation rules were integrated in the model (Launay et al. 2019). The regulation rules include minimum compensation discharge in the by-passed Rhône River, maximum discharge through the plant and bypass canal, and stage–discharge relation in the reservoir. Theoretical rules like this do not reflect the real fluctuations due to hydropeaking and maintenance during routine operation. Input data include water discharges monitored by the Rhône national company (CNR), FOEN (Swiss Hydrological Services) and Regional Directorate of Environment, Development and Housing (DREAL). The SPM concentrations were also provided for the five investigated tributaries (Arve, Fier, Guiers, Bourbre and Ain Rivers). The SPM concentrations of the Arve and Fier Rivers have been continuously recorded at turbidity

stations operated by the OSR since 2012 and 2014, respectively (Thollet et al. 2018). Temporary stations have been operated in the Guiers, Bourbre, and Ain Rivers during limited periods. When SPM measurements were not available as calibrated turbidity records, we used SPM concentrations inferred from discharge-SPM concentration relations (sediment rating curves) estimated by Launay (2014). The 1-D numerical model of the Rhône River was used to decompose the flow hydrographs according to water inputs from the different tributaries. This decomposition process provided the relative contributions of the water flowing at the downstream monitoring station at Jons. The SPM fluxes were equally decomposable. All the details on the 1D hydro-sedimentary model are available in Launay et al. (2019).

For each SPM sampling period on the Rhône at Jons station (Table 2), a simulation over the sampling period was used to decompose the flow hydrograph according to water discharge (Fig. 2a) and the SPM fluxes, accounting for SPM propagation and sediment deposition and erosion. The SPM concentration distributions were considered homogeneous, with a mean diameter $d = 20 \mu\text{m}$ (Fig. 2b). Calibration of the hydro-sedimentary parameters was performed according to Guertault (2015). During the dam flushing event, the SPM re-suspension from the hydropower scheme reservoirs was calibrated against the SPM records measured downstream of the dams.

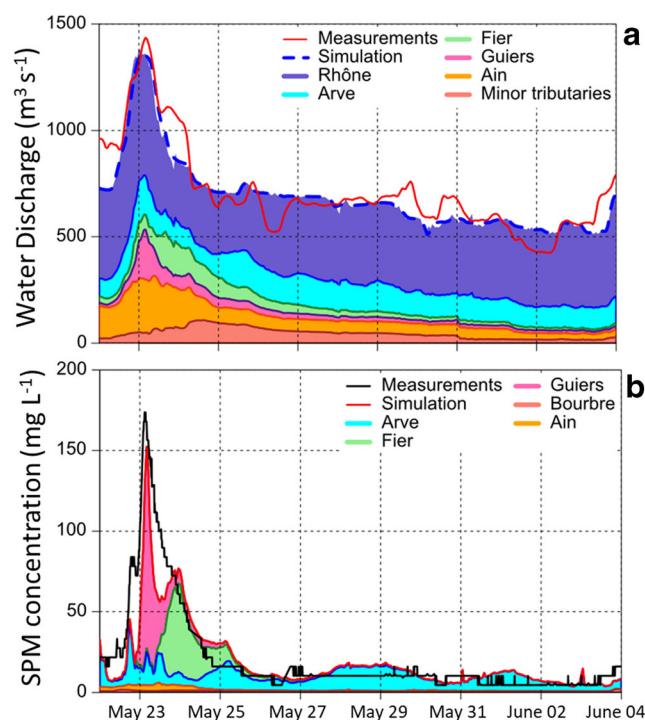


Fig. 2 Example of results from the 1-D hydro-sedimentary numerical model in the upper Rhône River: (a) discharge hydrograph decomposition; (b) SPM concentration decomposition

3 Results

3.1 Major and trace element concentrations in total and HCl-extracted SPM fractions

Means and standard deviations of element concentrations in SPM from the five potential sources (Ain, Arve, Bourbre, Fier, and Guiers Rivers) and at the mixing station are detailed in Table 3. The lowest total element mean concentrations were measured on SPM from the Guiers and Ain Rivers. In contrast, the highest total element mean concentrations were obtained on the Arve, the Bourbre, and the Fier Rivers. Total mean concentrations of Al, Co, Fe, Li, Mg, Mn, Ti, and V were higher in SPM of the Rhône at Jons station than in the five tributaries.

Similar to total element concentrations, the Guiers River displayed the lowest HCl-extracted mean concentrations for several elements. The Arve and Fier Rivers displayed only the one highest HCl-extracted SPM concentration for As and Sr, respectively. In contrast, the Bourbre River displayed the highest HCl-extracted mean SPM concentrations for Ba, Cr, Cu, Mn, P, Pb, and Zn. On the Rhône at Jons station, HCl-extracted concentrations of Al, As, Co, Fe, Li, Mg, Ni, Ti, and V were higher than in the five tributaries.

The mean HCl-extracted element fractions in SPM from the Rhône River and its tributaries represented from 1% to 86% of the total element mean concentrations depending on the element (Fig. 3). The HCl-extracted fraction ranged from 1% to 25% for Na, Ti, Al, Li, V, Cr, Ba and As. For Fe, Ni, Mg, and Co, the mean contribution of the HCl-extracted fraction was higher, ranging from 29% (Fe) to 47% (Co). Finally, Zn, P, Cu, Pb, Mn, and Sr displayed the highest HCl-extracted fractions, ranging from 57% to 86%.

The PCA analysis on total and HCl-extracted element concentrations (SI 3) suggests that the variabilities of the element concentrations in SPM of the Bourbre, Ain, and Fier Rivers were lower than those observed for the Guiers and Arve River. At this point, this analysis also shows that the raw (without

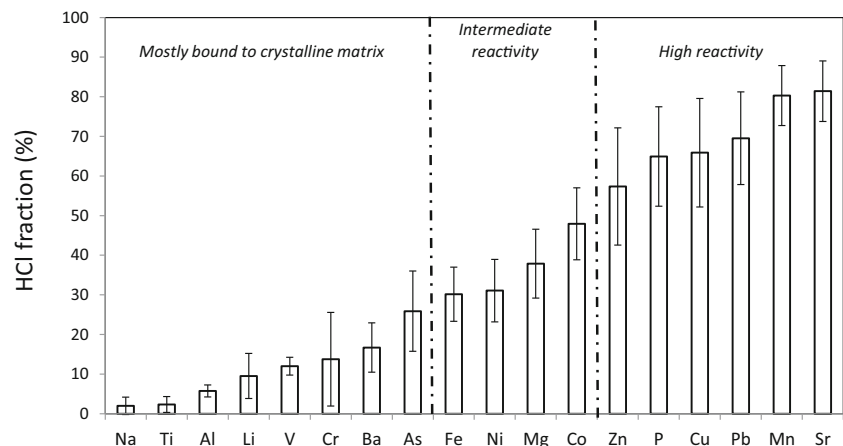
particle size correction and tracer selection) fingerprint of the Bourbre River seems to be distinct from those of the Ain and Guiers Rivers and from those of the Arve and Fier Rivers.

3.2 Effect of particle size distribution on element concentrations and range test

When the correlation between element concentrations and particle size distribution was significant (Pearson test, $R > 0.7$, $p < 0.05$; SI 2), a correction factor was applied. As an example, plots of element concentrations versus median particle diameter (expressed as the logarithm of D50) are showed for total and residual As concentrations in SPM samples from the Ain River (Fig. 4, Table 4, SI 3). In SPM samples from the Arve River, total concentrations for eight elements (Co, Cu, Fe, Mg, Mn, Pb, V, and Zn) displayed significant relationships with particle size, and only Mg and V concentrations in the residual fraction showed a significant relationship with particle size. In the Bourbre River, total concentrations for 12 elements (Al, As, Ba, Co, Cr, Fe, Li, Mg, Mn, Ni, Ti, and V) were significantly correlated with particle size. A significant relationship was also displayed for these elements in the residual fraction except for Ba, Mn, Cu, and Zn (Table 4). For the Fier River, 13 elements (Al, As, Ba, Co, Cr, Cu, Fe, Li, Mn, Pb, Ti, V, and Zn) and 11 elements (Al, As, Cr, Cu, Fe, Li, Mg, Pb, Ti, V, and Zn) were significantly correlated to particle size in the total and in the residual fractions, respectively. For all these elements and according to the SPM fraction considered (total or residual), a correction factor was applied (Table 4). For the Guiers River, only Li was correlated with particle size in total and residual fractions.

The conservativity test performed for each individual sample at Jons station showed that all elements were within the acceptable range of the corresponding elements concentrations in SPM samples of the five potential sources, except for Al, Ba, Fe, and Li in the total fraction and, for Al, Li, and Mg in the residual fraction (Table 4).

Fig. 3 Mean contributions and standard deviation ($n = 76$) of the HCl-extracted fraction of total element concentration in SPM for all stations (Rhône at Jons and 5 main tributaries)



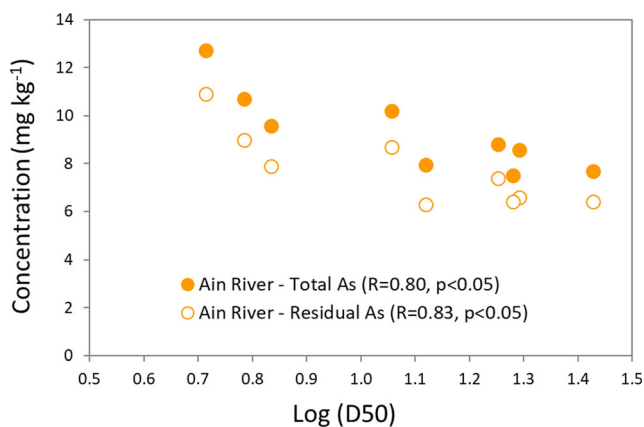


Fig. 4 Relation between As concentrations in total and residual SPM fractions as a function of median particle size (D50) of SPM on the Ain River

3.3 Discrimination of tracers

The DFA applied to element concentrations in SPM sources, previously corrected by particle size, indicated that five elements allowed to correctly discriminate the five tributaries. For the total fraction, the selected elements were Mg, Ni, Sr, Mn, and As (Table 4). For the residual fraction, Ba, Fe, Ni, Mg, and V were selected as the most discriminating elements for all the samples except JON_16 and JON_17 for which Ba, Cu, Ni, and Sr were selected as the most discriminant elements (Table 4).

3.4 Relative contribution of SPM sources: Fingerprinting and 1-D hydro-sedimentary approaches

The concentrations of the five elements for the total (Mg, Ni, Sr, Mn, and As) and residual (Ba, Fe, Mg, Ni, and V or Ba, Cu, Ni, and Sr) fractions were used separately in the mixing model to estimate the SPM contributions of each tributary for 15 SPM samples retrieved at Jons station during three contrasting hydrological conditions: base flow, flood, and dam flushing event (Fig. 5).

In general, the relative contributions estimated from element concentrations in the residual fraction are in better agreement with the 1D hydro-sedimentary model than the relative contributions estimated from element concentrations in the total fraction. For four of the five SPM samples retrieved during the dam flushing event, the relative SPM contributions were mainly represented by the Arve River, whether using total or residual element concentrations (Fig. 5a). The contributions of the Arve River were lower for the fingerprinting method using total element concentrations (50% to 76%) than when using element concentrations in the residual fraction (57% to 88%). For JON_21 sample, both fingerprinting methods estimated similar contributions (33 and 37%) for the Guiers River. However, the fingerprinting model using

the total fraction estimated a main contribution of the Ain River (42%) with a low contribution of the Arve River (8%). By contrast, the fingerprinting model using the residual fraction estimated higher and similar contributions from the Arve (33%) and the Bourbre (26%) Rivers.

For base flow conditions (Fig. 5b), the fingerprinting method using the residual fraction displayed a main contribution (75% to 81%) from the Arve River. These contributions are in agreement with those obtained with the 1D-hydro-sedimentary model that estimated that SPM in transit at Jons station were mainly originated from the Arve River (70% to 91%). Concerning the fingerprinting method using the total fraction, the Arve River was also determined as the main source of SPM, but with lower contributions, ranging from 44% to 58%. The lower contributions of the Arve River were counterbalanced by a higher contribution of the Fier River, ranging from 26% to 33%.

For flood events (Fig. 5c), both fingerprinting approaches using total and residual fractions estimated a lower contribution of the Guiers River for JON_01 sample, with a relative contribution of 5% and 8%, respectively, in comparison to 49% according to the 1-D hydro-sedimentary model. The fingerprinting method using the total and residual fractions estimated a higher contribution for the Fier (44% and 36%, respectively) and Arve (26% and 32%, respectively) Rivers. For JON_15 sample, both fingerprinting approaches estimated a main contribution from the Fier River (74% and 67%), which is consistent with the contribution of the Fier River (55%) estimated by the 1D hydro-sedimentary model. However, while the 1D-hydro-sedimentary model estimated that the Ain River was the second main source of SPM (28%), both fingerprinting methods estimated a lower contribution from the Ain River (< 3%), and a higher contribution from the Arve River (18% and 30%). For JON_23 sample, the fingerprinting approach using the residual fraction estimated that the Arve River was the main source of SPM (73%), which is in agreement with the relative SPM contribution estimated for the Arve River by using the 1D-hydro-sedimentary model (92%). By using total element concentrations, the fingerprinting method estimated that the contribution of the Arve River was lower (60%), with a higher contribution from the Fier River (20%).

For JON_30 sample, the fingerprinting approach using the residual fraction displayed that SPM contributions were equally distributed between the Arve (33%), Guiers (27%), Fier (18%), and Bourbre (17%) Rivers. The fingerprinting method using the total fraction estimated similar contributions for the Fier River (34%), Arve River (20%), and Ain River (30%). Similar and according to the 1D-hydro-sedimentary model, relative contributions were fairly distributed between the Fier (35%), Ain (33%), and Guiers (24%) Rivers.

Table 4 Correction factors applied for elements displaying a significant correlation between log (D50) versus element concentrations in total and residual fraction of SPM for the Ain, Arve, Bourbre Fier, and Guiers Rivers, results of conservativity test, and discriminant factor analysis obtained after particle size correction for the two studied fractions

Element	Element concentration in residual fraction															
	Total element concentration		Ain River		Arve River		Bourbre River		Fier River		Guiers River		Excluded after conservativity test		Cumulative source correctly classified (%)	
	Correction factor	Ain River	Arve River	Bourbre River	Fier River	Guiers River	Excluded after conservativity test	Ain River	Arve River	Bourbre River	Fier River	Guiers River	Excluded after conservativity test	All samples	All samples excepts JON_16 and JON_17	Cumulative source correctly classified (%)
Al		1.36	1.25				Jon_06, 08, 16, 17, 25, 26, 27, 28	0.99								68
As		0.99	1.43	1.19			JON_06, 08, 16, 17, 26									86
Ba		1.14	1.18													
Co		1.31	1.47	1.22					1.18							
Cr		1.17	1.17						1.07	1.10						
Cu		3.04	1.00						1.13	1.20						93
Fe		1.34	1.48	1.21			JON_16, 17	1.13	1.17	1.40					91	
Li		1.56	1.25	1.12			JON_06, 16, 17		1.18	1.17	1.13		JON_16, 17			
Mg		0.99	1.20	1.35					1.15	1.01			JON_16, 17		52	
Mn		1.38	1.48	1.35												
Na																
Ni			1.66						1.29						95	91
P																
Pb		1.01	1.22							1.01						
Sr																98
Ti			1.10	1.12					1.10	1.12						
V		1.32	1.61	1.30				1.13	1.19	1.21					98	
Zn		1.00	1.33						1.22	1.18						

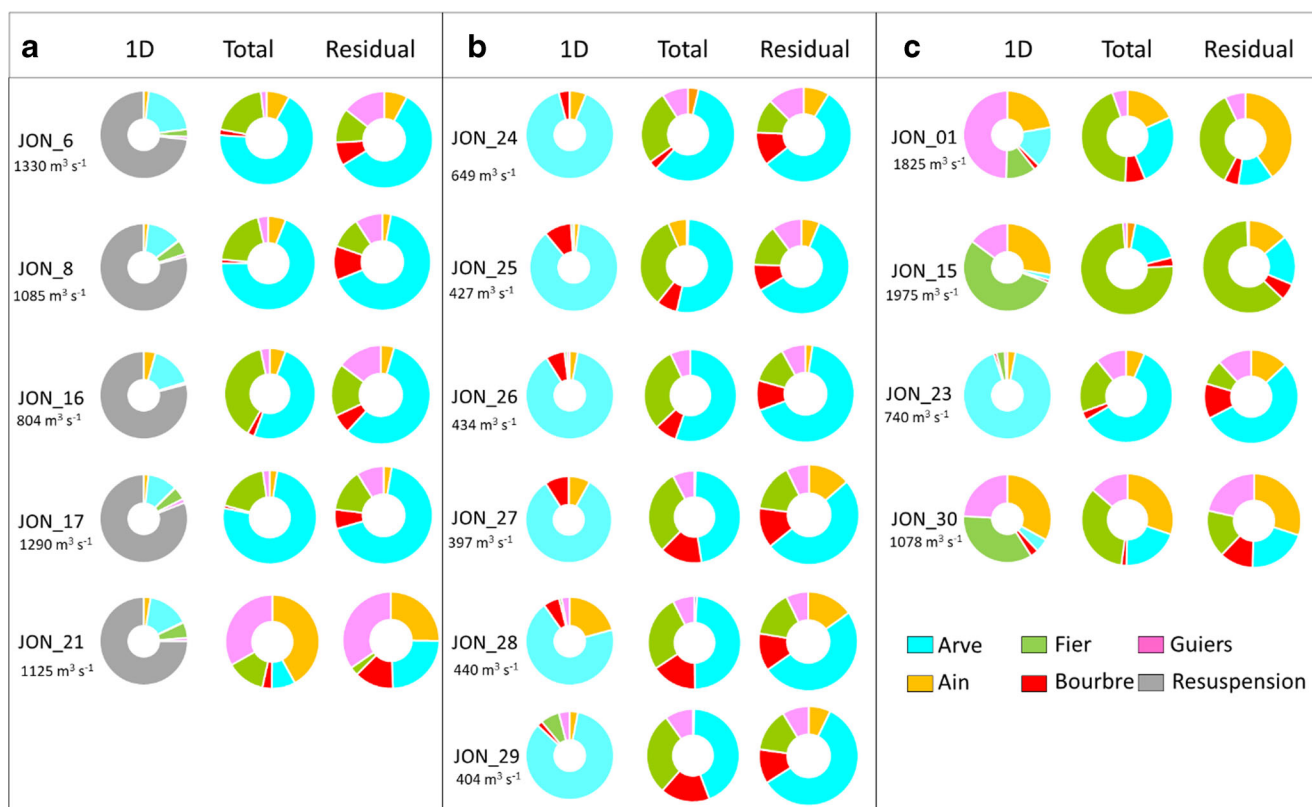


Fig. 5 Relative SPM contribution at Jons station (Rhône River) for SPM collected during dam flushing (**a**), base flow (**b**), and flood events (**c**), estimated by the 1D hydro-sedimentary model and by the fingerprinting approach using element concentrations in the total or in the residual SPM fractions

The results of the Monte Carlo simulations showed that confidence intervals were very low (< 4%) for each contribution, whether using total or residual fractions (Fig. 6).

4 Discussion

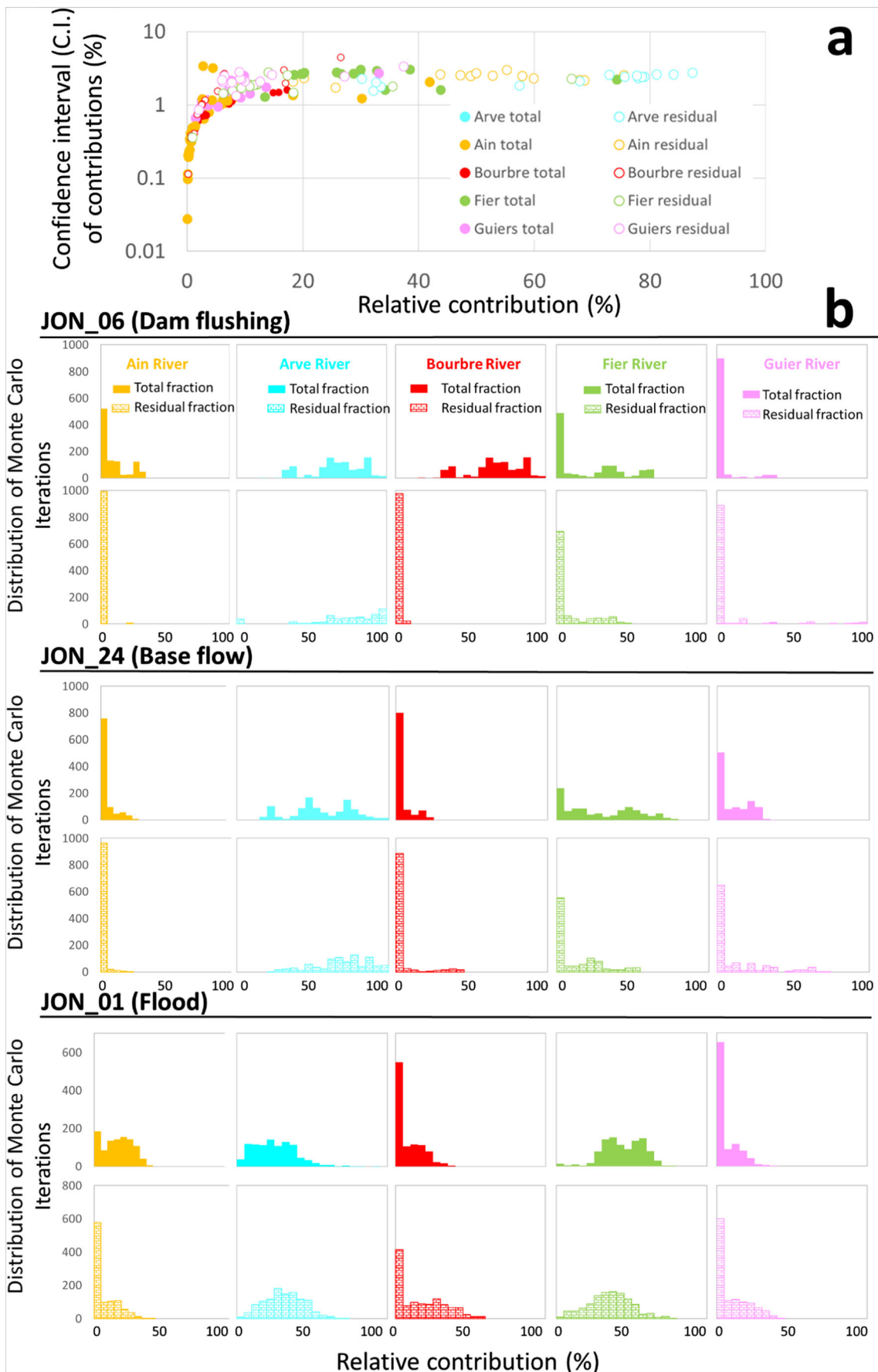
4.1 Relevance of using element concentrations in the residual fraction of SPM to assess sediment sources

In SPM from the Upper Rhône River, the HCl-extracted fraction represented less than 25% of the total concentration for Na, Ti, Al, Li, V, Cr, Ba, and As, suggesting that the total concentrations of these elements could be used as conservative tracers, due to their low reactivity. This result could be surprising for Na, since this element is widely recognized to be non-conservative and is often removed from sediment fingerprinting studies, notably in the fine particles (< 63 μm) of sediment samples (Gholami et al. 2019). As mentioned by Negrel et al. (2015), soil weathering causes a rapid flushing of Na as dissolved ions. This non-conservative behavior results in the depletion of particulate Na during SPM transport, which is mainly derived from weathering of aluminosilicate phases (dissolution of plagioclase) and evaporites (Negrel et al. 2015). However, through X-ray diffraction (XRD)

analysis, Slomberg et al. (2016) reported that mineral composition of SPM sampled at Jons station was mainly represented by five minerals, including the albite (a feldspar plagioclase mineral). If these minerals are still present in SPM at Jons station, this suggests that they are resistant to weathering. This hypothesis is consistent with the study of Aström et al. (1998) who demonstrated that in soils and sediment, elements such as Na are only to a limited extent extracted in acid aqua regia (1 HCl/3 HNO₃, v/v), so mainly associated with “resistant” minerals such as feldspar. Thus, the low reactivity of Na (a major constituent of albite mineral) in SPM in Upper Rhône tributaries is consistent since HCl 1 M extraction is weaker than an aqua regia extraction.

A second group of elements, Fe, Ni, Mg, and Co, displayed an intermediate reactivity, with a HCl-extracted fraction ranging from 29% to 47%, which is consistent with the results obtained by El Nemr et al. (2006), with HCl-extracted fractions of 43% for Fe and 58% for Ni in fine sediments of the Suez Gulf. This higher reactivity suggests that attention should be paid before using the total concentrations of these elements to assess sediment sources since they are subjected to change during their transport.

For Zn, P, Cu, Pb, Mn, or Sr, the use of total concentration as conservative tracers may be even more problematic due to their high reactivity, since the HCl-extracted fraction was



◀ **Fig. 6** Confidence interval (C.I.) of each relative SPM contributions (%) obtained after Monte Carlo simulations by using element concentrations of total and residual SPM fractions (a). Example of Monte Carlo distribution (1000 iterations) for JON_06 (Dam flushing), JON_24 (Base flow) and Jon_01 (Flood) samples (b)

above 50% (Fig. 3). These results are comparable to those obtained by Dabrin et al. (2014) on SPM from the Garonne River, with contributions of the HCl-extracted fraction representing between 55% and 78% of the total Cu and Pb concentrations, respectively. Indeed, Cu, Pb, or Zn concentrations in SPM are affected by pollution or anthropogenic-based processes, leading to an increase of concentrations in the particulate available fractions (Choi et al. 2012).

Since the tracer selection for SPM source discrimination was only done on a statistical basis (DFA), our results showed that for total element concentrations, four (Mg, Ni, Mn, and Sr) of the five elements selected by the DFA (Mg, Ni, Sr, Mn, and As) were intermediate (Mg and Ni) to highly reactive (Mn and Sr; Fig. 3). This highlights that a common statistical analysis enabled selecting the best combination of elements, without taking their potential reactivity into account. On the contrary, As was mainly bound to the crystalline matrix and was also selected as a fingerprinting tracer. Arsenic is commonly used in fingerprinting approaches (e.g. Theuring et al. 2015), corroborating the idea that total concentrations of As could be used as fingerprinting property.

For the residual fraction, the DFA selected five elements (Ba, Fe, Mg, Ni, V, or Ba, Cu, Ni, and Sr) including two low reactive elements (Ba and V), which confirms the relevance of using elements mainly associated with the crystalline matrix of the sediment. The DFA analysis applied on the residual fraction singled out highly-reactive elements such as Sr and Cu, which were determined as highly reactive elements in the total fraction, with HCl-extracted contributions of 82% and 65%, respectively. Hence, highly reactive elements that should not be considered relevant for fingerprinting approaches could be considered discriminant fingerprinting properties by using the residual fraction concentrations.

Our approach could be compared to another promising method proposed by Maher et al. (2009) who performed magnetic measurements on untreated and acid-treated samples (HCl 12 M) of river channel, estuarine, and inner shelf sediments. Despite the measurements of only four magnetic parameters (magnetic susceptibility, anhysteretic remanent magnetization, saturation remanence, and remanence ratios), they showed that the use of acid-treated samples eliminated any influence of post-depositional processes that may modify the initial signature of sediment source. As previously mentioned by Collins et al. (1997), since the fine particle fractions are more geochemically active, they are likely to discriminate sources more robustly. However, this benefit can be counterbalanced because these finer particles are also more

subject to transformation and non-conservative behaviors during transport. Most fingerprinting studies use major and trace element concentrations in the total fraction of SPM or sediments after a total digestion (typically by using HCl, HNO₃ and HF or aqua regia; e.g. Le Cloarec et al. 2011; Evrard et al. 2011). Authors of these studies assumed that the total element concentrations in SPM or sediments are conservative during transport, settlement, and remobilization. However, several elements are known to be highly reactive, such as flocculation or precipitation of Fe and Mn, co-precipitation with Fe/Mn oxides of trace metals (e.g. As and V), dissolution, or formation of authigenic iron sulfide in reduced or anoxic conditions (e.g. Morse 2002). Despite their potential non-conservative behaviors, these elements are still used in the literature and in total fraction of our study. Collins et al. (2017) mentioned that despite risks of misinterpretation, some studies have included tracers prone to transformations such as P, as identified by Owens et al. (2000). To assess element concentrations in the most conservative fraction of SPM or sediment, several extraction methods were developed, allowing extracting major and trace elements from different matrices (e.g. exchangeable fraction, carbonates, Fe/Mn oxides, and organic matter) of the sediment (Tessier et al. 1979). Extraction of elements from sediment using HCl 1 M is empirically defined as an extraction allowing to assess elements potentially bioavailable (Bryan and Langston 1992). Hydrochloric acid is a strong acid that can solubilize the most reactive phases of the sediment: its reducing properties allow extracting elements from Fe/Mn oxides, and it is efficient to decompose labile organic matter and amorphous sulfides (Snape et al. 2004). The concentration of 1 M HCl is sufficient to buffer the dissolution of carbonates but is low enough to limit the extraction of the residual fraction (Snape et al. 2004). As a result, the difference of element concentrations between total extraction (by using HCl, HNO₃ and HF) and 1 M HCl extraction represents element concentrations in the residual fraction that is mainly primary and secondary minerals of SPM and sediment. This fraction is considered conservative under environmental conditions encountered in the different compartments of the river and avoids any influence of post-depositional processes that may alter the initial signature of SPM.

By using HCl 1 M, all the particulate reactive carrier phases of elements that could be modified at short or long temporal scale are broken down (Sutherland et al. 2002). This includes adsorbed elements, elements bound to carbonates, to Fe/Mn oxide/hydroxides, and to sulfides. The extraction using HCl 1 M is used to assess acid volatile sulfides (AVS) and simultaneously extractable metals (SEM), conditioning the mobility and toxicity of elements in anoxic sediments (Di Toro et al. 1992). This ability to break down sulfides is interesting to assess sediment sources during flood or dam flushing events, as it overcomes transformed properties of elements co-precipitated with iron and sulfide under anoxic conditions

(e.g. sediments stored behind dams). Moreover, Aagemian and Chau (1976, 1977) demonstrated that the use of 0.5 N HCl satisfied the essential requirement of minimal dissolution of the silicate detrital lattice and produced the highest contrast between anomalous and background samples. However, Cooper and Morse (1998) reported that HCl 1 M was not sufficient to break down Cu and Ni sulfide minerals and particulate organic matter. Thus, it would be interesting to improve our methodology, as inferred from the residual fraction of SPM, by adjusting a more appropriate soft extraction allowing to also break down particulate organic matter and/or by adapting soft extractions according to each element. In this way, Dold et al. (2003) demonstrated that H₂O₂ extraction (in water bath) was able to dissolve organic matter and supergene Cu-sulfides.

4.2 SPM source apportionment: Strengths and weaknesses of fingerprinting approach compared to 1D hydro-sedimentary approach

Five samples of SPM were retrieved at Jons station during the dam flushing event (Jon_6, JON_8, JON_16, JON_17, and JON_21; Table 1, Fig. 5). Both total and residual fractions fingerprinting approaches estimated that SPM from Jons station were mainly constituted by SPM from the Arve River, except for one sample (JON_21). This result is consistent with the literature since the Arve River is known to carry about 500,000 t year⁻¹ of flysch and molasses particles, 50% of which being trapped behind the Verbois dam (Grimardias et al. 2017). For this event, the hydro-sedimentary model simulated the SPM contribution of each tributary and allowed to estimate that the main input of SPM at Jons station originated from dam flushing re-suspension, with contributions ranging from 69% (JON_16) to 79% (JONS_8 and JON_17; Fig. 5a). This may suggest that the relative contribution of re-suspended sediment estimated by the 1D-hydro-sedimentary model may be attributed to SPM coming from the Arve River. Indeed, by adding the SPM flux from the Arve River and the re-suspended sediment contribution, the 1D hydro-sedimentary model estimated that the Arve River contributed about 90–95% of SPM fluxes at Jons during this event. For JON_21 sample, the relative SPM contributions estimated from the fingerprinting and 1D hydro-sedimentary approaches do not match. This could be explained by the sampling method, since this sample was collected with a particle trap exposed during the first days of the dam flushing event. We can assume that, in the first days of the event, the particle trap integrated re-suspended sediment from different origins and/or that it induced a bias during SPM sampling in these unusual conditions (Masson et al. 2018). Nonetheless, except for this sample, our results highlighted the complementarity of the fingerprinting and hydro-sedimentary approaches: while the 1-D hydro-sedimentary model allowed to estimate the

contribution of sediment re-suspension, the fingerprinting approach using residual fraction allowed to determine its primary origin.

For all the six samples retrieved during base flow conditions (Fig. 5b), the relative contribution of the Arve River obtained using the residual fraction (75–81%) was in better agreement with results obtained by the 1D hydro-sedimentary (70–91%) model than those obtained using the total fraction (44–58%). Indeed, fingerprinting using the total fraction estimated that about 26% to 33% of the contribution was represented by SPM from the Fier River. By contrast, the results for the fingerprinting approach using the residual fraction of SPM displayed relatively high contributions (75–81%, Fig. 6) from the Arve River. Indeed, in base flow conditions, Rose et al. (2017) demonstrated that SPM were represented by autochthonous SPM mainly made of particulate organic matter and co-precipitated metals. This corroborates the idea that fingerprinting properties in SPM sampled during low flow conditions do not reflect the composition of SPM coming from erosional sources, suggesting using element concentrations in the residual fraction, as it is more representative of eroded material. Thus, according to the fingerprinting approach using element concentrations in the residual fraction, SPM at Jons station mainly originated from the Arve River during base flow discharge. To date, no accurate estimation of SPM flux contributions for a given time for the Upper Rhône basin has been published yet. Based on discharge and SPM measurements at Jons station and tributaries, Launay (2014) estimated that for an entire hydrological year, the Arve River contributed to 45% of SPM fluxes despite a low water discharge contribution (14%).

At last, for samples retrieved during flood events (Fig. 5c), the fingerprinting approach using the residual fraction estimated a contribution of the Arve River similar (73%) to the 1D hydro-sedimentary model (91%) for JON_23 sample. In comparison, the fingerprinting model using the total element concentrations estimated a contribution of 60% for the Arve River. For JON_1 sample and according to the 1-D hydro-sedimentary model, the period of sampling was characterized by a main SPM input from the Guiers River with a contribution of 49%. However, both fingerprinting approaches and 1D hydro-sedimentary model do not match each other to estimate the SPM contribution from the Guiers River. For JON_15 sample, results obtained for both fingerprinting approaches were in agreement with contributions from the Fier River of 74% for the approach using total element concentrations and 67% for the approach using element concentrations in the residual fraction of SPM. This is in agreement with the SPM contribution estimated by the 1-D hydro-sedimentary model with a main contribution of the Fier River (55%). By contrast, both fingerprinting approaches estimated that the Arve River was the second main input of SPM (18% and 30%) with a low contribution of the Ain River (<3%), while the 1-D hydro-

sedimentary model estimated that the second contributor was the Ain River (28%). Two hypotheses could explain these differences: first, the underestimation of the estimated SPM flux by the fingerprinting models suggests that during these high hydrological conditions, SPM from the Ain River were not well homogenized in the cross-section of the Rhône downstream of the confluence between the Rhône and the Ain Rivers. Indeed, the sampling station at Jons is located in the middle of the Rhône River, upstream of its separation in two main channels (the “Miribel” and “Jonage” channels) and located only at 6 km downstream the confluence of the Rhône and Ain Rivers (Launay 2014; Fig. 1). We hypothesized that during floods of the Ain River, particles are still flowing on the right side of the Rhône River and are poorly sampled by the particle trap at the Jons station. This hypothesis is corroborated by the study conducted by Bouchez et al. (2010), which showed that lateral mixing downstream confluences in large rivers are at least of several tens of kilometers. The second hypothesis is related to an overestimation of the Arve River input: since this tributary is the main input of SPM through an entire hydrological year (Launay 2014), we could assume that bed sediment is mainly represented by SPM from the Arve River. During high water discharge, a fraction of SPM sampled at Jons station may be represented by bed-sediment remobilization with a composition strongly derived from the Arve River. For JON_30 sample, the 1-D hydro-sedimentary model estimated that SPM contributions were equally distributed between SPM inputs from the Fier (35%), Ain (33%), and Guiers (24%) Rivers. During this high hydrological condition, the estimation of SPM contribution through both fingerprinting approaches was more uncertain. The two fingerprinting approaches estimated a non-negligible contribution from the Guiers River (14% and 27% respectively) and from the Fier River (34 and 18%). However, in the same way as for JON_15 sample, both fingerprinting approaches overestimated the Arve River contribution (20% and 33% respectively for total and residual fractions). During flood, when SPM contribution is not predominantly represented by one tributary, the fingerprinting approach slightly differed from the 1-D hydro-sedimentary model, probably (i) by overestimating the Arve River contribution due to re-suspension of bed sediment (mainly composed by sediment from the Arve River) and (ii) by a sampling method not suitable to take into account SPM inputs from the Ain River.

5 Conclusion

Trace and major element concentrations were determined in SPM sampled during contrasting hydrological conditions in the Upper part of the French Rhône River and its main tributaries, the Arve, Ain, Bourbre, Fier, and Guiers Rivers. Using a total (HCl, HNO₃, and HF) and a soft extraction (HCl 1 M

of elements in SPM, we highlighted the potential reactivity of major and trace elements in SPM for a large-sized catchment such as the Upper Rhône hydrosystem. Through the use of a mixing model combined with a Monte Carlo procedure, using total and residual element concentrations, we estimated the SPM contributions of the main tributaries of the Upper Rhône River at Jons station for contrasting hydrological conditions. This methodology is proposed to avoid taking into account the most reactive part of the sediments whose geochemical signature can change during transport. The cross-validation of the results allowed a critical interpretation of the results obtained by the use of fingerprinting approach. By comparing the mixing model results of both fingerprinting approaches with those of a 1-D numerical hydro-sedimentary model, we showed that it was more relevant to use element concentrations in the residual fraction of SPM to identify SPM sources at large-watershed scale. This study demonstrated that in base flow conditions, SPM fluxes at Jons station were mainly made of SPM originating from the Arve River. We further showed that this fingerprinting approach could discriminate the SPM origin of the sediment re-suspended during dam flushing events and was complementary to the 1-D hydro-sedimentary model for such events. A next step will be to validate this geochemical approach by applying it to the entire Rhône watershed and to sediment cores to assess the present and historical relative SPM contributions of the Rhône tributaries and sub-catchments to the Mediterranean Sea.

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