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Chronology and sources of trace elements accumulation in the Rhône pro-delta sediments (Northwestern Mediterranean) during the last 400 years

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Abstract :

The Rhône pro-delta sediments receive the particulate inputs from the Rhône River, the largest freshwater discharge of the Western Mediterranean Sea. Trace element (TEs: Ag, Cd, Co, Cr, Cu, Ni, Pb, and Zn) concentrations and stable Pb isotope ratios were determined along a 7.7-meter-long sediment core collected from the Rhône prodeltaic lobe, archiving the deposits of the last 400 years. Trace element mean concentrations during the pre-industrialized era (before 1850 AD) were no different from the average composition of Earth's upper crust, except for Cr. Principal component analysis, performed on TEs and tracers, suggests three origins for TEs that we identified as (i) marine biogenic material (Ag, Cd), (ii) ultramafic rocks (Li, Cr, Ni), and (iii) other clay minerals (Co, Cu, Pb and Zn). During the pre-industrial period, several changes in the TE/Al ratios coincided with flood events or/and modifications in the channelization of the Rhône River mouth. Some frequencies in the TEs temporal variations allow us to hypothesize some influence of climate variation. Sediments deposited after 1850 AD exhibited concentrations of certain TEs exceeding 1.3 to 3.5 times the concentrations of the pre-industrial era (Ag > Cd > Pb > Cu > Zn). Principal component analysis, performed on post-1850 AD data, allows to distinguish pristine and "anthropogenically impacted" TEs. Pristine TEs, such as Co, Cr, and Ni, are linked with clay markers (Al, Li and Fe), Co and Ni are associated with Mn-enrichment, whereas Cr is correlated with Fe. "Anthropogenically impacted" elements consist of Ag, Cd, Cu, Pb, and Zn. Vertical profiles show that TE contamination has increased dramatically after the IInd World War and reached a maximum between 1970 and 1980, consistently with anthropogenic atmospheric emissions changes. Sediment deposited on the Rhône prodelta at the beginning of XVIIth Century have 206/207Pb ratios close to values of natural Pb (1.200), whereas sediments deposited after 1850 AD are clearly contaminated by European gasoline and industrial Pb pools (<1.180). In addition, peak-events can be related to flooding periods, natural and man-induced channelization of the Rhône River mouth, or diagenetic processes.

Highlights

► Trace elements are accumulated in the Rhône prodelta sediments. ► Preindustrial sediments recorded floods and channelization of the Rhône River. ► Contamination reached a maximum between 1970 and 1980 AD.

1. Introduction

As stated by the international Geotraces program (www.geotraces.org), many trace elements (TEs) are critical for marine life, and therefore influence the functioning of ocean ecosystems and the global carbon cycle. Indeed, they are micronutrients essential to life (e.g. Fe, Zn, Cu), tracers of modern processes in the ocean (e.g. Al, Mn), and are significantly perturbed by human activities (e.g. Pb). In the Western Mediterranean, the biogeochemical cycling of TEs is largely governed by atmospheric inputs (Migon et al., 2002). However, the influence of rivers on TE distributions is discernible in the sediments of the adjacent continental shelf (Ferrand et al., 1999; Roussiez et al., 2005 and 2006; Radakovich et al., 2008). According to the Mermex Group review (The Mermex Group, 2011), present and past inputs of riverine particulate TEs to the Mediterranean, especially anthropogenic TEs, need to be more precisely estimated.

Estuary and delta sediments are suitable environments to reconstruct historical accumulation of elements and anthropogenic chemicals from continental and marine sources during last centuries or even millennia (e.g., Valette-Silver, 1993; Santschi et al., 2001; Callender, 2003). In particular, this is the case for the Rhône prodeltaic prism, which receives particulate inputs from the Rhône River (RR), and has archived its inputs for centuries (Fanget et al., 2013). Studying this area provides a unique opportunity to document the chronology and the sources of the TE inputs from the largest riverine sediment source to the Mediterranean Sea.

Here, we document the temporal accumulation of TEs (Ag, Cd, Co, Cr, Cu, Ni, Pb and its stable isotopes, and Zn) in sediment recovered in the Roustan lobe within the Rhône prodelta area, located on the continental shelf of the North side of the Western Mediterranean (Fig. 1). Based on a multiproxy study, including major elements and radiometric analyses, we have been able to identify the major changes in TE accumulation in the Rhône prodelta sediments during the last four centuries. These changes are discussed in terms of hydrological regime, channelization of the RR mouth, climate change, and TE contamination history.

2. Study area

The RR and its tributaries drain various catchments including sedimentary rocks, granitic, and metamorphic rocks from the Alps, Massif Central, and Vosges. In addition, RR carries alluvium

from one of the most industrialized and urbanized European catchments (The Mermex Group, 2011). The RR sediment load has varied significantly during the past few centuries. Since the XIXth Century, it has decreased by a factor close to ten, mainly because of stabilization of mountain slopes due to decline of rural agriculture and reforestation, and to a lesser extent because of damming and dredging activities (Pont et al., 2002; Maillet et al., 2006a; Provansal et al., 2014). However, since the completion of damming of the Nile in 1970, the RR is the largest source of continental sediment for the Mediterranean Sea. The annual solid discharge during the last decades was highly variable, from 2 to $20 \cdot 10^{12} \text{ g yr}^{-1}$ (Mtons yr^{-1}) (Pont et al., 2002; Antonelli et al., 2004; Sabatier et al., 2006; Launay, 2014), with an average flux of $3.2 \cdot 10^{12} \text{ g yr}^{-1}$ for the last 30 years (Gairoard et al., 2012). Sediments delivered by the RR are entrapped in the prodelta, with sediment accumulation rates varying from 50 cm yr^{-1} at the mouth to 0.2 cm yr^{-1} at 20 km seaward (Calmet and Fernandez, 1990; Charmasson et al., 1998; Radakovitch et al., 1999; Miralles et al., 2005). Eighty percent of particulate load of the RR is delivered during flood events (Pont et al., 2002), creating stratified structure in the prodelta sediments (Fanget et al., 2013) which is superimposed over longer-term trends of accumulating sediments.

According to the review of the centennial hydrological changes by Fanget et al. (2013), the chronology of floods is quite well documented from 1500 to 1995 AD. First, a mean of 4 to 5 major floods occurred per decade between 1500 and 1650 AD. Second, from the XVIIth Century climatic minimum to the XIXth Century, corresponding to the Little Ice Age (LIA), the frequency of floods increased, with 10–20 major floods in 1701–1710, 1751–1760, 1771–1780 and 1811–1820 AD. The flood frequency has decreased since the end of LIA (Pichard, 1995), from 8–9 per decade at the ends of the XIXth Century, to 2–3 per decade at the end of the XXth Century. Since that time, the 2003 AD flood and its associated seabed morphological changes have been extensively studied (Eyrolle et al., 2006; Maillet et al., 2006b; Miralles et al., 2006b; Antonelli et al., 2008). The geomorphology of the RR mouth has been modified several times as a result of natural stresses and/or human-induced channelizations. After the major floods between 1709 and 1711 AD, the Bras de Fer lobe formed during the LIA (Arnaud-Fassetta, 1998) was displaced eastward (Fig. 1). Between 1711 and 1852 AD, the RR displayed three mouths (Piémanson, Roustan and Pégoulier mouths, Fig. 1). In 1855 AD, in order to improve fluvial navigation, the Piémanson and Roustan mouths were closed, and in 1892 AD the Roustan mouth was re-opened for navigation and the Pégoulier channel was progressively filled. Since 1892 AD, the Roustan mouth is the only exit of

the Grand Rhône (Fig. 1), the foremost distributary channel of the RR discharging into the Mediterranean.

3. Material and Methods

3.1. Core collection

This study is based on a multi-proxy analysis of a 7.71 m-long piston core (KS57, Fig. 1) collected on the Roustan prodeltaic lobe (43°17.10'N; 4°50.97'E) at 79 m water depth during the RHOSOS cruise in September 2008 aboard R/V Le Suroît. The core was sliced every 2 cm from 2 cm below the sediment-water interface (SWI) to 100 cm, then every 4 cm down to the bottom of the core, which was 767 cm below the SWI. Subsamples were frozen (−18°C), freeze-dried and stored under cold (4°C) and dark conditions until analysis.

3.2. Chemical analyses

Iron (Fe) and Al concentrations were determined by atomic absorption spectrophotometry (AAS, SpectrAA 600[®], Varian). Aluminium (Al), Co, Cr, Cu, Li, Ni, Zn, and Pb (including its stable isotopes) analyses were performed after total dissolution of the sediment with a mixture of HCl, HNO₃, and HF in hermetically sealed Teflon bombs according to the protocol described by Loring and Rantala (1990) and modified by Chiffoleau et al. (2004). For Ag and Cd, in order to avoid interferences with ICP-MS, the sediment was digested using only HCl and HNO₃. All reagents used were SupraPur[®] (Merck). Concentrations were determined using inductively coupled plasma mass spectrometry (ICP-MS, Element X Series[®], Thermo Electron Corporation). The results were validated using certified reference materials (CRM), MESS-3 and BCSS-1 from the National Research Council of Canada. A blank sample and CRM were included with each batch of 15 samples in the total digestion procedure and then analyzed with ICP-MS and AAS. The blank values were always below the detection limits. Values obtained for major element analysis were always within the range of certified values (Table S1).

Organic matter (OM) is estimated using organic carbon (C_{org}) content. Total carbon (C_t) and organic carbon (C_{org}) were determined using freeze-dried and homogenized subsamples of sediments with an elemental analyzer (Model CN 2000[®], LECO) after acidification with 2M HCl (overnight, at 50 °C) for C_{org} in order to remove carbonates (Cauwet et al., 1990). The precision for

C_{org} and C_t analyses was $\pm 2\%$. Concentrations are expressed as the weight percent of dry sediment (% dry weight). Calcium carbonate content was calculated from mineral carbon ($C_t - C_{\text{org}}$) using the molecular mass ratio $\text{CaCO}_3:\text{C} = 100:12$.

3.3. Numerical treatment, statistical analyses, and normalization

We applied Principle Component Analysis (PCA), a multivariate technique in which the numbers of variables is reduced to a smaller number of factors that describe the principal variability or joint behavior of the data set. Geometrically, this new set of variables represents a principal axis rotation of the original coordinate axes of the variables around their mean (Jackson, 1991; Huang et al., 2010). The statistical computations were performed with XLSTAT® software from Addinsoft.

Spectral analysis is performed using the Lomb-Scargle Periodogram, a generalization of the Fast Fourier Transform to data sets with non-uniform sampling period (<http://www.peranso.com/>). The false alarm probability (p-value) for every point in the periodogram is estimated in order to determine the periods of significant cycles in the data. Peak frequencies are found using a first-order difference scheme that identifies local maxima in the periodograms, and errors in the location of peaks are estimated as the distance halfway between a local maximum and the nearest local minimum in the periodogram, or one-quarter of the natural line-width of frequency peaks, whichever is larger. In an ideal case, these two estimates are approximately equal.

In order to test if the observed variations in the profiles were the result of mineral changes within the cores, the trace element concentrations were geochemically normalized by the use of tracer elements (Kersten and Smedes, 2002). Aluminum was chosen as the granulometric normalizing element to take into account the clay fraction with its high specific surface, which favors metal binding for Co, Cr, Cu, Ni, Pb, V and Zn (e.g., Windom et al., 1989).

4. Results and discussion

4.1. Characterization of sediment core

The dating of core has been established based on ^{210}Pb , ^{137}Cs vertical profiles and microfossil assemblages (ostracods, benthic foraminifera, and coccoliths) in combination with sedimentological

proxies (Fanget et al., 2013). Herein, we present our geochemical results based on this well-constrained chronology. This 7.67 m core covers a period of just over 400 years. The first 3.75 m below the SWI has accumulated since the middle of the XIXth Century, corresponding roughly to the beginning of the industrial era in Europe. Figure S1 gives the relationship between depth and date of the deposits in core.

During the modern history of the subaqueous Rhône delta, both fluvial and marine inputs have occurred. Fanget et al. (2013) proposed that the structure of core KS57 can be summarized by four different periods (Intervals I–IV), spanning a time range between ~1600 AD and the sampling date (i.e., 2008 AD), corresponding to different positions of distributary-channels induced by climate events and/or by human activities. Low RR influence is observed between 1600–1711 AD (Interval I) and 1855–1892 AD (Interval III). These periods correspond to the time when the RR was flowing into the Mediterranean through the Bras de Fer channel (i.e., at the westernmost side of delta system) and the Pégoulie channel (i.e., at the easternmost side of delta system) respectively (Fig. 1). Conversely, significant RR influence can be detected between 1711–1855 AD (Interval II) and 1892–Present-day (Interval IV). During Interval II, the RR had three mouths (Piémanson, Roustan, and Pégoulie mouth) and the sediments accumulated on the Grand Rhône lobe; during Interval IV, the Piémanson et Pégoulie mouths were artificially closed and the sediments accumulated on the Roustan lobe (Fig. 1). The strongest riverine influence is recorded when the RR mouth was within the modern Roustan channel, i.e., just in front of the KS57 core location.

The Al, Fe and Li profiles confirm and allow us to refine the previously described chronological patterns (Fig. 2abc). Low concentration values of these clay mineral proxies occurred around 1700, 1855, 1895, and 1995 AD, dates which match the boundaries between Intervals I, II, III, and IV, and close to the SWI respectively. These low values are characteristic of low clay contribution, which suggests the presence of sandy layers, such as the one already pointed out at ~270 cm below the SWI (~1900 AD) by Fanget et al. (2013). These structures indicate that sedimentation changes have occurred during channel modifications and flooding events. Firstly, several changes occurred between 1700 and 1710 AD, i.e. at the boundary between intervals I and II (Fig. 2). In 1702, huge floods occurred in Haute Provence, following which the Bras de Fer Branch (Fig. 1) was progressively filled with RR alluviums, and in 1706, the Launes channel (east of Bras de Fer) was created (Pichard, 1995; Provansal et al., 2003). Secondly, after 1890, the Roustan mouth was opened for navigation and the Pégoulie channel was progressively filled in.

Thirdly, and more recently, huge floods of the RR were recorded during the period from October 1993 to February 1994 (e.g., Arnaud-Fassetta, 2003). During these three periods, the concentrations of Al and Fe were sharply depleted (Fig. 2ab). In addition, within Interval III, we observed Li-enriched sediment (Fig. 2c). In this case, Li enrichment alone (i.e., without Al or Fe) would not trace clay mineral as usually expected (Loring, 1991), but may indicate an increasing influence of marine material. This hypothesis is supported by the presence of marine ostracod assemblages within Interval III (see *figure 7* in Fanget et al., 2013). This connection between the abundance of marine ostracods and the Li-specific enrichment is also observed in the first part of Interval I, corresponding to the period 1600 and 1670 AD; indeed, abrupt changes can be seen in the Li/Al ratio profile (Fig. S2).

Some insights into the early diagenetic processes occurring in the core, which may contribute to the conditions of TE accumulation on the Roustan lobe, can be deduced from the OM vertical distribution in the core. Indeed, C_{org} concentrations decreased with depth in the first ~270 cm below SWI, i.e., within Interval IV (Fig. 3). Below, C_{org} remained quite unchanged down to the bottom of the core. The redox potential discontinuity was at 2.7 cm below the SWI for a sediment core taken in May 2008 at the same location (79 m) on the Roustan lobe with a multitube-corer, which preserved the SWI. This result is consistent with the oxygen penetration depth (OPD) of 5 mm measured *in situ* by microelectrodes in this sediment (Cathalot et al., 2010). This result is consistent with the extensive microelectrode measurements in Roustan lobe, which indicate that oxygen rarely penetrates more than 5 cm in the sediment (Cathalot et al., 2010). Thus, we can conclude that an important fraction of the OM mineralized under anaerobic conditions, as suggested by Garcia-Garcia et al. (2006). In the Interval IV, i.e., between 0 and 270 cm, the carbonate concentration (Fig. S3) is consistent with the results of Rassmann et al. (2016), with a rapid decrease just below the SWI (corresponding to aerobic oxidation) followed by an steady increase (corresponding to anaerobic degradation). Below ~270 cm, carbonate content steadily decreased from 45 % to 35 % with depth (Fig. S3). This feature of carbonate (*in situ*) dissolution suggests the persistence of OM degradation (e.g., Emerson and Bender, 1981). Both autochthonous biogenic and continental detrital carbonates (from calcareous outcrops in the RR basin) can contribute to the carbonate content of the sediments of the prodelta region, and it is known that the carbonate dissolution rate depends upon its mineral origin: aragonite, which constitutes most of carbonate deposited in shallow environments, is more rapidly dissolved than calcite (Mucci, 1983). It is interesting to note that

according to Fanget et al. (2013), core KS57 mainly consists, from its bottom to ~270 cm (Intervals I and III), of homogeneous silty clay with abundant iron sulfide, and between ~270 cm and the SWI (Interval IV) of laminated grayish and brownish silty clay. In brief, our data suggest that the degradation of OM is active in Interval IV, but weak below. As expected, the OM degradation rate is diminished in sulfidic sediments. However, the diagenetic context is more complex, since the Mn/Li profile show peaks at 47, 270 and 610 cm (Fig. S4). These peaks suggest the presence of relicts of oxic SWI lying at these depths, which would correspond to drastic changes in the sedimentary conditions. Indeed, a Mn-enriched layer points to Mn^{IV} precipitation as Mn oxihydroxides, when dissolved Mn^{II} diffusing from below encounters molecular O_2 diffusing from the underlying water column (e.g., Burdige and Gieskes, 1983). These oxihydroxides are known for their efficient adsorptive properties for TEs (e.g., Gadde and Laitinen, 1974). The main peak, at ~270 cm, corresponds to the reopening of the Roustan channel at the end of the XIXth Century; the peaks at 47 and at 610 cm correspond to the huge flooding periods, 1993-1994, and at the very end of the XVIIth Century respectively (Fig. S4). The observation of multiple Mn peaks in sediment is not uncommon, and has been interpreted as the result of non-steady-state diagenesis due to changes in the carbon cycle (e.g., Deflandre et al., 2002; Kuzyk et al., 2011) or in turbidite sequences (e.g., Colley et al., 1984).

4.2. Trace elements behavior before industrialized era

Table 1 summarizes statistics on TE concentrations measured in sliced layers of core. We have arbitrary defined, as usually admitted, the European industrial era as the Second Industrial Revolution, i.e., a period starting 1850 AD. We checked this choice using $^{206/207}\text{Pb}$ and $^{206/204}\text{Pb}$ isotopic ratios to qualify metal anthropization. $^{206/204}\text{Pb}$ were higher than 18.67 before 1850 AD, ranging between 18.85 and 18.67, whereas after this date, values were lower than 18.67, down to 18.33 (Fig. 4). On the other hand, in spite of a slight steady decrease of the $^{206/207}\text{Pb}$ isotopic ratios from ~1.20 to ~1.93 corresponding to the period between 1600 and 1850 AD (especially during the XVIIth Century), the beginning of the huge decrease in the ratios (down to 1.175) also occurred around 1850 AD. Thus, TE mean concentrations during the pre-industrialized era were in most cases low, but for Ag ($0.10 \pm 0.02 \mu\text{g g}^{-1}$), Cd ($0.17 \pm 0.02 \mu\text{g g}^{-1}$) and especially Cr ($70 \pm 4 \mu\text{g g}^{-1}$), higher than the average composition of Earth's upper crust (EUC, Table 1). The mean concentration of these TEs are in the range of pre-industrial concentrations found in the deepest part of the

sediment cores collected in the Cap de Creus canyon along the Northwestern Mediterranean margin (Cossa et al., 2014). Relatively high Cr concentrations ($\sim 70 \mu\text{g g}^{-1}$) were also measured during Rhône river flooding events (Radakovitch et al., 2008), and could be the result of naturally enriched minerals. Indeed, relatively high Cr contents are found in pristine suspended load of the Po River originating from the Western Alps as a result of the erosion of ultramafic rocks (ophiolite) (Amorosi, 2012). Ag and Cd pre-industrial concentrations are not yet documented for the RR basin, and more research is needed in that respect. TE concentrations were quite correlated with each other in the pre-industrial period, suggesting, at a first approximation, common origins ($p < 0.01$, Table 2). Ag is an exception since it is only significantly correlated ($p < 0.01$) with Cd. However, based on the results of a PCA, this allows sorting out three groups of TEs based on their common variations (Table 3a, Fig. S5a). One group combines Li, Cr and Ni, another combines Co, Cu, Pb and Zn, and the third combines Ag, Cd and carbonates. Al and Mn are both associated with the first and the second group, and the second and the third, respectively (Fig. S5). It can be reasonably speculated that the first group indicates ultramafic rocks origin, the second other clay minerals, and the third a marine biogenic component.

4.3. Temporal variations before industrialized era

Temporal variations of TE/Al, during the 1600-1850 AD period, are illustrated in figure 5. In this graphical representation, in order to remove the effect of sediment texture on TE concentrations, we have normalized TE concentrations with Al concentration. Several peaks of TE/Al ratios can be identified in relation with particular events documented on Table 4. Event 1, with small peaks of all TE/Al ratios, coincides with the very cold and long 1620-21 winter. Events 2 and 4 are related to the changes in RR channelization that occurred between Intervals I-II, and II-III (Fig. 5), respectively. In addition, Event 2 is coincidental with a Mn-rich layer (Fig. S4), that indicates an episode of abrupt sedimentation. Event 3 occurs during major inundations that have been recorded in the 1774-1777 period (Table 4).

4.4. Trace elements and climatic variations

Spectral analysis carried out on data in the pre-industrial case identifies very few significant periodicities on a per-element basis. Note that this does not mean such cycles are not present, but rather reflects the fact that insufficient data are available to confidently distinguish them from noise.

However, similarities in the spectra of elements grouped as discussed above support the grouping of elements by common origin, and are suggestive that significant peaks may be identified if additional data could be collected. In the ultramafic rocks group, for example, peaks corresponding to 65 ± 10 year, 21 ± 1 year, and 14 ± 0.3 year cycles line up in all three spectra (Fig. S6). The coincidence of sub-significant peaks in the clay minerals group also suggests the possibility of 65 ± 10 and 14 ± 3 year cycles (Fig. S7). Among the marine biogenic group, only the 65 ± 10 year cycle is coincident in all of them (Fig. S8).

Interestingly it appears that most of the frequencies here identified (i.e., 14-17, 21, 65-year cycles) correspond to the yearly mean sunspot numbers identified by Lomb (2013), including the classical decadal frequency and the Gleissberg cycles (64-110 year) for the period 1700 and 2011 AD. Also, interesting to note is that the Atlantic Multidecadal Oscillation, which govern westerly circulation in the Atlantic and influences the European climate, presents a periodicity ranging from 65 to 90 years (Knudsen et al., 2011; Olsen et al., 2012). This suggests that the oscillations in TEs of the deposited sediment on the Roustan lobe are not due to chance but reflect the climatic variations that govern drought and flood, and consequently the erosion of the catchment basin of the Rhône River. Such a process has recently been suggested to occur in the Parana delta (Milana and Kröhling, 2015), whereas Park and Chang (2013) have observed coincidence between sunspots and precipitation rhythms. Longer time series are required for verifying this hypothesis.

4.5. Trace elements anthropization

Data clearly show that several TE concentrations in the sediment core layer, deposited after 1850 AD, were significantly greater than those calculated for pre-industrial era. These elements show increases in mean concentrations of 3.5 times for Ag, ~2 times for Cd and Pb, and ~1.3 for Cu and Zn, higher than the pre-industrial values. Conversely, Co, Cr, and Ni mean concentrations remain roughly unchanged (Table 1). A PCA, performed on post-1850 AD data, allows separation of several groups of TEs. Pristine TEs, such as Co, Cr, and Ni, are linked with clay markers (Al, Li and Fe), Cr and Ni being associated with Mn, whereas Cr is correlated with Fe. Another group consists of “anthropogenically impacted” elements (Ag, Cd, Cu, Pb, and Zn), within which Ag is correlated with C_{org} (Table 3b, Fig. S5b). Stable isotopic compositions of Pb shows highly significant ($p < 0.01$) differences (1.196 ± 0.003 vs 1.1843 ± 0.004 for 206/207 ratios (Table 1). These results clearly indicate that recent sediments deposited in the Roustan lobe are contaminated

with Ag, Cd, Cu, Pb and Zn, and have been for at least 150 years. This conclusion, which is consistent with the RR as the main source of TE contamination in the Gulf of Lions sediments (e.g., Roussiez et al., 2005, 2006; Radakovitch et al., 2008; Cossa et al., 2014), can reasonably be extended to the whole of the Rhône pro-delta sediments.

4.5. Chronology of TE accumulation during the industrialized era

Nine peak-events (Table 4) were recorded in the TE/Al chronological profiles (Fig. 6). Event 5, occurring during Interval III, is enriched with TEs having an anthropogenic component (Ag, Pb and Zn), and is coincidental with the 1889-91 AD period of high flooding. Event 6 is a typical Mn-enriched peak, associated with Co, Ni, and Cd, relict of the SWI exiting before the re-opening of the Roustan channel, which brought new river sediments to the KS57 site on the prodelta lobe. Also noteworthy is the Co and Ni association with Mn, which is a common feature present at the SWIs (Goldberg, 1954; März et al., 2011; Friedrich and Catalano, 2012). Events 7-8 correspond to significant inputs of anthropic TEs, which may be related to the December 1926, November 1944 AD, February 1945 AD and November 1963 AD floods (Table 4). Trace element contamination increased dramatically after the IInd World War and peaked around 1977 (Fig. 6). The occurrence of TE concentration maxima between 1960 and 1980 AD is a common chronological feature in Europe, in North America and, even further, in the contaminated aquatic sediments of the entire Northern hemisphere (in review by Callender, 2003). Several authors suggested that coal burning is the reason for this temporal pattern (e.g., Muller et al., 1977 cited by Callender, 2003). In areas neighboring the Rhône basin, coal burning and smelters influence has been shown for Pb in the Geneva Lake sediments (Monna et al., 1998), and in some Gulf of Lions canyons and Pierre-Blanche lagoon (Miralles et al., 2006a; Elbaz-Poulichet et al., 2011; Cossa et al., 2014). Similar chronology is also observed for Pb, Cd and Zn in Arctic lake sediment (Hermanson, 1991). In North America, temporal distributions of sedimentary Pb in sediment cores from urban reservoirs and lakes also peak between 1960 and 1980 AD (in review by Callender, 2003). The reconstructed anthropogenic Pb emissions in France also peaked between 1960 and 1980 (Ferrand et al., 1999; Miralles et al., 2006a). Compiling the TE emissions from the pre-industrialized era to 2010 AD by gathering information from various sources (Nriagu, 1979, Nriagu and Pacyna, 1988, CITEPA, 2014) we have been able to reconstruct the TE atmospheric emission changes in Europe since 1875 (Fig. 7). The temporal trends obtain for Cd, Cu, Pb and Zn are very consistent with the TE/Al ratio

chronologies recorded here (Fig. 6). This result suggests that the anthropogenic TEs accumulated in the Rhône prodelta sediments are mainly from atmospheric deposition on the RR catchment area. Copper contamination in France originates mainly from copper sulfate, used as fungicides on vineyards, known as “Bordeaux mixture”. This “Bordeaux mixture” is progressively replaced by Ag bullet for treating vines. Copper contamination also originates from train catenaries and brake pad erosion (CITEPA, 2014). Chromium results mainly from electrical steelworks, from foundries, and from glass production. Cadmium is a natural back product of phosphate fertilizers (Otero et al., 2005). For the most anthropogenically impacted TEs (Pb and Zn), it is generally thought that the origin of the anthropogenic increase of Pb was emissions by leaded gasoline burning and regional industry, and that the origin of the Zn was a combination of urban runoff and coal burning (e.g., Cossa et al., 2014).

4.6. Lead sources

Stable isotopic ratios of Pb allow examination of the origins of Pb. Figure 8 illustrates the relationship between $^{206/207}\text{Pb}$ and $^{206/208}\text{Pb}$ in the core and provides comparable data to that obtained in the Cap de Creus canyon located at the western end of the Gulf of Lions (Cossa et al., 2014). Natural, industrial, and gasoline additive isotopic Pb values are also given (Ferrand et al., 1999; Miralles et al., 2006a; Komarek et al., 2008). Firstly, it appears that sediments deposited on the Roustan lobe before 1850 have isotopic ratios close to the values of natural Pb, with values of 1.200 at the bottom of the core (765 cm, i.e., ~1600 AD), whereas sediments deposited since 1850 AD, with $^{206/207}\text{Pb}$ values down to 1.175, are clearly contaminated by European gasoline Pb and industrial Pb pools (Fig. 8). Secondly, $^{206/207}\text{Pb}$ ratios in the sediments accumulated since 2000 AD (0-20 cm) are higher than in subsurface layers in the last two decades of the last century (1.184 ± 0.0017 vs 1.179 ± 0.0023). These lower values are close to those given by Ferrand et al. (1999), i.e., 1.178-1.181, for sediments collected nearby between 1983 and 1994. These observations confirm that Pb contamination of the RR particulate discharge reached a maximum at the end of the XXth Century, but has been decreasing since the beginning of the XXIth Century. Using a $1/\text{Pb}$ vs $^{206/207}\text{Pb}$ model, which allows determination of the isotopic composition of pollutant lead as the intersection with the X-axis (e.g., Ferrand et al., 1999), we obtained statistically significant ($p < 0.01$) Pb isotopic signatures (the end-member for $1/\text{Pb} = 0$) of 1.183 ± 0.002 and 1.175 ± 0.002 for the pre-industrial and industrial eras, respectively. Interestingly the intercept remains 1.175 after the banning of Pb in

gasoline in Europe, i.e., 2000 AD, suggesting another anthropogenic Pb source. Aerosols, the $^{206/207}\text{Pb}$ ratio values of which are less radiogenic (< 1.16 according to Ferrand et al., 1999), certainly contribute a low value for the industrial period. However, relatively high $^{206/208}\text{Pb}$ is associated with low $^{206/207}\text{Pb}$ (Fig. 8) suggesting that industrial Pb, accumulated in soils and sediments of the RR basin has been, and still remains, a massive source of Pb for the Roustan lobe sediment.

The shape of the $^{206/207}\text{Pb}$ profile (Fig. 4) exhibits a decreasing trend in ratios between Interval I and II, separated by a bump between 1700 and 1740 AD. Based on sediment samples from Interval I only, we calculate a Pb isotopic signature (the end-member for $1/\text{Pb} = 0$) of 1.191 ± 0.002 , which is significantly lower ($p < 0.01$) than the signature obtained for the pre-1850 AD period (see above). These features suggest (i) that an input of low radiogenic Pb has accompanied the flood or/and channelization change that occurred at the end of the XVIIth Century (see Event 2 on Table 4 and Fig. 5), and (ii) that Pb contamination slowly started a hundred years before the beginning of the 2nd Industrial Revolution (1850 AD).

5. Summary and Conclusions: The Rhône prodelta anthropization in the context of Mediterranean Sea

Trace element concentrations in surface coastal sediments are largely documented in the Western Mediterranean and the Eastern Mediterranean (e.g., Scoullou and Oldfield, 1986; Abdel-Moati and El-Sammak, 1997; Abi-Ghanem et al., 2009 and 2010; Schintu et al., 2009 and 2015; Mzoughi and Chouba, 2011; Roussiez et al., 2011). Concentrations range between geological backgrounds, with the example of Sardinia coasts (Schintu et al., 2009), and very high concentrations. In the case of Pb, the increases reach up to 30-fold increase in the sediment of Elefsis Gulf, subjected to intense anthropogenic pressures of the cities of Athens (Scoullou and Oldfield, 1986), or up to 70-fold increase in the sediment of the Manzala lagoon sediments (Nile delta), directly linked with the Greater Cairo metropolitan city (Abdel-Moati and El-Sammak, 1997).

Historical reconstructions of the anthropization are less numerous. They systematically testify temporal trends for Pb and Zn starting between 1850 and 1920 in relation with the industrialization history of each area. In some cases, the metal contamination peaks between 1950 and 1970 in

conjunction with coal utilization (e.g., Azoury et al., 2013). In other cases, there is no evidence of any decline of the concentrations in recent years (e.g., Palanques et al., 1998 and 2008). In the Po delta sediments (Northern Adriatic Sea), most Pb and Zn concentration-depth/year profiles in cores showed an upward increase from 1860 and a still significant anthropogenic supply at the time of sampling (Romano et al., 2013). In Llobregat prodelta sediment, near the city of Barcelona, the sharpest increase in anthropogenic Cu, Pb and Zn corresponds to the period 1920-1960, with 1.5 to 10-fold concentration increases (Palanques et al., 2008). In the nearby Besos prodelta sediments, the sharper increases in metal concentration also occurred in the 1920s and the 1960s, but no evidence of current reduction of the anthropogenic inputs was observed (Palanques et al., 1998). In both cases, these increases in anthropization coincide with the industrial development and population evolution in Catalonia. Similar evolution was found in Ligurian canyons sediment located near the French-Italian boundary (Martin et al., 2009 and Heimbürger et al., 2012). They contrast with the sediments of the Rhône prodelta presented here, some Gulf of Lions canyons (Miralles et al., 2006a; Cossa et al., 2014), the Pierre Blanche lagoon (Elbaz-Pouchet et al., 2011) both located within the Gulf of Lions, and even the remote Lebanon continental shelf (Azoury et al., 2013) where the sedimentation rates are high enough to point out that anthropization has reached maxima within the second half of the XXth Century.

Here, we show that Rhône River prodeltaic prism has archived riverine and marine sedimentary inputs for the last 4 centuries. During the pre-industrialized era (i.e., before 1850 AD) TE concentrations (with the exception of Cr) were no different from the average composition of Earth's upper crust. Three origins for TEs were identified: (i) ultramafic rocks (Li, Cr, Ni), (ii) other clay minerals (Co, Cu, Pb, and Zn), and (iii) marine biogenic matter (Ag, Cd). During the pre-industrial period, several peaks in the TE/Al ratios coincided with flood events or/and changes in the channelization of the Rhône River mouth. In addition, frequency analysis using Lomb-Scargle Periodograms is suggestive that some oscillations in TEs of the deposited sediment reflect climatic variations, such as the Atlantic Multidecadal Oscillation, that ultimately govern water precipitations and consequently the erosion of the catchment basin of the Rhône River.

Sediment deposited after 1850 AD has some TE concentrations exceeding up to 3.5 times those of the pre-industrial era (Ag>Cd-Pb>Cu-Zn). Principal Component Analyses allows identification of two groups of TEs during the industrialized era: (i) Pristine TEs, such as Co, Cr, and Ni, are linked with clay markers (Al, Li and Fe), and (ii) "Anthropogenically impacted" TEs (Ag, Cd, Cu,

Pb, and Zn). TE contamination increased dramatically after the Second World War and reached a maximum between 1970 and 1980, consistently with atmospheric emissions changes.

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Figure captions

Fig. 1. Detailed morphology of the Rhône subaqueous delta and main onshore morpho-sedimentary features from Fanget et al. (2013).

Fig. 2. Iron (Fe), Aluminium (Al) and lithium (Li) vertical profiles in sediment core KS57 during the last 400 years.

Fig. 3. Organic carbon (C_{org}) vertical profiles in sediment core KS57 during the last 400 years.

Fig. 4. Lead (Pb) stable isotope ratios vertical profiles in sediment core KS57 during the last 400 years.

Fig. 5. Trace element (TEs) vertical profiles in sediment core KS57 during the last 400 years.

Fig. 6. Trace element (TEs) vertical profiles in sediment core KS57 during the industrialized era.

Fig. 7. Trace element (TEs) emission increase in Europe relatively to the year 1875 AD. Diagram built from Nriagu (1979), Nriagu and Pacyna (1988) and CITEPA (<http://www.citepa.org/>).

Fig. 8. $^{206/207}\text{Pb}$ vs $^{206/208}\text{Pb}$ relationships in sediment core KS57 before and after 1850 AD. Equivalent data from the Cap de Creus (CdC) sediments (from Cossa et al., 2014) are given for comparison.

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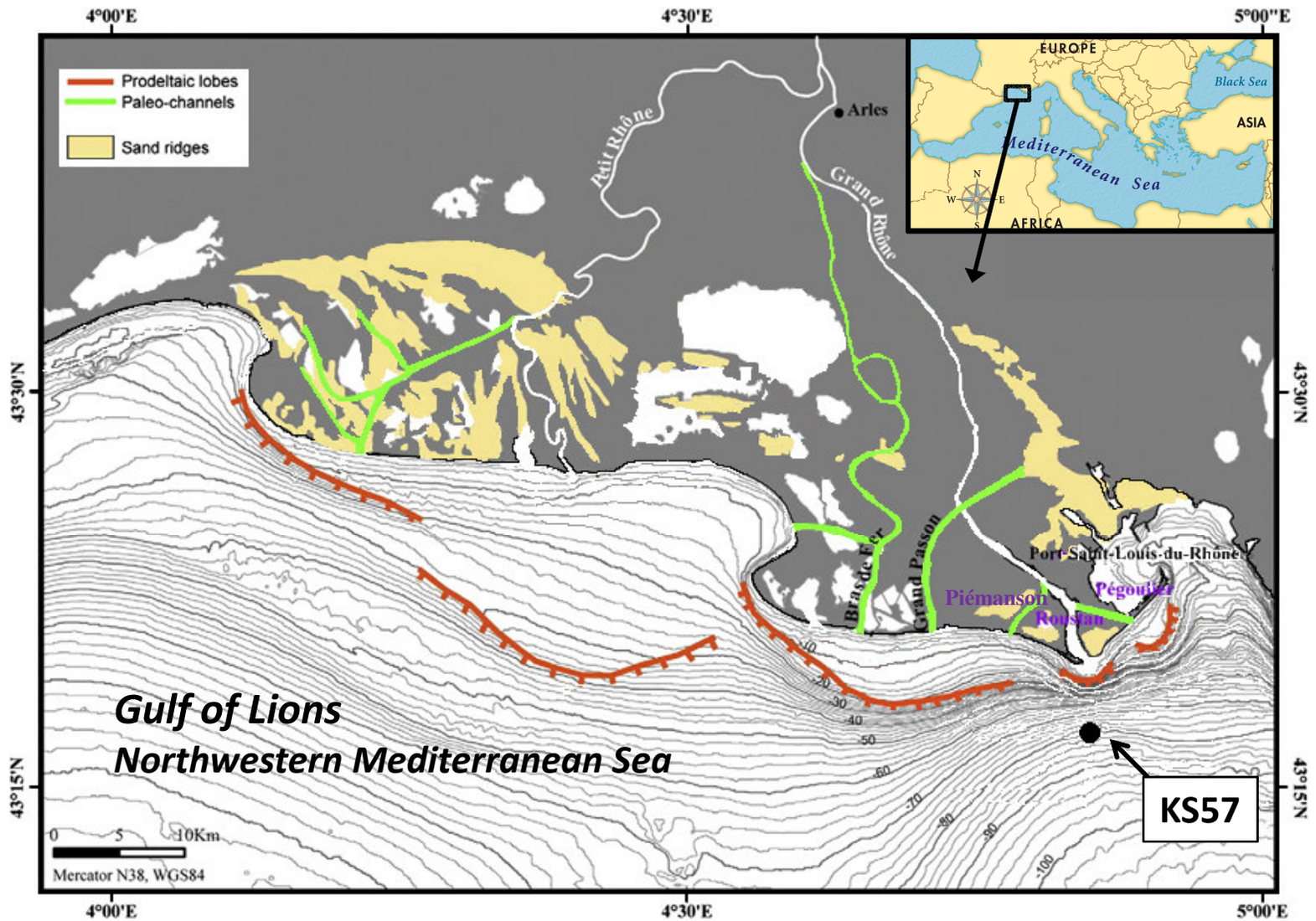


Fig. 1

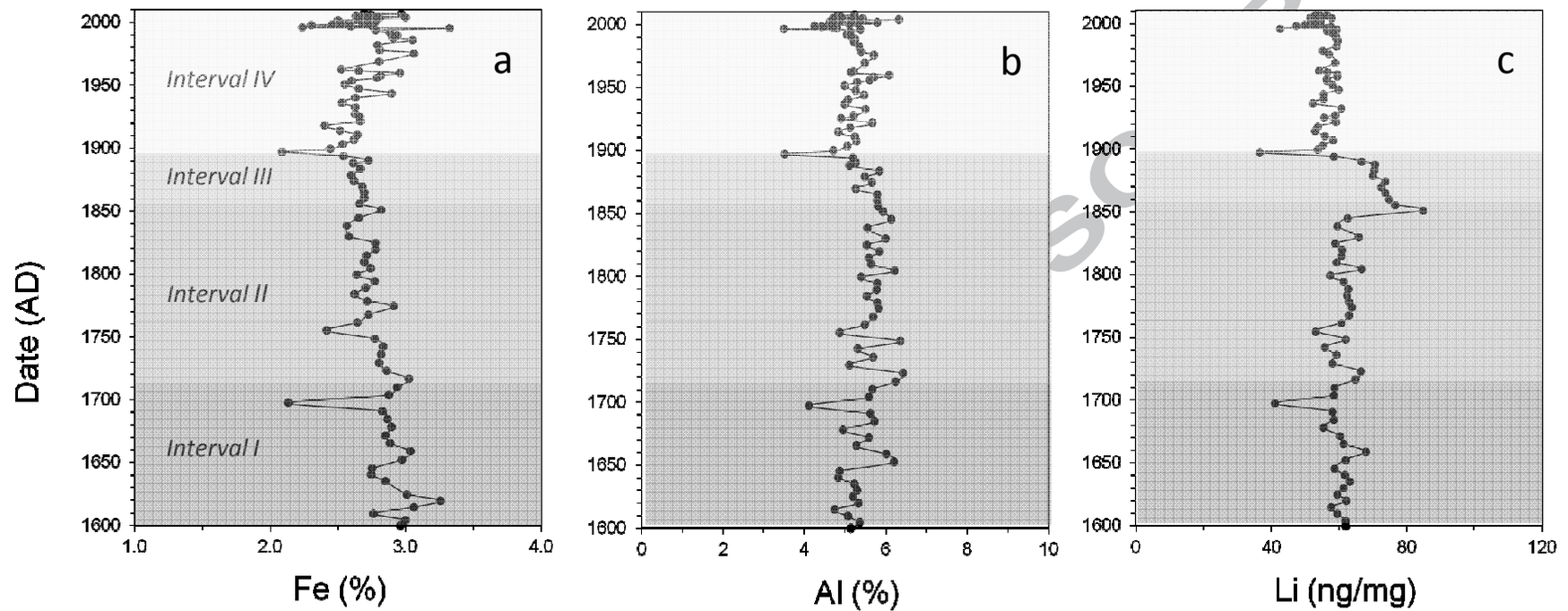


Fig. 2

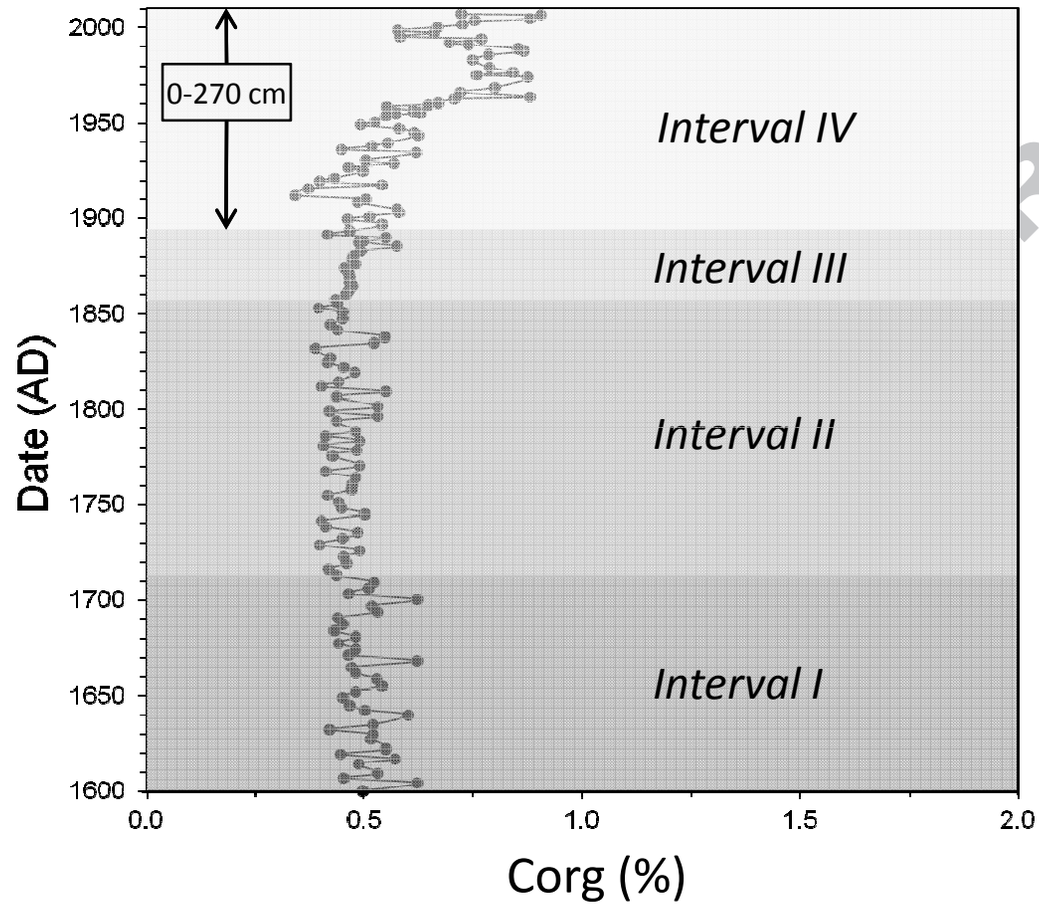


Fig. 3

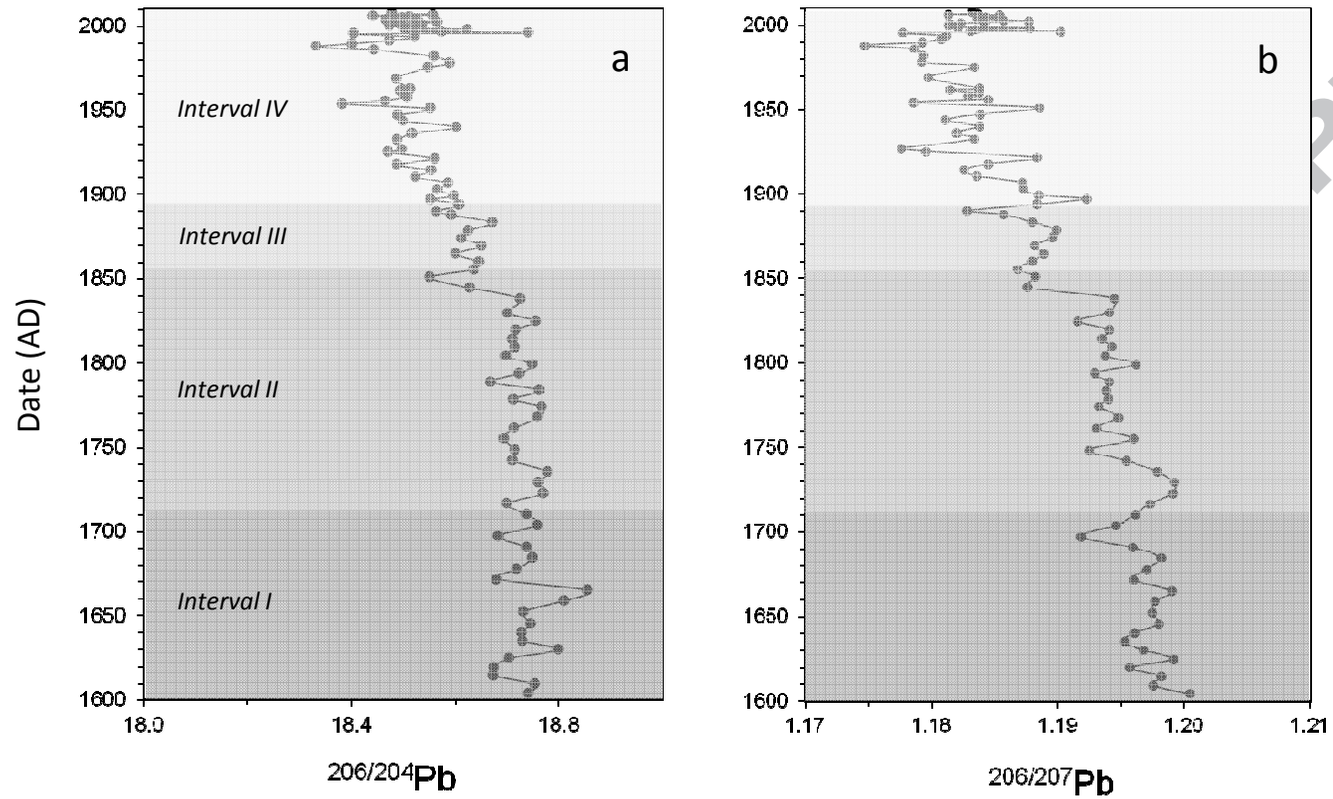


Fig. 4

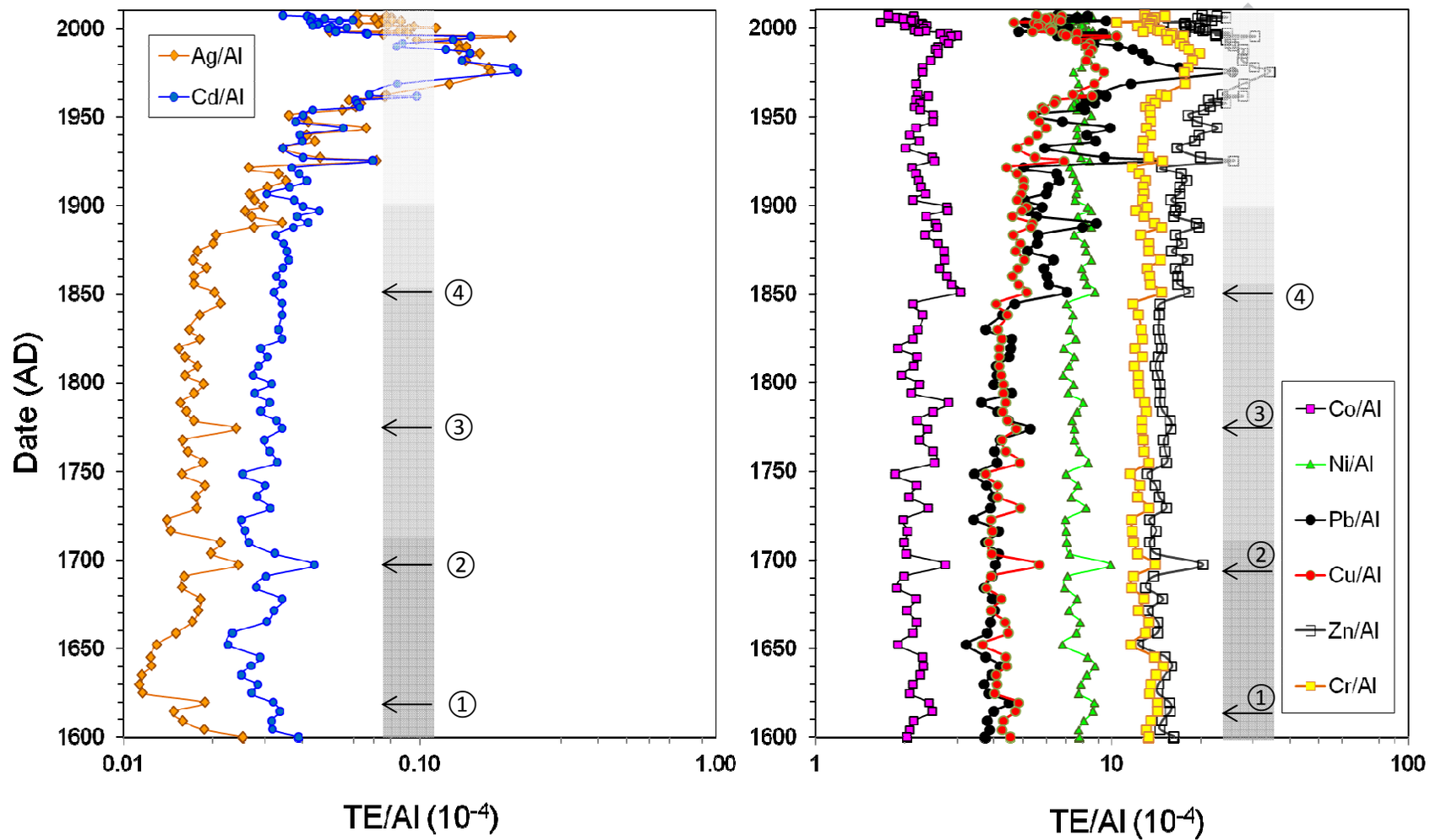


Fig. 5

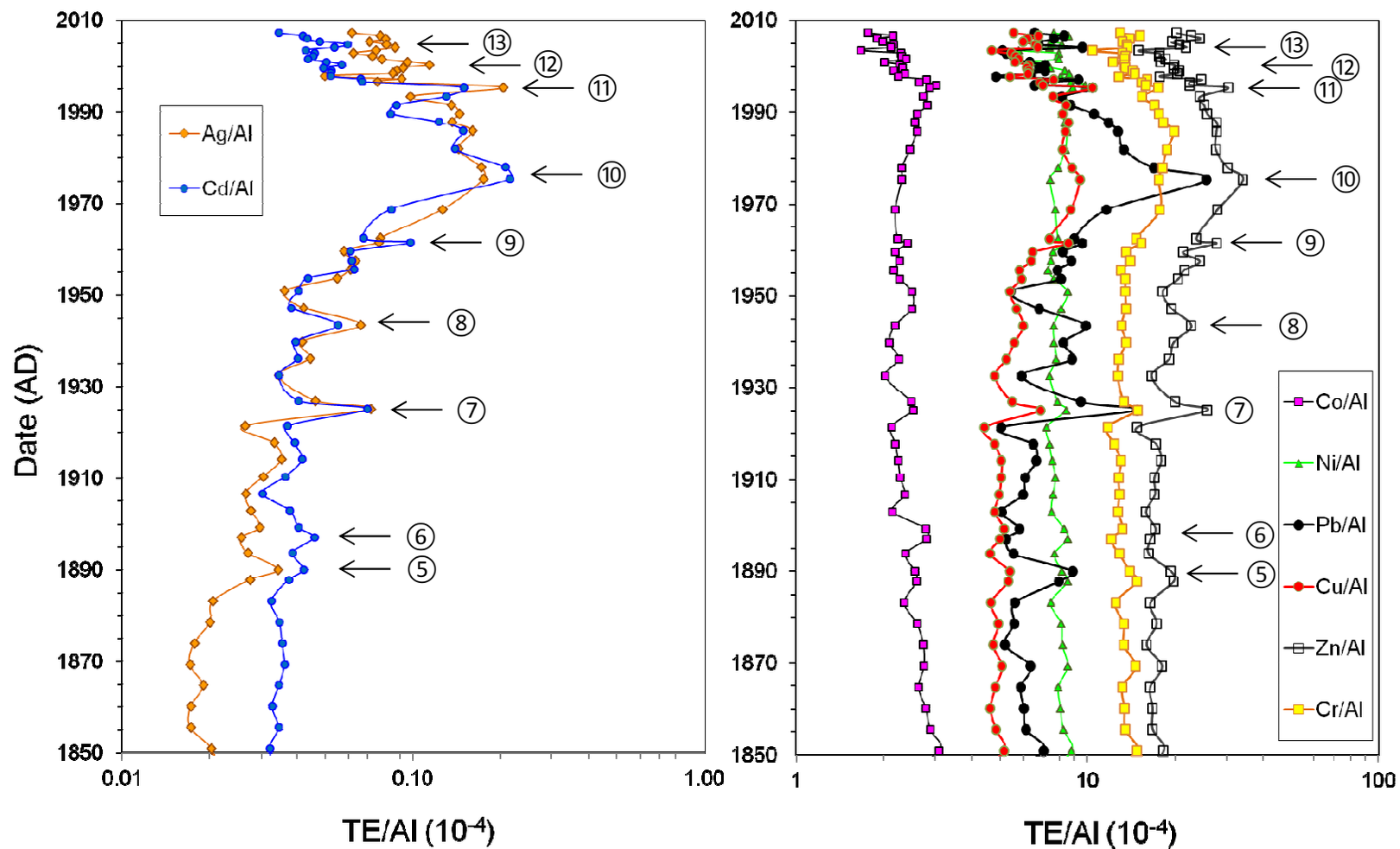


Fig. 6

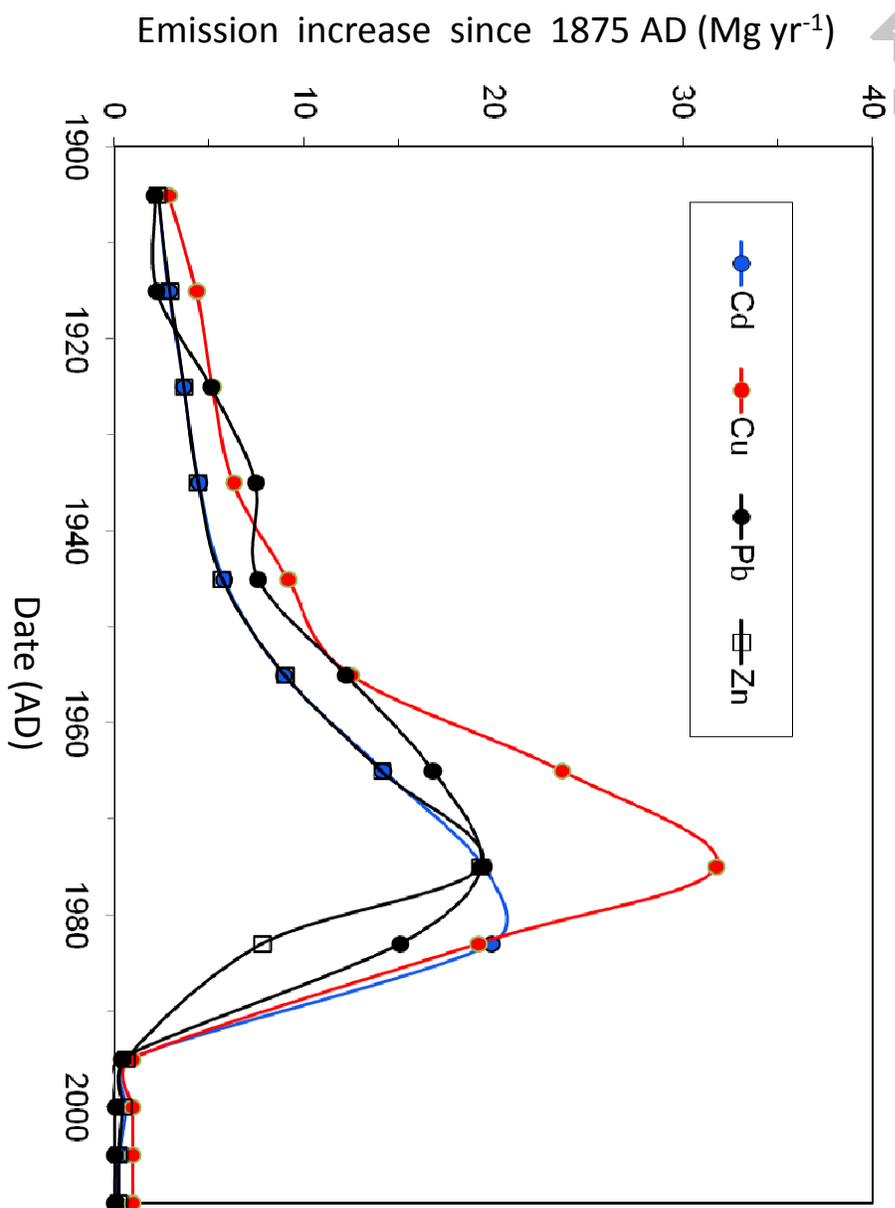


Fig. 7

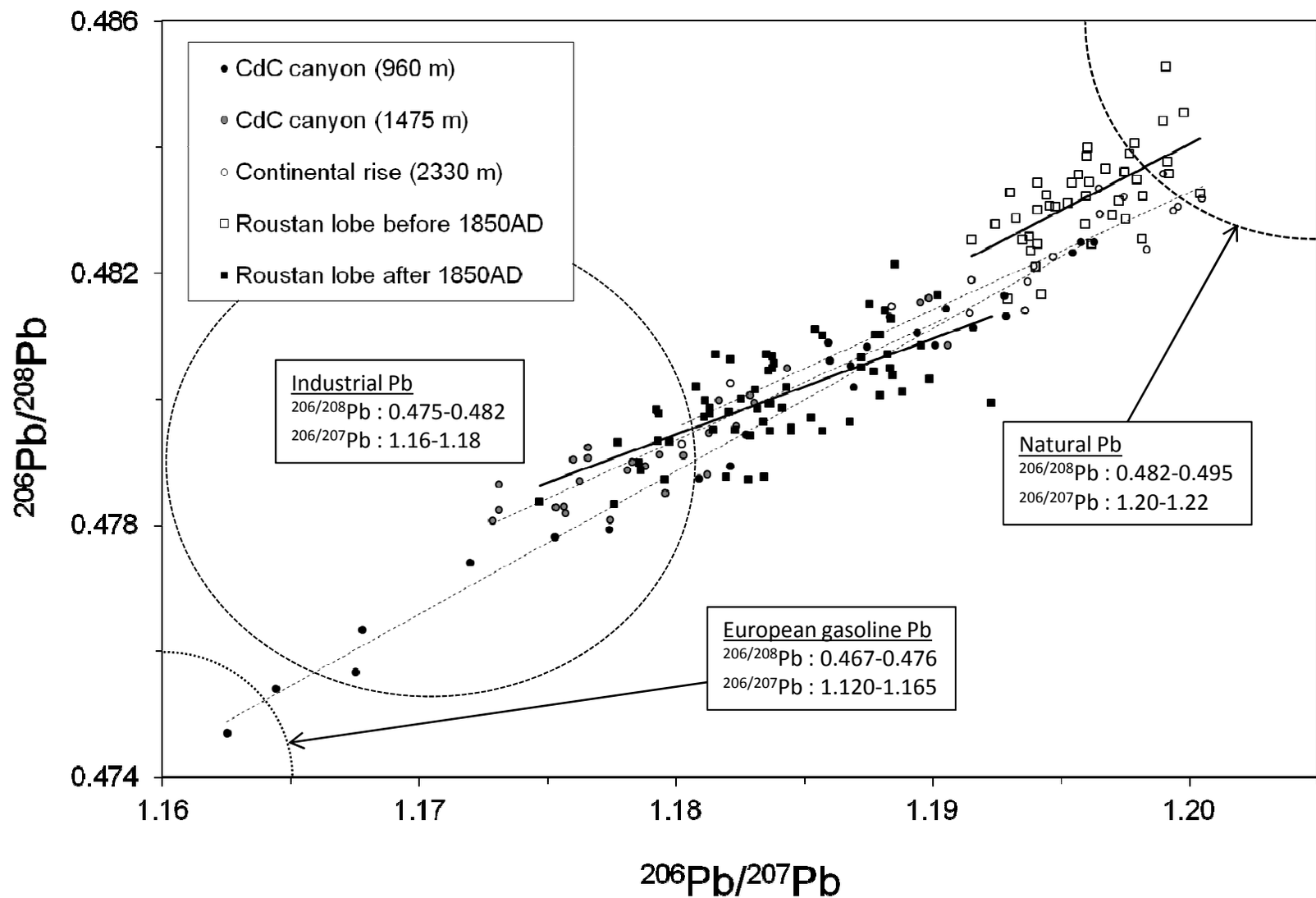


Fig. 8

Tables

Table 1. Summary statistics on trace element concentrations and Pb stable isotopic ratios in the KS57 core. Mean_a: arithmetic mean; Mean_g: Geometric mean (used for log-Normal frequency distributions); SD: Standard deviation; N: number of sediment layers analyzed. EUC refers to average composition of Earth's upper crust (values compiled by Li, 2000).

Table 2. Pearson correlation coefficients between element concentrations for the pre-industrial period. Bolt values refer to the significance of the correlation ($p < 0.01$).

Table 3. Principal component analyses. Correlation coefficients between variables and the first components (F1 to F3), (a) before and (b) after 1850 AD. In brackets, the percentages of the total variability explained by each component. Bolt values refer to the significance of the correlation ($p < 0.01$).

Table 4. Presence of peaks in the TE/Al ratio distribution profiles. Peak numbers are those given on Figures 5 and 6. Number of "x" is proportional to the amplitude of the peaks. No water discharge value is available before 1850. (1) Association Nazairienne de Généalogie. Epidémies et famines en France : Climats et épidémies. <http://angeneasn.free.fr/epidemies.htm>; (2) Fournet (21012) ; (3) Champion (1862) ; (4) Compagnie Générale du Rhône. <http://www.ecologie-et-progres.com/crue.htm>; (5) Pardé (1928) ; (6) Pardé (1948).

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Table 1. Summary statistics on trace element concentrations and Pb stable isotopic ratios in the KS57 core. Mean_a: arithmetic mean; Mean_g: Geometric mean; SD: Standard deviation; N: number of sediment layers analyzed. EUC refers to average composition of Earth's upper crust (values compiled by Li, 2000).

	Ag	Cd	Co	Cr	Cu	Ni	Pb	Zn	^{206/207} Pb	^{206/208} Pb	^{206/204} Pb
	μg/g	-	-	-							
Mean _a (1850-2008)	0.35	0.32	12.34	73.47	31.96	41.09	41.71	107.50	1.1839	0.4800	18.5306
SD (1850-2008)	0.24	0.22	1.83	11.72	7.83	3.68	19.14	25.05	0.0036	0.0008	0.0722
N (1850-2008)	61	61	62	62	62	62	62	62	62	62	62
Mean _a (1600-1850)	0.10	0.17	12.09	70.23	23.54	41.82	22.31	80.57	1.1955	0.4831	18.7276
SD (1600-1850)	0.02	0.02	1.09	3.83	1.68	1.84	2.91	5.25	0.0025	0.0008	0.0430
N (1600-1850)	43	43	43	43	43	43	43	43	42	42	42
Mean _g (1600-2008)	0.18	0.22	12.14	71.56	27.73	41.27	30.85	94.07	1.1886	0.4813	18.6098
Median (1600-2008)	0.14	0.20	12.00	70.70	26.40	41.35	28.85	88.20	1.1880	0.4807	18.5995
N (1600-2008)	104	104	105	105	105	105	105	105	104	104	104
EUC	0.05	0.10	10-18	35	25	20-56	17-20	71	-	-	-

Table 2. Pearson correlation coefficients between element concentrations for the pre-industrial period. Bolt values refer to the significance of the correlation ($p < 0.01$).

	C _{org}	Carbonates	Ag	Cd	Co	Cr	Cu	Li	Mn	Ni	Pb	Zn	Al
C _{org}	1												
Carbonates	-0.41	1											
Ag	-0.29	0.18	1										
Cd	-0.27	0.36	0.63	1									
Co	-0.18	0.40	0.17	0.31	1								
Cr	0.01	0.17	-0.09	-0.03	0.52	1							
Cu	-0.22	0.29	0.36	0.45	0.69	0.56	1						
Li	0.11	0.07	-0.04	0.03	0.42	0.93	0.53	1					
Mn	-0.26	0.54	0.34	0.31	0.55	0.22	0.57	0.14	1				
Ni	-0.03	0.08	0.01	0.03	0.71	0.81	0.66	0.69	0.31	1			
Pb	-0.29	0.47	0.40	0.48	0.42	0.50	0.64	0.48	0.60	0.33	1		
Zn	-0.15	0.22	0.36	0.51	0.57	0.56	0.84	0.61	0.39	0.60	0.72	1	
Al	-0.25	0.32	0.26	0.17	0.38	0.68	0.50	0.71	0.36	0.47	0.64	0.55	1
Fe	0.02	-0.33	-0.03	-0.26	-0.17	0.40	-0.06	0.37	-0.13	0.17	0.09	-0.12	0.25

Table 3. Principal Component Analyses. Correlation between variables and the first four components (F1 to F3). In brackets the percentages of the total variability explained by each component. Bolt values refer to the significance of the correlation ($p < 0.01$).

Before 1850	F1 (42%)	F2 (19%)	F3 (9%)
C _{org}	-0.283	-0.451	-0.282
Carbonates	0.458	0.498	-0.169
Ag	0.372	0.558	0.461
Cd	0.456	0.631	0.165
Co	0.751	0.056	-0.464
Cr	0.757	-0.601	-0.005
Cu	0.874	0.081	-0.137
Li	0.720	-0.603	0.079
Mn	0.623	0.380	-0.139
Ni	0.734	-0.433	-0.296
Pb	0.807	0.188	0.273
Zn	0.853	0.045	-0.041
Al	0.754	-0.192	0.321
Fe	0.064	-0.598	0.645

After 1850	F1 (49%)	F2 (28%)	F3 (9%)
C _{org}	-0.405	0.001	0.805
Carbonates	0.356	-0.667	-0.521
Ag	0.771	-0.556	0.164
Cd	0.827	-0.335	0.272
Co	0.520	0.742	0.153
Cr	0.964	0.068	0.007
Cu	0.930	-0.271	0.137
Li	0.370	0.865	-0.200
Mn	0.071	0.905	0.119
Ni	0.691	0.657	-0.011
Pb	0.829	-0.209	0.163
Zn	0.944	-0.211	0.110
Al	0.544	0.519	-0.217
Fe	0.865	-0.005	-0.117

Table 4. Presence of peaks in the TE/Al ratio distribution profiles. Peak numbers are those given on Figures 5 and 6. Number of “x” is proportional to the amplitude of the peaks. No water discharge value is available before 1850. (1) Association Nazairienne de Généalogie. Epidémies et famines en France : Climats et épidémies. <http://angeneasn.free.fr/epidemies.htm>; (2) Fournet (21012) ; (3) Champion (1862) ; (4) Compagnie Générale du Rhône. <http://www.ecologie-et-progres.com/crue.htm>; (5) Pardé (1928) ; (6) Pardé 1948).

Peak	Ag/Al	Cd/Al	Pb/Al	Zn/Al	Cu/Al	Co/Al	Cr/Al	Ni/Al	Documented flooding	RR discharge ($\text{m}^3 \text{s}^{-1}$)	Interval	Date (AD)	Remarks	Ref.
1	xx	x	x	x	x	x	x	x	?		I	~1620	Cold and long winter 1620-21	1, 2
2	xx	xx		xx	xx	xx	x	xx	Nov. 1694		I-II	~1695	Mn-enrichment	3
3	xx		xx	x	x				1774-77		II	~1775		3
4	xx	xx	xx	xx					?	1856: 11640	II-III	~1850	Milenium record	3
5	xx		xxx	xx					Janv. 1889, Sept. 1890, Oct. 1891	1889: 8780; 1891: 7800	III	~1890		4
6		x				x		x	Nov. 1896	9060	III-IV	~1896	Mn-enrichment	4
7	xxx	xxx	xxx	xxx	xxx	x	x	x	Dec. 1926	7400	IV	~1926		5
8	xx	xx	xx	xx					Nov. 1944, Fev. 1945	> 7500	IV	~1944		6
9		xx	x	x	x				Nov. 1963	> 3660	IV	~1964		4
10	xxxx	xxxx	xxxx	xxxx	xxxx				Nov. 1977	8690	IV	~1976		4
11	xxx	xxx	xx	xx	xx	x	x	x	Oct. 1993, Janv. 1994, Nov. 1996	9800, 11000, 8980	IV	~1995	Mn-enrichment	4
12	xx		xx						Sept., Nov. 2002	10500, 10200	IV	~2000		4
13		xx	xx						Dec. 2003	11500	IV	~2004		4