

ROLE OF DESERT AEROSOL IN METAL FLUXES IN THE MEDITERRANEAN AREA

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ABSTRACT (220 parole)

Aerosols and precipitations collected over Sardinia are mainly controlled by two end-members: Saharan and European. European background aerosols in Sardinia show a 3 to 25-fold decrease in EF_{crust} values for Cd, Zn, Pb, and Cr compared with coastal stations in the Western Mediterranean. Partitioning of total atmospheric deposition between the soluble and the insoluble phases shows that Al, Si, Fe and Pb originating from the atmosphere are mostly in the insoluble form. For Ca, Cd, Cr metals the dissolved fraction represents 50-90 % of the total input.

Aerosol dissolution experiments performed at particle concentrations ranging from 3 to 300 mg l^{-1} show Cd and Pb exhibiting low solubility at the high mass particle concentration. Solubility of Pb increases with the EF_{crust} values and the finest grain-size of the aerosols ($<1\mu\text{m}$), and is less affected by pH. Al and Fe solubilities appear to be mainly controlled by the particulate load. Dissolution of both metals decreases significantly from 30% (13 % for Fe) at aerosol loads lower than 5 mg l^{-1} , to less than 1% for both metals, at total aerosol concentrations greater than 50 mg l^{-1} , which are encountered during Saharan transport.

Fluxes (dry +wet) of metals at Sardinia show that $\cong 30\%$ for Cd and Pb, to more than 80 % for Fe and Al are due to Saharan contribution.

INTRODUCTION

There is increasing recognition that the atmosphere can be an effective, and in some cases dominant, transport path for pollutants entering the coastal and ocean (Duce et al., 1991(1)). Trace metals are emitted by industries activity from populated areas around the Mediterranean (MED) basin and from the north part of Europe (European background=BKG). In addition, with seasonal character, “pulses” of Saharan (SAH) dust carry large amounts of metals of natural origin to the MED and adjacent continental regions. The two sources are chemically very different, in particular with respect to their trace metal content. The metals attached to the aerosols can reach the sea surface following different pathways: (i) by dry deposition in which aerosols are delivered directly to the sea surface by air-sea route and trace metal solubility is constrained by aerosol-sea water reactions; (ii) by wet deposition in which aerosols reach the sea surface by the indirect air-rain water-sea water route, following in-cloud or below-cloud precipitation scavenging processes.

Several authors (Guerzoni et al., 1997(2); Loye-Pilot & Martin, 1996 (3); Bergametti et al., 1989(4) concluded that the two major factor controlling variations in particulate trace metal concentrations in the western MED are: (i) precipitation scavenging which affects both crustal and anthropogenic trace metals; and (ii) the inputs of SAH dusts, which affects crustal trace metals.

Global estimates of dust deposition to the MED Sea range significantly from ~25 to ~100 Tg yr⁻¹ (D’Almeida, 1986 (5); Prospero, 1996 (6)). Much of the variation results from the difficulty in obtaining good estimates of dust deposition. Despite the shortcomings in these estimates, it is shown for many metals (e.g., Al, Pb, Mn, Fe) that the global flux from the partial dissolution of eolian dust in surface waters is much greater than that carried by Rhone and Po rivers for Western MED and Adriatic Sea respectively (Guieu et al., 1997 (7); Rossini and Guerzoni, 1996 (8)). In the open seas, atmospheric input may be therefore the only source of bioavailable metals to the marine system, and those metals can play an important role in the marine biological productivity (Duce and Tindale, 1991 (9); Young *et al.*, 1991 (10)). In recent years several authors have studied the trace metal dry, wet and/or total deposition in the Western MED (Dulac et al., 1989 (11); Remoudaki et al., 1991 (12); Migon et al., 1991 (13), 1997 (14) and a very large data set obtained during the EROS 2000 project has permitted to establish a reliable dissolved/particulate input budget to the NW MED (Guieu et al (1997 (7)).

There has been considerable debate as to metal solubility estimates (Jickells, 1995 (15) although a reasonable consensus on the range of solubilities observed for environmentally important metals in rainwater has now been reached (Losno et al., 1993 (16); Lim et al., 1994 (17); Spokes et al., 1994 (18)). Previous studies have also emphasised the importance of pH and aerosol type in controlling aerosol dissolution (Chester et al., 1993a (19); Lim et al., 1994 (17); Spokes et al., 1994 (18)). This study aims to (i) identify how concentration, mineralogy, grain size and chemistry of the insoluble fraction control aerosols metal partitioning solid and solution phases, and to (ii) estimate the importance of desert dust in the atmospheric trace metal input.

SAMPLING AND ANALYTICAL METHODS

Sampling

Chemical data of the soluble and insoluble fraction of aerosols collected at a land-based coastal station in SE Sardinia, Western MED (9° long E, 39° lat N) (Fig. 1) are presented. A total 88 samples (55 aerosol; 33 rain) were collected during the period October 1990/April 1992 and presented here. Aerosol samples were collected with high volume (60 m³ h⁻¹) Sierra Andersen[®] sampler, using filters made of polyester as well as Whatman 41 filters. Sampling time was 72 hours. A detailed description of sampling site and characteristics of the aerosols can be found in Guerzoni et al. (1997) (2). Three separate quarters of the filters were then used for three separate analyses: (i) acid digestion for total trace metal analyses; (ii) water extraction for major anions; (iii) leaching experiment for dissolved trace metal analyses. Filter chemical blanks were determined from filters that were taken into the field and handled in the same manner as those used for sample collection.

Rain samples were collected using a Wet & Dry (ARS 200 MTX[®]) automatic collector, and the sample is filtered through a 0.2 µm diameter Nuclepore[®] polycarbonate preweighted filter within few hours, and subdivided into two phase fractions, soluble and insoluble. The insoluble fraction was dried in a dry-box and re-weighted (to obtain the total particulate concentration). Blanks were run periodically, by making all operations with only Milli-Q[™] water instead of samples. Filtering was done with a Millipore[®] Sterifil unit, equipped with a pre-filter on the cover, to avoid contamination with ambient air. The analyses of the filtrate and residual fraction on filters defined the dissolved and particulate metal concentrations.

All manipulations were conducted in clean rooms equipped with a laminar flow hood.

Trace metal dissolution experiments

The laboratory leaching experiments were designed for the extraction of the water soluble components of the particles collected on filters.

In the experimental procedure portion of filters (one quarter) on which aerosol had been collected were folded and placed in acid-cleaned Teflon bottles (250 ml). We used different particulate amounts, choosing from our filter library appropriate concentration of particulates onto the filter for each sample. The particulate concentrations were known by separately weighing another portion of the filter. Filters were leached with 100 ml of ultrapure deionised water (Milli-Q[™]), on a reciprocating shaker at high speed for one hour. Then the filters were removed and the solutions were filtered through a 0.2 µm Nuclepore[®] polycarbonate filter, and the metals recovered in the filtrate were operationally considered as being dissolved. Dissolution in rain water and sea water of metals attached to aerosols occurs in short time (Statham and Chester, 1988 (20); Guieu et al. 1994 (21) therefore, we assumed that the exposure time chosen for this experiment allows to simulate the natural conditions. The dissolution experiments were performed at particle concentrations ranging from ≈3 to ≈300 mg l⁻¹, at an initial pH of the Milli-Q[™] water of 5.0-5.5. After filtration, the solution pH was measured and the filtrate was then acidified to pH 2. The pH of the added water was always roughly the same so the final pH, after leaching, was altered by aerosol itself, thus obtaining the range of

experimental pH values. The solutions were then analysed for trace metals using flame and flameless atomic absorption techniques, as appropriate.

Trace metal analysis

Another quarter of the aerosol filter was digested following dissolution in HNO₃/HF in teflon bombs. Trace metal concentrations were determined by graphite furnace atomic absorption spectrophotometry (GFAAS). Sampling methods, sample treatment and analytical methods are described in detail in Guerzoni et al. (1993 (22); 1997 (2) and Molinaroli et al. (1993) (23).

To determine the precision and the accuracy of trace metal analyses by GFAAS, certified standard solutions were regularly analyzed throughout the period of study. Results showed that analytical accuracy was of the order of $\pm 5\%$, and precision was of the order of ± 5 and 10% based on analysis of replicate samples and both solid and aqueous standards. Detection limits (DL), defined as three times the standard deviation of the blank, were calculated from replicated analysis of ultrapure deionised water (Milli-Q™) at pH 5.6, the matrix used in the experiment. For the elements of interest, DL values were: Al and Si $0.5 \mu\text{g l}^{-1}$, Cd $0.04 \mu\text{g l}^{-1}$; Ca, Cr, Cu and Fe $1 \mu\text{g l}^{-1}$; Pb $0.4 \mu\text{g l}^{-1}$.

Determination of blanks

Operational field blanks were performed using precipitation and aerosol sampling protocols at sampling sites by deploying and momentarily exposing "clean" rain collectors and filters between precipitation and aerosol sampling events. Blank concentrations of dissolved Al, Fe, Cd and Pb ranged between ± 5 and 10% trace metal concentrations in the precipitation analysed. For particulate trace metals, clean filters were taken through the in HNO₃/HF digestion, and the resulting blank solution were analysed. As a percentage of trace metal concentration in particulates, filter blanks were as follows: Al, Ca, Fe, Mg, Si, ± 5 ; Cr, Pb, Zn $\pm 10\%$; Cd, $\pm 12\%$. All trace metal concentrations reported here are blank corrected

RESULTS AND DISCUSSION

Measurement in air and rain

The atmosphere is a very dynamic compartment of the earth system and the concentrations of reactive gases and particulate matter are highly variable in space and time. This variability ranges from rapid day to day changes in aerosol lead concentration as a result of changes in atmospheric transport and removal (Fig. 2a), to massive changes in dust loadings over Sardinia, a result of seasonal dust outbreaks in Africa (Fig. 2b). The peculiar influence of dryness on aerosol samples collected on MED islands was discussed by Bergametti et al. (1989) (4) and was also found in Sardinia, where a three-fold increase in mass loading for the samples collected with more than three rainless days was shown (Guerzoni et al., 1995) (24).

Previous work (Guerzoni et al., 1997) (2) has shown that at sites relatively far from pollution sources it is possible to de-couple the desert-derived "end-member" aerosol

from the "BKG" component. A multiple approach can be adopted to achieve this: (i) the use of air-mass back trajectories to define source regions (Harris et al., 1982) (25); (ii) the fact that much of the SAH dust over the MED Sea is transported in the form of "pulses", and the identification of these "pulses" offers a potential way of isolating the desert-derived component; (iii) other combined criteria which could be used to classify BKG and SAH aerosols include the use of mineralogical markers (Molinaroli, 1996) (26), Al/Si ratio (Bergametti et al., 1989) (4) and Ca content (Molinaroli et al., 1993) (23). Finally, for precipitation, (iv) pH values were also used to improve classification, since the alkalization effect of SAH dust in rain has been well described, both in Sardinia (Caboi et al., 1992 (27); Le Bolloch and Guerzoni, 1995 (28) and elsewhere (Löye-Pilot et al., 1986 (29); Glavas 1988 (30); Roda et al., 1993 (31).

The SAH outbreaks in our data set represent less than 15% of the total sample (7 out of 55) for the aerosols, and slightly more than 30% of the rainwater samples (10 over 33). The strongest episodes occurred in two periods: October-November, and March-April, and principally with mixed, SW and E-SE winds.

In Table 1 we have listed the particulate load and grain-size characteristics of the aerosols collected at Sardinia. The data are divided in two groups: SAH, mainly desert-enriched aerosol, and BKG, mainly anthropogenic-maritime enriched aerosol. From this table it is apparent that the soluble fraction, mainly composed of sea-salts (70-95%), is similar in SAH and BKG aerosols (25 and 16 $\mu\text{g m}^{-3}$), whilst the most important difference is in the insoluble (mineral) fraction, highly enriched in SAH episodes. Grain-size data also show an enrichment in fine-grained fraction ($<1\mu\text{m}$) in the BKG, as compared to coarser enrichment (2-8 μm) in the SAH aerosols. Table 2 lists the average concentration of particulate load encountered in Sardinia during the sampling period, where a range of more than two orders of magnitude is apparent, both in aerosol and rainwater samples.

In Tables 3 and 4 the relevant metal data of aerosols and precipitation collected at Sardinia are listed together with the average crustal abundance (EF_{crust}), and the percentage of dissolved fraction. The data set given in the two tables provide examples of the contrasting atmospheric chemistries of the trace metals at the Sardinia site.

(i) The SAH samples have always higher values, of all the crustal elements (Al, Ca, Fe, K, Mg, Si) than BKG, both in aerosols and rainwater.

(ii) Al, Fe and Si are mostly in the insoluble form (range of solubility 0.4-10%), whilst Ca, K and Mg are mainly (50-90%) in the dissolved fraction.

(iii) Cd, Cr and Pb have lower solubilities in aerosol as compared to rainwater, both in SAH and BKG samples.

Mixture of Crustal/European aerosol (EF 's)

The degree to which a trace metal in aerosol or rain water is enriched or depleted, relative to the crustal source, can be assessed by using an enrichment factor (EF_{crust}). For crustal sources, Al, is normally used as the source indicator element and the Earth's crust as the source material (Taylor, 1964) (32). The EF_{crust} value is then calculated according to an equation of the type:

$$EF_{\text{crust}} = (C_{xp}/C_{Alp}) / (C_{xc}/C_{Alc})$$

where (C_{xp} and C_{Alp}) are the concentrations of a trace metal x and Al , respectively, in the aerosol, and (C_{xc} and C_{Alc}) are their concentrations in average crustal material.

When the concentrations of trace metals in aerosols are plotted on EF diagrams the overall population can be described in terms of mixing of anthropogenic-rich components, which have high EF_{crust} values ($EF_{crust} > 10$) for all trace metals and crust-rich components, which have low EF_{crust} values ($EF_{crust} < 10$) for all trace metals. Enrichment Factors of the Sardinia samples are reported in Tab. 3 and 4.

Previous work (Chester et al., 1993a) (19) has described the "chemical character" of the MED atmosphere containing both crust derived and urban derived aerosols. At land-based stations on the northern part of West MED the aerosols have European-dominated "chemical character" similar to that of other Western European aerosol population (i.e. North Sea). In contrast, aerosol from the SAH region to the South have a crust dominated "chemical character". Thus, the chemical composition of the MED particulate aerosol can be considered to be largely constrained by the extent to which "BKG" material is perturbed by mixing with desert material.

In Figure 3 our data from Sardinia are displayed on the EF diagrams together with trace metals in aerosol at two coastal sites (Blanes and Cap Ferrat) (Chester et al., 1993a) (19). For all the aerosols Fe displays low EF_{crust} values (Fig.3A). Therefore, the Fe is a crust-dominated trace metal under all conditions in the aerosol from the MED and surrounding regions.

On the EF diagram Zn (Fig. 3B); and Pb (Fig. 3C) both plot in two fields One is the "mixing relationship" field, and the other is where this field is perturbed at the upper right hand corner as a result of the localised inputs of urban-rich material at the Blanes and Cap Ferrat sites. Thus, over the MED and surrounding regions inputs of crustal material are not sufficient to force Zn and Pb to behave as a crust-dominated trace metal. The Sardinia samples fall in the "mixing relationship" field; i.e. the Sardinia aerosol shows none of the "extreme" perturbations resulting from the local inputs of either anthropogenic-rich or crust-rich material such as those shown by aerosols collected at the Blanes and Cap Ferrat sites. Thus, the $W \Rightarrow E$ fall off in both (i) trace metal concentrations and (ii) trace metal EF_{crust} values along the length of the Western Mediterranean results, at least in part, from the input of local anthropogenic-rich material, which most strongly affects Cd, Zn and Pb, at the Blanes and Cap Ferrat sites; inputs which are not recorded in the aerosols collected at the Sardinia site in sufficient quantities to perturb the "mixing relationship" between crustal and anthropogenic material. In terms of its "chemical character", therefore, the Sardinia aerosol may be distinguished from those aerosols collected at the more mainland coastal sites surrounding the Western Mediterranean.

Dissolution in water of metals associated to aerosols

In order to study the role of desert particulates in the dissolution processes, we have performed a leaching experiment, concentrating on a few metals of both crustal (Al, Fe) and anthropogenic (Cd, Pb) origin and trying to identify process which affect dissolution. The reason of this kind of study is related to the findings of Chester *et al.* (1993a) (19), who have shown that a perturbation in the urban/crustal aerosol ratio results in a change in the dissolved/particulate partitioning. The leaching experiment is

also connected to the recent work of Spokes and Jickells (1996) (33), who identified additional processes related to particulate load which affect solution phase removal. Studies of the chemical and physical properties of atmospheric particles of different areas of the world (Rahn, 1976 (34); Prospero, 1981 (35); Duce *et al.* 1991 (1) have allowed the following conclusions to be drawn: (i) the large particle fraction ($> 1\mu\text{m}$ aerodynamic diameter) derives mostly from natural sources in arid areas of the world and is responsible for the release of Al, Fe, Si and P to ocean waters. (ii) Smaller particles ($< 1\mu\text{m}$) are predominantly of anthropogenic origin and lead to the enrichment of elements such as Pb, Cu, Zn, Cd, N, As and Hg, especially in the Northern Hemisphere. (iii) Both size fractions contribute Mn, V and Cr to surface waters.

Some problems arise in assessing the trace metal solubility for “dry” deposition samples, especially from large natural particles. Using theoretical deposition velocities will give a sensible estimate of the dry deposition flux. The question is how to assess metal solubility for this flux, since dry deposition is particle-sized constrained and is more “crustal” in character than the aerosol from which is derived. Because of this, using data for the parent aerosol may considerably over-estimate the trace metal solubility in “dry” deposition.

To partially overcome this problem, Chester *et al.* (1993b) (36) have outlined a relationship between the EF_{crust} value and the seawater solubility, where Cu, Zn and Pb have higher seawater solubilities in the urban-rich than in crust-rich population. Therefore it should be possible to correct our laboratory data by using the EF values.

Leaching experiment

We have been conducting particle/water interactions studies using aerosol concentrations, pH conditions and timescales representative of those in rainwater. This study used aerosols that can be considered SAH end-members (whilst not “pure” SAH dusts), as well as aerosols with an anthropogenic-dominated signature, with all the intermediate mixed samples. We decided not to adjust the pH, in order to avoid alteration in the dissolution rates of some minerals (Maring and Duce, 1989 (37); Wieland and Stumm, 1992 (38) and to observe the effect of aerosol itself on the pH.

We are aware that there are several inherent difficulties in applying the results to the “real” world, in particular:

(i) the experimental system does not allow us to reproduce completely the pH ranges likely to be encountered by aerosols, but rather assesses whether metal solubility is affected by particle mass, grain-size, and minerals, at pH values higher than 5.0-5.5;

(ii) the results of the experiments can be related directly to the rain water, but it is more difficult to extrapolate them to dissolution in sea water;

(iii) there might be a fractionation from the parent aerosol to the dry deposition, with a possible depletion of Cd and Pb in the latter one.

Al and Fe

Our Al and Fe solubilities are wide ranging (Al: 0.3-28%; Fe: 01-13%) and resulted to be mainly controlled by the particulate load. The percentage of aluminium and iron dissolution decreases significantly as particulate loading increases. Many studies have reported anomalous metal solubility behaviour dependent not only on the type of

particle surfaces present, but also on absolute particle concentrations (e.g. Stumm, 1992 (39); Morel and Gschwend, 1987) (40). In our experimental design, aluminium dissolution decreases significantly (Fig. 4a) from more than 20% at aerosol loadings lower than 5 mg l⁻¹, to less than 1%, at total aerosol concentrations greater than 100 mg l⁻¹, which are mass particle values usually detected during the SAH dust outbreaks. Same pattern was observed also for iron (Fig. 4b), even with much fewer samples analysed. At aerosol loadings less than 20 mg l⁻¹, an average of 6% of iron present in aerosol undergoes dissolution with this value decreasing rapidly to average 1% at total aerosol concentrations greater than 200 mg l⁻¹. In the case of iron, our results are in agreement with the findings of Spokes and Jickells (1996) (33). Those authors added varying amounts of Saharan aerosols (up to a particulate load of 140 mg l⁻¹) and showed that the percentage iron dissolution decreased significantly as particulate loading increased, although apparently not under solubility control. On the other hand, the same authors (33) showed a linear relationship of dissolved manganese and aluminium with particulate loading, with no evidence that the dissolution was under solubility control.

Our dissolved Al data showed a non linear increase in solution concentration with particulate load (Fig.4c) and the reason of the percentage aluminium dissolution decrease (Fig. 4a) with increasing particulate load is unclear but may result from:

(i) aerosols acting as adsorption substrates, at high particulate loadings, with the resultant removal of dissolved metals from seawater (36), rather than providing a source through desorption;

(ii) increased solid aggregation with increasing particulate load leading to a net decrease in the number of Al desorption sites (33);

(ii) partial formation of Al hydroxide compounds. Losno et al (1993) (16) have shown that at pH >5, which are our experimental conditions, the precipitation of Al hydroxide compounds (e.g. Al(OH)₃, gibbsite) or a trivalent basic insoluble Al salt is likely.

Results for Al are in contrast with Wieland and Stumm (1992) (38) who show that aluminium dissolution from kaolinite, which represents 20-30% of the total clay mineral of SAH samples (Molinarioli, 1996) (26), increases linearly with suspended load.

We have calculated K_d values for Al using the formulation of Hydes and Kremling (1993) (41):

$$C_d = \frac{f[S]}{(1 + K_d S)}$$

where: C_d= dissolved metal concentration (g ml⁻¹); F(S) = geochemically available metal in solid (g ml⁻¹); K_d = distribution coefficient (ml g⁻¹); S=particulate load (g ml⁻¹) to predict dissolved metal concentrations given a knowledge of particulate load.

Calculated K_d values were not constant, and range from 2 x 10⁵ to 6 x 10⁶ (assuming all the aluminium in the aerosol is available for dissolution), showing some anomalous particle concentration effect. The higher K_d values were at S < 25 mg l⁻¹ with some decrease with increasing to S > 100 mg l⁻¹, probably due to colloids (Honeyman and Santchi, 1988 (42). It was impossible to fit the data from this study to the Hydes and

Kremling equation, at any single K_d value, and we could not try to fit the data by using a more appropriate fraction (e.g. loosley bound) since we did not performe any solid state speciation experiment on this aerosols (36).

Further experiments are needed, in order also to check the role of previously weathered primary minerals (Giusti *et al.*, 1993) (43), and of the grain-size of the particles on the particle load control on aluminium solubility.

Pb and Cd

Previous work has shown that at Sardinia Cd and Pb are of predominantly anthropogenic origin. In Figure 5 the dissolved fraction of Cd and Pb is plotted versus the particulate load. Solubilities of the two metals are also quite different. Solubility for Pb range from ≈ 1 to $\approx 50\%$, and there is a clear exponential inverse relationship with the mass particle content of the aerosol (Fig. 5a), with some evidence that the dissolution was under solubility control: a decrease from 100 to 40 mg l^{-1} in the particulate load content lead to an increase in solubility from 5 % to 40% . Whilst only a limited number of samples (30 over 55) were analysed for Cd, due to technical problems, all of them show a solubility in the range from 70% to almost 100%. In particular, only one SAH dust was available for this leaching experiment, and therefore the linear relationship shown in Fig 5b is mainly related to anthropogenic-dominated aerosols. Our data, even if limited confirm the findings of Chester *et al.* (1993a) (19), who has already noted that Cd is relatively very soluble ($> 80\%$) from both the crust-rich and the urban rich aerosols.

In Fig. 6 we have plotted the % of dissolved lead with grain-size $< 1\mu\text{m}$, the crustal Enrichment Factor (EF_{crust}) and pH.

The solubility of Pb shows a good linear relationship with the finest grain-size fraction, ($r= 0.69$, $p<0.001$), with an evident increase in solubility with higher content of clay fraction ($<1\mu\text{m}$) of aerosol (Fig. 6a). The solubility of Pb appears also to be highly correlated ($r= 0.70$, $p<0.001$) to the EF_{crust} value (Fig. 6b).

Chester *et al.* (1993a) (19) showed that Pb has higher EF_{crust} value and higher solubility in the urban-rich than in the crust-rich aerosol population. However they pointed out that in “mixed” aerosols re-scavenging by crustal material can affect Pb solubility, as a proportion of the metal initially released is taken back out of solution.

In Fig. 6c the percentage dissolution of lead as a function of pH is shown, and in the same figure we have also illustrated the theoretical pH/solubility curve studied by different authors, both in rain and in aerosol (Chester *et al.* 1993a (19); Lim *et al.* 1994 (17); Spokes and Jickells, 1996 (18)). The relationship between Pb solubility and rainwater pH exhibits the classical pH adsorption edge, which is typical of adsorption/desorption processes at particulate-water interfaces. However, some authors note that the low solubility region is defined by a relatively small number of points, reflecting the few cases studied of rain with $\text{pH} > 6$ (Lim *et al.* 1994 (17); Spokes *et al.* 1994 (18)). This observation is important, given that the rains normally contain variable amounts of particulate material and that particle concentrations, grain-size, and physical surface characteristics (scavenged clays or silicate mineral aerosol) may also be important parameters in adsorption models of trace metals in rain (Lim *et al.*, 1994 (17)).

From Fig. 6c we can see that most of the samples plot inside the pH/solubility “area” we have drawn with $\pm 10\%$ of tolerance with respect to the ‘classical’ curve. Two groups of samples plot out this area. Few samples have lower than expected solubilities at pH value $\cong 5.5$. Those samples are ‘mixed’ aerosols (see e.g. Chester et al., 1993a (19) terminology), where some re-scavenging from crustal material is possible. In this case the crustal material eventually does not directly affect the solubility of the anthropogenic dominated part of the mixed aerosol, but behave as an external factor taking back from solution a proportion of the metal initially released. On the other hand, another group of samples plot out the adsorption curve, with higher solubilities as expected at pH values > 6.5 . A mineralogical inspection of the samples showed that those aerosols contains cerussite minerals, derived by sphalerite and galena old mine tailings, located at about 1-200 km away, that very rarely reach the sampling sites. Those samples were already described by Guerzoni et al., (1997) (2), and have average grain-size of 2-4 μm . Those minerals are very often composed of carbonates, that can eventually leach out part of the lead, at pH values as high as 6.5-7.5 (Caboi et al., 1992) (27). After having discarded the cerussite-enriched samples the pH/solubility correlation is quite good ($r = -0.60$, $p < 0.001$).

Therefore, in order to understand which were the most important parameters influencing the lead partitioning we have applied a multiple regression analyses to the reduced aerosol data-set, without the cerussite-enriched samples. The percentage of dissolved lead was taken as dependent variable, and we compared the relative importance of EF_{crust} , grain-size $< 1\mu\text{m}$, and pH in the variation of solubility. The results of this simple model showed that the variation in the dissolution of lead depends more from EF_{crust} , and grain-size ($\sim 40\%$ each) and less from pH ($\sim 20\%$). This is evidently not a general conclusion, but is useful to highlight the multiple approach needed to understand the processes which alter metal solubility of aerosols.

Metal fluxes at Sardinia

Several authors have attempted to calculate metal fluxes from the atmosphere and have encountered several difficulties. Whilst the evaluation of the wet input, calculated from rainwater concentrations and rainfall amount is generally reliable, the estimation of dry deposition is more problematic. Despite the shortcomings in these estimates, Guieu et al (1997) (7) have made a compilation of available data on wet and dry fluxes around the Western MED Sea. Those authors have shown for many trace elements (e.g., Al, Pb, Mn, Fe) that the global flux from the atmosphere entirely dominates the total external input of pollution-derived elements, and that the atmosphere is also very important for elements of terrigenous origin such as Al and Fe.

In this paper fluxes were calculated from rainfall and aerosol data listed in Tables 3 and 4. For precipitation samples the “wet” flux (e.g. in $\text{mg m}^{-2} \text{yr}^{-1}$) was calculated by multiplying the metal concentration (in $\mu\text{g l}^{-1}$) by the amount of rain (litres) and dividing by the area of the sampler (0.066 m^2). For the aerosols, the “dry” flux (F_x) (e.g. $\text{mg m}^{-2} \text{yr}^{-1}$) was calculated according to the following equation: $F_x = c * \underline{V}_d$, where \underline{C} is the metal concentration value in ng m^{-3} and \underline{V}_d is the deposition velocity in cm sec^{-1} . The deposition velocities used here were chosen within the range of literature values (Dulac

et al., 1989 (11); Slinn and Slinn, 1980 (44); Migon et al., 1997 (14). The values were 2 cm sec^{-1} , for Al, Ca, Fe, Si, metals that derive principally from aerosols largely dominated by mineral particles, with a median grain-size of 2-10 μm . The selected deposition velocity for Cd and Pb was 0.2 cm sec^{-1} , because those are elements of predominantly anthropogenic origin, associated to finer grain-size ($< 1\mu\text{m}$), and 1.0 cm sec^{-1} was chosen for Cr, that appears to be related to both size fractions (Giusti et al., 1993) (43).

From aerosol and precipitation data collected at Sardinia we have calculated the dissolved and total fluxes separately, and the fraction of these fluxes that can be attributed to SAH dust.

As previously mentions, some problems arise in assessing the “dry” deposition of the anomalously enriched elements such as Cd and Pb. We are aware that there is a fractionation from the aerosol to the dry deposition, which is largely particle-sized mediated. This fact is probably not so important in the estimate of the dry fluxes of non-enriched elements (Al, Ca, Fe, Si and Cr). On the contrary, the dry fluxes of Cd and Pb may be considerably over-estimated, and our data can be seen as an upper limit, in case of SAH enriched episodes.

Model estimates yield a mineral dust deposition from North Africa of 170 Tg yr^{-1} to the Atlantic, 25 Tg yr^{-1} to the MED and 5 Tg yr^{-1} to the Caribbean (Prospero, 1996) (6). There is also an extremely large seasonal and inter-annual variability, in dust deposition and we might expect a comparable variability in consequent ocean effects.

Estimates ($\text{mg m}^{-2} \text{ yr}^{-1}$) of the fluxes (dry+wet) of soluble and insoluble fractions and of the percentage that are due to Saharan dust contribution, for samples collected at Sardinia are listed in Table 5. On the basis of these data, the trace metals can be divided into three broad groups.

Group 1: which includes Al, Fe and Si. For these trace metals the SAH fluxes over the total atmospheric fluxes exceeds 70%, and the the insoluble fraction is prevalent.

Group 2: which includes Ca and Cr. The contribution to the total deposition made by SAH is ~60%, and the dissolved fraction decreases from Ca (59%) to Cr (38%).

Group 3: which includes Cd and Pb. The contribution to the total deposition made by SAH is ~30%, but the particulate/dissolved partitioning in the SAH and BKG varies between the two trace metals. For Pb the particulate fraction due to SAH is almost 40%, whilst the dissolved fraction is less than 10%. For Cd the dissolved fraction dominates the flux ($64 \text{ vs } 7 \mu\text{g m}^{-2} \text{ yr}^{-1}$) and the fraction due to SAH is almost one third of the total.

CONCLUSIONS

1. Data on aerosol and precipitation collected at a remote site in Sardinia are presented. The partitioning of total atmospheric deposition between the soluble and the insoluble phases shows that Al, Si, Fe and Pb originating from the atmosphere are mostly (>90%) in the insoluble form. For the other metals studied (Ca, Cd, Cr), the dissolved fraction represents 50-90 % of the total input.
2. In terms of its "chemical character", the Sardinia aerosol may be distinguished from those aerosols collected at the two sites of Blanes (Spain) and Cap Ferrat (France). In particular, the BKG aerosols show a 3 to 25-fold decrease in EF_{crust} values for Cd, Zn, Pb, and Cr compared with coastal stations in the Western Mediterranean.
3. We have presented a series of novel experiments, carried out in closed system, which explored the role of particle concentrations in constraining metal solubilities. The principal conclusions are:
 - (i) at high particle concentrations the solubilities of Al, Fe, Pb and Cd decrease;
 - (ii) Al and Pb solubilities decrease with increasing pH values;
 - (iii) the solubility of Pb increase with increasing EF values and decreasing particle size
4. Estimates of the fluxes (dry +wet) of metals show that $\cong 30\%$ for Cd and Pb, to more than 80 % for Fe and Al are due to Saharan contribution.

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Tab. 1- Particulate load and grain-size characteristic of Saharan and Background aerosols (from Guerzoni *et al.*, 1993 (22)).

		Saharan	Background
Particulate load	$\mu\text{g m}^{-3}$ soluble	25	16
“	$\mu\text{g m}^{-3}$ insoluble	26	4
< 1 μm	%	8	24
1-2 μm	%	21	20
2-8 μm	%	40	22
8-32 μm	%	27	27
>32 μm	%	4	7

Tab. 2 - Concentration of particulate load in air and rain at Sardinia (from Guerzoni *et al.*, 1997 (2))

	air ($\mu\text{g m}^{-3}$)	rain (mg l^{-1})
Background	1-3	0.1-10
Desert-enriched	20	10-50
Saharan outbreaks	44	50-500
Maximum recorded	110	510

Tab. 3 - Relevant data of the aerosol collected at Sardinia. The data are divided in two groups: Saharan, mainly desert-enriched aerosol, and Background (mainly anthropogenic-maritime enriched aerosol). Mean total concentrations, enrichment factors (EF_{crust} , see text for explanation) and the percentage of dissolved fraction are indicated, on Saharan and Background samples respectively.

Element	Saharan (n=7)			Background (n=48)		
	Conc. ($ng\ m^{-3}$)	EF_{crust}	Sol. (%)	Conc. $ng\ m^{-3}$	EF_{crust}	Sol. (%)
Al	2929	---	(1)	210	---	(6)
Ca	3364	3	(81)	629	15	(96)
Fe	1138	1.7	(0.2)	84	8.3	(10)
Si	5875	0.8	(1)	609	1.3	(6)
Na	5066	11	(87)	4084	268	(95)
K	753	1.5	(50)	280	23	(83)
Mg	1271	2	(90)	615	30	(95)
Cd	0.08	12	(57)	0.17	80	(88)
Cr	3.3	1.1	(20)	0.5	3.3	(52)
Pb	25.3	66	(9)	11.7	550	(20)

Tab.4 - Mean total concentration ($\mu g\ l^{-1}$) and enrichment factors (EF_{crust} , see text for explanation) of major elements and trace metals for rainwater, divided in Saharan and Background. In brackets the percentage of dissolved fraction. (*) Particulate load is in $mg\ l^{-1}$.

Element	Saharan (n=10)			Background (n=23)		
	Conc. ($\mu g\ l^{-1}$)	EF_{crust}	Sol. (%)	Conc. ($\mu g\ l^{-1}$)	EF_{crust}	Sol. (%)
Al	2780	---	0.4	228	---	6
Ca	6530	4	92	1430	11	98
Fe	1523	0.8	0.8	136	0.9	10
Si	7894	0.8	5	1021	12	32
Na	15890	21	99	7725	125	99
K	909	1.3	39	331	5.8	59
Mg	860	0.9	52	226	2.9	70
Cd	0.15	22	87	0.10	183	90
Cr	0.91	0.2	49	0.48	16	63
Pb	1.9	3	32	13.9	281	77
Particulate load*	93		59	67		94

Tab. 5 - Estimates ($\text{mg m}^{-2} \text{ yr}^{-1}$) of the fluxes (dry+wet) of soluble and insoluble fractions and of the percentage that are due to Saharan dust contribution for samples collected at Sardinia.

Element	Insoluble	ins-Sah	%ins-Sah	Soluble	sol-Sah	%sol-Sah	Total	tot-Sah	%tot-Sah
Al	891	734	0.82	16	6	0.36	907	739	0.82
Ca	175	154	0.88	1734	1029	0.59	1909	1183	0.62
Fe	406	330	0.81	10	2	0.18	416	332	0.80
Si	2143	1666	0.78	166	49	0.30	2309	1715	0.74
Cd	0.005	0.002	0.45	0.050	0.014	0.29	0.05	0.016	0.30
Cr	0.36	0.26	0.72	0.26	0.10	0.38	0.62	0.36	0.58
Pb	1.19	0.49	0.41	1.10	0.10	0.09	2.29	0.59	0.26

FIGURE CAPTIONS

- Fig.1 The Mediterranean basin with the location of sampling station.
- Fig. 2- Variation of aerosol lead (above, ng m^{-3}) and mineral suspended particulate (below, $\mu\text{g m}^{-3}$) concentrations over a seven-months period (Oct '90-Apr '91) at a remote site in Sardinia (data from Guerzoni et al., 1997), illustrating the large-scale short-term variability.
- Fig. 3 - EF diagrams for Fe, Zn and Pb for the Sardinia particulate aerosol (\blacktriangle) superimposed on data from two mainland coastal stations: Blanes (+) and Cap Ferrat (o) (from ref. 19).
- Fig. 4 -Percentage dissolution of aluminium (above) and iron (below) in aerosols collected at Sardinia station as a function of particulate load. Filled dots represent SAH samples.
- Fig. 4c Percentage dissolution of aluminium in aerosols collected at Sardinia station as a function of pH
- Fig. 5 - Percentage dissolution of lead (above), cadmium (below) in aerosols collected at Sardinia station as a function of particulate load. Filled dots represent SAH samples.
- Fig. 6 - Percentage dissolution of lead as a function of grain-size fraction $<1\mu\text{m}$ (above), of enrichment factor (centre) and pH (below) in aerosols collected at Sardinia. Open squares are the cerussite-enriched samples, and filled triangles are mixed aerosols (see text for explanations).

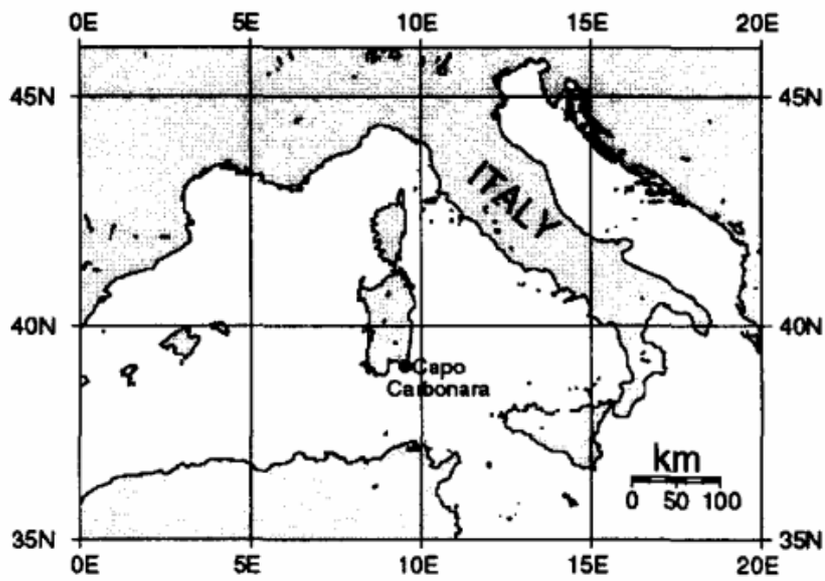


FIG. 1

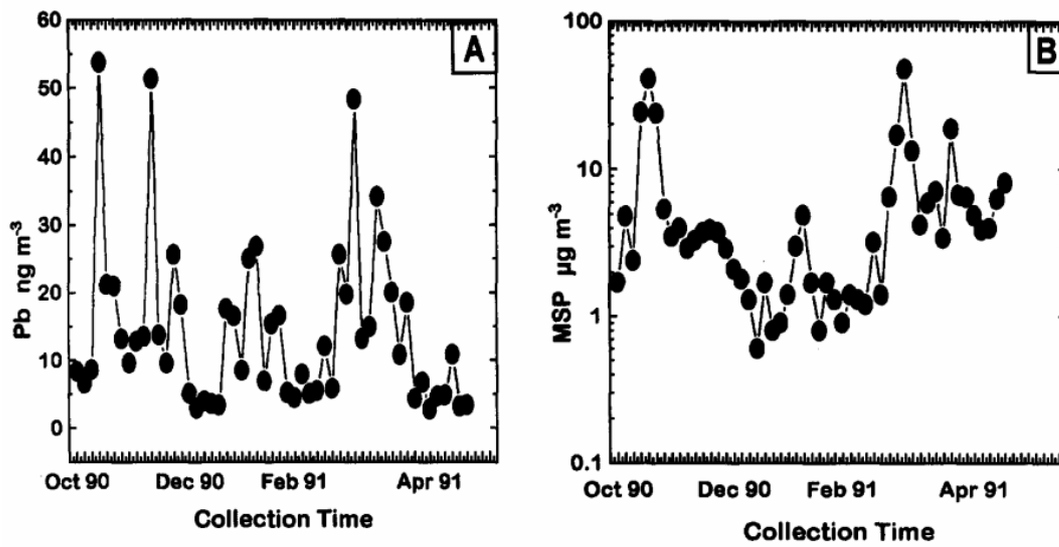


FIG. 2

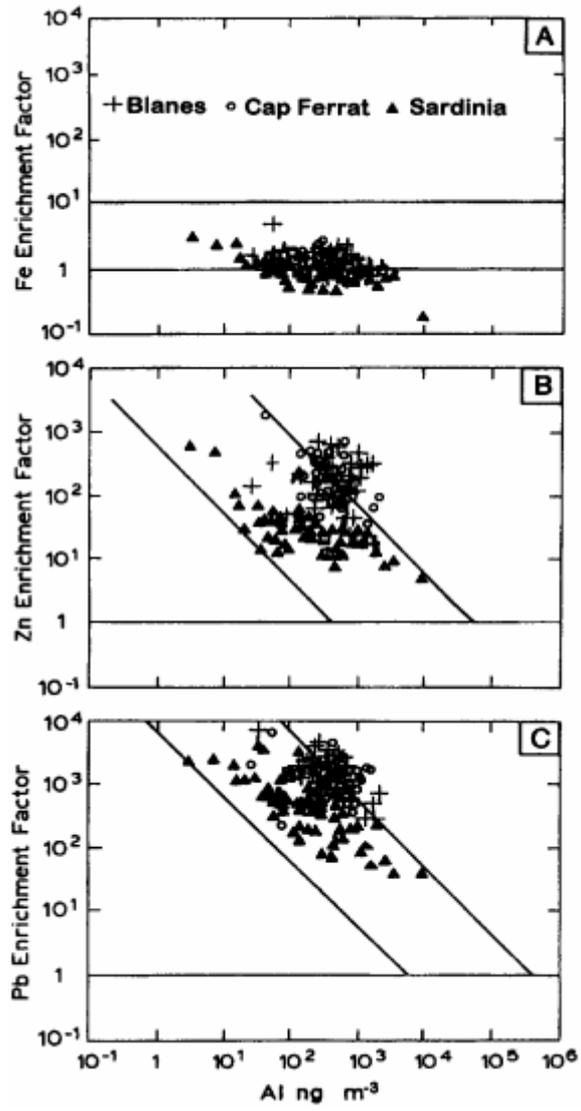


FIG: 3

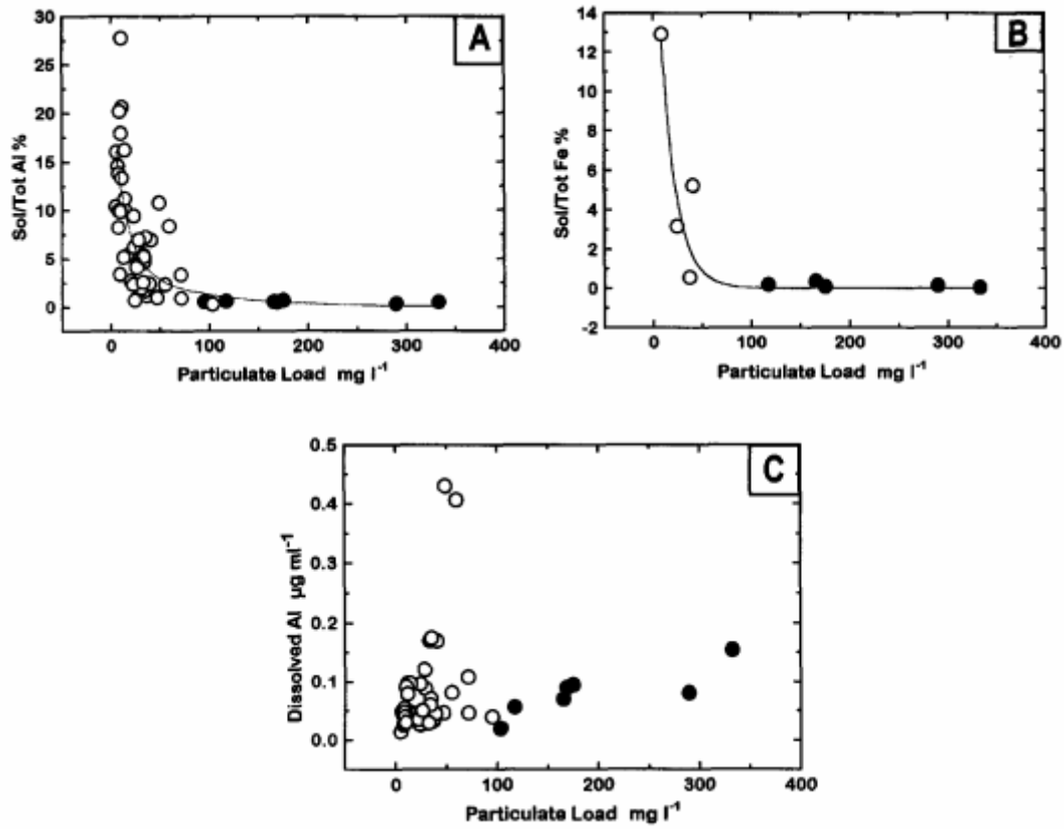


FIG: 4

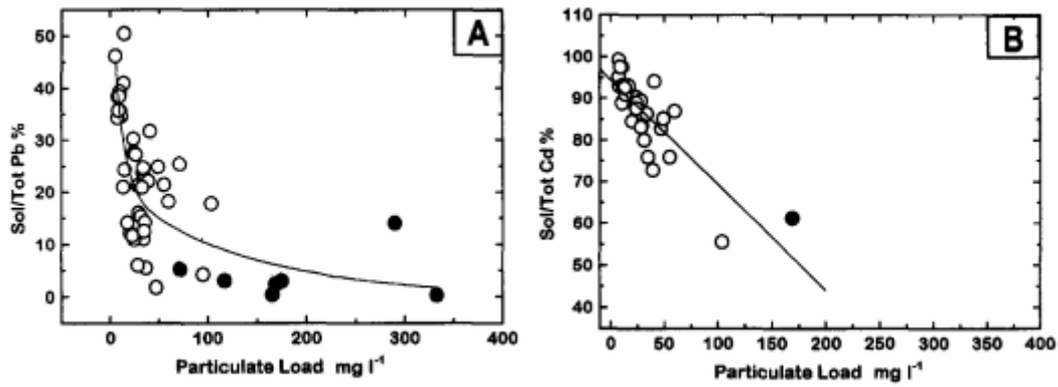


FIG: 5

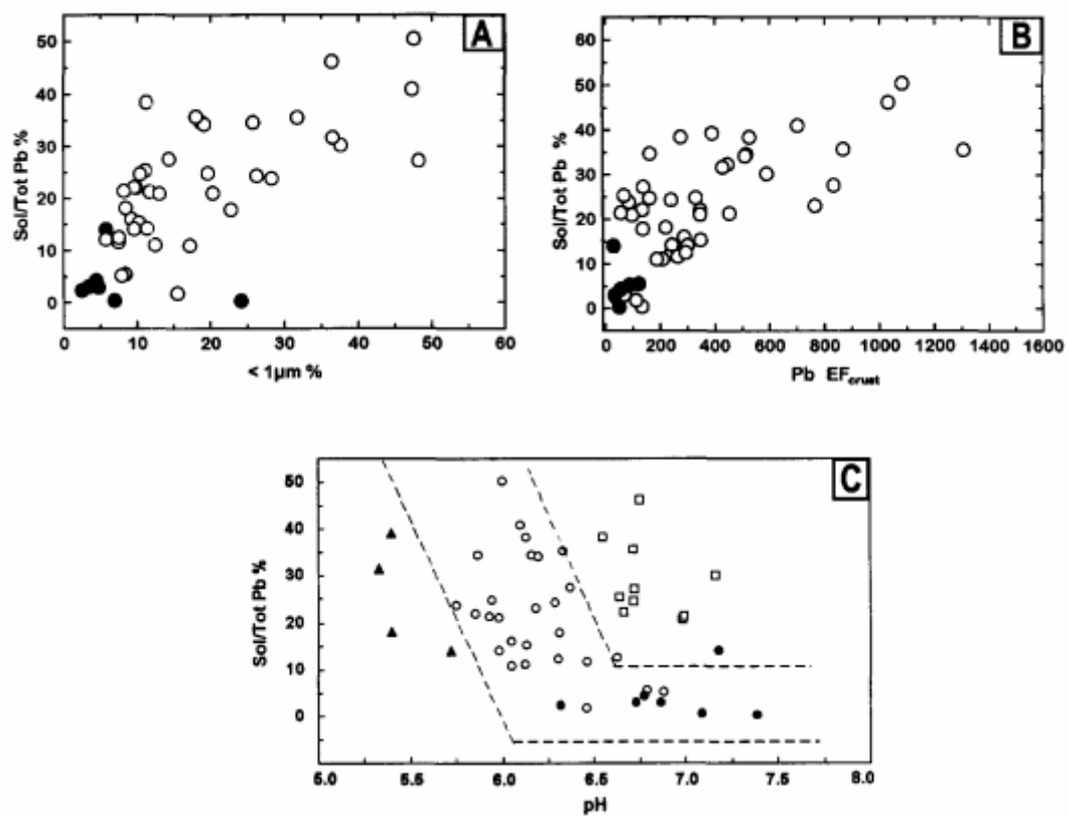


FIG. 6