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Geochemistry of natural and anthropogenic fall-out (aerosol and precipitation) collected in N W Mediterranean: two different multivariate statistical approaches.

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ABSTRACT

The chemical characteristics of the mineral fractions of aerosol and precipitation collected in Sardinia (NW Mediterranean) are highlighted by means of two multivariate statistical approaches. Two different combinations of classification and statistical methods for geochemical data are presented. It is shown that the application of cluster analysis subsequent to Q-Factor analysis better distinguishes among Saharan dust, Background pollution (Europe-Mediterranean) and Local aerosol from various source regions (Sardinia). Conversely, the application of simple cluster analysis was able to distinguish only between aerosols and precipitation particles, without assigning the sources (local or distant) to the aerosol. This method also highlighted the fact that crust-enriched precipitation is similar to desert-derived aerosol.

Major elements (Al, Na) and trace metal (Pb) turn out to be the most discriminating elements of the analysed data set.

Independent use of mineralogical, granulometric and meteorological data confirmed the results derived from the statistical methods employed.

Keywords: multivariate statistical analysis, aerosol, precipitation, geochemistry Mediterranean.

INTRODUCTION

Statistical analysis in atmospheric studies is a well-known and well-developed field (Morandi et al., 1987; Hopke, 1988; Van Malderen et al., 1992). This study presents an example of the use of cluster analysis (CA), Q-Factor analysis (Q-FA) and discriminant analysis (DA) in atmospheric geochemistry, in an attempt to classify mixed aerosol and precipitation samples on the basis of their major element and trace metal chemistry.

Atmospheric inputs to the Mediterranean are of particular interest, in view of the different areas of aerosol production on the north and south belts.

The principal source of natural "crustal" material in the Mediterranean is the Sahara, while industrialized and semi-industrialized Northern Europe is a source of anthropogenic "background" material to the atmosphere. These inputs have an important influence on both the mineralogical and chemical composition of the Mediterranean aerosol. Previous works have shown that the bulk aerosol in the lower troposphere in this area has two main components: sea salt aerosol and mineral aerosol (Bergametti et al., 1989; Correggiari et al., 1989; Chester et al., 1993a).

These aerosols are removed from the atmosphere to the sea surface by a combination of 'dry' (i.e., not involving aqueous phase) and 'wet' (precipitation scavenging) depositional modes, and these modes are geochemically different with respect to the solutions with which the aerosols come into contact. Estimates of dust transport and deposition in the Mediterranean have been made by the Group of Experts on the Scientific Aspects of Marine Pollution (GESAMP, 1989) using a model. The temporal variability of dust deposition rates may vary greatly because of the strong seasonal cycle in dust concentrations and the large seasonal variability of precipitation rates (Prospero, 1996).

The present study is based on a collection of six-month aerosol samples and one-year precipitation samples at a remote site. An overall sampling program was applied to investigate the geochemistry of natural (Saharan dust) and anthropogenic (Background) aerosols and to separate local versus distant inputs to the Western Mediterranean (Guerzoni et al., 1997). If and how crust-enriched precipitation is similar to desert-derived aerosol was also verified.

Two different ways of treating the data sets were planned, as follows:

- 1. a rank transformation was applied to the data and subsequently CA on the ranked data, followed by DA on the CA groups.
- 2. the data matrix was first reduced on the basis of Q-FA on the ranked data; then CA was applied to samples, with factor loading as new variables, and then DA on the CA groups.

The aims of this paper were to present the geochemical composition of insoluble aerosol and suspended particulate in precipitation and to show two different multivariate statistical approaches to the data sets; and then to translate statistical results into meaningful geological statements. The contributions of local versus distant sources were particularly highlighted.

SAMPLING

Chemical data of the insoluble fraction of aerosols collected at a land-based coastal station in SE Sardinia, Western Mediterranean (9° long E, 39° lat N) (Fig. 1) are presented, together with

measurements on mineral particulate deposited with precipitation. A total of 86 samples (55 aerosol; 31 rain) were collected during the period October 1990/October 1991, and 69 of these (55 aerosol; 14 rain) are presented here. The remaining 17 rain samples were not analysed due to the low concentration of the insoluble fraction ($<2 \text{ mg } \Gamma^1$).

In particular, aerosol sampling covered the period 1/10/1990-30/4/1991; precipitation was collected continuously during the period 1/10/1990-30/09/1991.

Two-to-three-day aerosol sampling (dry) and event-by-event rain collection (wet) were performed concurrently. Aerosol samples were collected with a high-volume ($60 \text{ m}^3 \text{ h}^{-1}$) Sierra Andersen[©] sampler, using filters made of polyester as well as Whatman 41 filters. Sampling time was approximately 72 hours (air volume sampled ~4000 m³ per sample).

Rain samples were collected using a Wet & Dry (ARS 200 MTX[©]) automatic collector.

ANALYTICAL METHODS

The filters used for aerosol sampling are washed in acidified DDW, dried overnight and weighed before sampling. After collection, the filters were left overnight in the same dry-box and reweighed (to obtain the total suspended particulate value). This ensures that all samples are brought to the same degree of dryness, thus avoiding weighing errors. Then the exposed Hi-Vol filters were sonicated in a Milli-Q DD water bath for 20-30 minutes to dissolve salts and to resuspend insoluble particles from the filter. Solutions were filtered (on pre-weighted 0.22 μ m diameter Nuclepore[©] polycarbonate filters). Particles on the filters were dried in a dry-box and reweighed (to obtain the mineral suspended particulate value).

Rain samples were filtered through a pre-weighed 0.22 μ m diameter Nuclepore[©] polycarbonate filter within a few hours of collection, and subdivided into two fractions, soluble and insoluble. The insoluble fraction was dried in a dry-box and re-weighed (to obtain the total particulate concentration value).

Major elements (Al, Ca, Fe, K, Mg, Na, Si) were analysed by atomic absorption spectrophotometry in insoluble aerosol particles and particles in precipitation. Trace elements (Cd, Cr, Pb, Zn) in the same samples were analysed in a graphite furnace.

Sampling method, sample treatment and analytical methods are described in detail in Guerzoni et al. (1993; 1997) and Molinaroli et al. (1993).

Back-trajectories from the sampling point were computed from analysed wind data along constant pressure surfaces, to help identify the potential source areas of the dust load. Trajectories were provided by NOAA (Harris, 1982).

STATISTICAL STRATEGY

The following steps summarise the strategy employed in the first study:

- 1. data were ranked to obtain a comparable unit of the variables, for representation of outliers and because the data were closed;
- 2. cluster analysis in *q-mode* (results cluster objects on the basis of values of variables) was performed using Ward's hierarchical agglomerative method and the squared Euclidean distance measure;
- 3. on the basis of the results of clustering, a number of sample groups was derived by applying a similarity level;
- 4. with this *a priori* criterion, linear discriminant analysis was employed to characterise sample groups in relation to variables with the greatest discriminating power. In the second study:

- 1. data were ranked as previously described;
- 2. factor analysis was used to reduce a large number of variables to a few uncorrelated variables (factors);
- 3. cluster analysis in *q-mode* was performed on significant factors (Ward's hierarchical agglomerative method and the squared Euclidean distance measure);
- 4. on the basis of the results of clustering linear discriminant analysis was employed.

This computational and statistical strategy allowed us to proceed gradually from simple data analysis to more sophisticated pattern recognition. In particular, all calculations in the analytical and multivariate statistics were based only on ranked data: each value was replaced by a number giving its place in the sequence from highest to lowest or vice versa (Swan and Sandilands, 1995). Ranked data allows application of methods with fewer assumptions about the underlying distribution (Lebart et al. 1984; Korhonen and Siljamäki, 1998). Rank transformation is also a suitable method for representing outliers and whenever standardisation is needed.

RESULTS AND DISCUSSION

Measurements in air and rain

Table 1 lists arithmetic and geometric means for insoluble fractions in aerosol and rain. The overall means are $6 \mu g m^{-3}$ and $26 m g l^{-1}$, with significant differences between samples collected from winds from the south, called Saharan, and samples from all other directions, called Background. This classification was also carried out prior to chemical analysis, using a suite of meteorological data, together with backtrajectories. Table 2 shows the wind component of particle concentrations in Saharan and Background aerosols. It may be seen that maximum load concentrations are related to S-SE and NW winds. Other authors have shown that southern winds in the Mediterranean carry great quantities of desert-derived particles (Chester et al., 1993b; Loÿe-Pilot and Martin, 1996). Dust transport across the northern boundaries of the Sahara into the Western Mediterranean and Europe is approximately 12% of the total, i.e. ~100 million tons. These estimates are subject to considerable uncertainties in relation to large-scale meteorological features (dry years, ITCZ latitudinal position) which give rise to strong interannual variability. Direct measurements (mainly bulk deposition; Loÿe-Pilot et al., 1995; Le Bolloch and Guerzoni, 1995) and historical inventories of occurrence of dust rain or dust haze in Europe (e.g. Bücher, 1989) indicate maximum fallout in spring and autumn.

Chemical data

Table 3 lists averages and geometric means of the insoluble fraction in aerosols and precipitation. The content of particulate concentration in rain is similar to that of aerosols for the major elements, and frequently lower for trace elements. The distributions of most elements are log-normal. These data have already been compared with other data from coastal areas in the Mediterranean (Corsica, Blanes, Tour du Valat, Cap Ferrat) (Guerzoni et al., 1996). The result is an increase in trace metal concentrations westwards along Mediterranean coastal areas.

First statistical treatment

Treatment was applied to the chemical data of the insoluble fraction, from aerosols and precipitation together. CA was applied to the ranked data, and results are shown in Fig. 2. Objects grouped together on numerical grounds form a cluster of points in the multivariate space spanned by the elements. Five groups emerge using the similarity value (Dlink/Dmax cut-off of 5). The main result of this kind of representation is that, based on their similarity coefficient, all samples fall in five groups, identifying aerosol depositional mode. This classification distinguishes rain samples (groups 2 and, to a certain extent, 5) from others, but mixes Saharan samples with other Local aerosols (group 1) and Mediterranean Background samples with ones from Local mining areas (groups 3 and 4). DA was applied to the 5 groups plotting in the space of the first two discriminant functions (Fig. 3). DA is used to distinguish statistically between two or more predefined groups of samples on the basis of multiple variables. It contains tests for establishing the rate of success for discriminating variables when they are combined into a discriminating function, and also criteria for controlling stepwise selection of variables according to their discriminating power. DA proceeds by calculating linear discriminant functions: linear combinations of the original variables which maximise the differences between the groups

$$F_k = a_1 V_1 + a_2 V_2 \dots$$

where V_1 are the elements and a_1 are the discriminant coefficients, the value of F for an object is called its discriminant score. These computations are described by Nie et al. (1975) and Rock (1988).

The elements with greater discriminating power were: (a) Pb for function 1, separating groups 2 (rain) from 3 and 4 (Background pollution and Local mining areas); (b) Al and Fe for function 2, separating groups 1 (Saharan and Local) from 5 (Rain and mixed aerosol). Inspection of the various aerosol groups with independent characteristics, i.e., grain-size, mineralogy, wind speed and direction, confirmed the unsatisfactory grouping of this classification.

Second statistical treatment

The second treatment was applied to the same ranked data set, using Q-FA as a first step. Q-Factor analysis transforms the data matrix using cos theta as similarity method, generating a matrix of m x m objects data. This technique applied to our data set was also useful in focusing better on the principal assemblages as source markers. Table 4 shows the factor score matrix, with indications of the percentage of information explained by each factor. It may be seen that approximately 10% of the information is lost in the next few steps. Three factors explain 90.5% of the variability and the first two factors represent 72.4% of total variability. Factor 1 (36.4%) represents crust-enriched aerosols, partly regional background and mostly Saharan dusts. Factor 2 is mainly representative of Background anthropogenic aerosols, mixed with Pb-enriched mining tails. Factor 3 typically has the signature of Local mining sources. Since some of the trace elements are common to Factors 2 and 3 (Pb, Cd and Zn in particular), Q-FA was not able to distinguish Local from distant sources. Back-trajectories (Harris, 1982) of samples closest to each group are shown in Fig. 4 and were used to trace possible sources.

The next step was to apply CA to the factor loading matrix. Results are shown in Fig 5. The procedure revealed the "highest" variability part of the data set, omitting the less

significant part (which may be defined as "background noise" and was therefore not particularly significant for the data set). In this case too, five groups were extracted by applying the same cut-off level. This classification was able to distinguish: (i) crustenriched aerosols, due to regional background plus Saharan dusts (group 2, corresponding to Factor 1 of Q-FA); (ii) samples of Background anthropogenic aerosols, mixed with Pbenriched mining tails (groups 3 and 4, corresponding to Factor 2). Instead, CA combined samples with the signature of Local mining sources (Factor 3) with other Local aerosols and Mediterranean Background (groups 5 and 1).

The five groups of samples obtained by clustering were used to perform DA. Results are shown in Fig. 6, in which discriminant scores are scatter-plotted for the two discriminant functions. Elements with greater discriminating power are: (i) Pb, Cd and Zn for function 1, separating especially group 2 (Saharan and rain) from 3 (Local mining areas), slightly from 4 (Background pollution), 5 (Local) and 1 (Mixed); (ii) Na for function 2, which clearly separates group 4 (Background pollution) from 5 (Local) and slightly from all the others. All discriminating elements therefore tend to discern Sardinian contributions from long-range ones.

In this case too, the significance of the five groups was checked with independent information, i.e., geological signature. In particular, Saharan dust was separated from other aerosols and put together with rain, most of them crust-enriched. Local inputs were included in group 3, from mining areas (old sphalerite and galena mining tailings, located about 100-200 km from the sampling site) and group 5 (Local outcrops). Group 4 represented long-range Background air pollution associated with N-NW trajectories. Group 1 contained mixed aerosols, in which the long sampling time (72 hours) favoured a complex mixture from different sources. Granulometric, mineralogical and meteorological data supported this interpretation. Grain-size analysis highlighted the well-known bimodality distribution of Saharan aerosols, combined with the typical mineralogical markers of desert provenance (i.e., palygorskite, kaolinite, rounded quartz, calcite) as well as wind direction (E-SE) and the highest mass loadings. As an example, Fig. 7 shows some cumulative volume grain-size distributions of Saharan aerosol events, two different episodes (in autumn and spring) showing distinct grain-size distributions (Molinaroli and De Falco, 1995). Samples were also examined by scanning electron microscopy (SEM); Fig. 8 shows three SEM photographs of typical mineral markers observed in Saharan and Local aerosols.

Wind speed (high: 15-18 m/s) and wind direction (west), together with very fine grain sizes (1-2 μ m) and mineralogical composition (talc, amphibole, Na-plagioclase, etc.) supported Local and regional inputs for groups 3 and 5.

CONCLUSIONS

Concurrent sampling of aerosol and precipitation is useful in comparing contents of the insoluble fraction of air particulate.

The statistical methods used in this study are well known, and commonly used in situations in which the interplay of many sources of variation do not allow an objective and comprehensive method of describing physical phenomena. Divergent results are obtained by applying two different combinations of classification and statistical methods to geochemical data.

In the first statistical treatment, aerosols and precipitation particles were better distinguished, but little information could be inferred for separating Local from distant sources.

In the second statistical treatment, the application of CA subsequent to Q-FA better distinguished among Saharan dust, remote Background pollution and Local aerosol sources, but some unexplained mixed aerosols still resulted. Subsequent DA better discriminated Sardinian contributions from long-range ones; moreover, crust-enriched precipitation was classified together with Saharan dust, indicating that they are similar to desert-derived aerosols.

Granulometric, mineralogical and meteorological data confirmed the results of the statistical methods employed in the second treatment.

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FIGURE CAPTIONS

- Fig. 1 Mediterranean basin and location of sampling station.
- Fig. 2 Dendrogram produced by clustering 69 samples. Cluster analysis was performed on ranked data matrix using Ward's hierarchical agglomerative method and squared Euclidean distance measure. Numbers (1-5) indicate clusters identified by the classification.
- Fig. 3 Discriminant score scatterplot for two discriminant functions: classification of five groups of samples obtained by cluster analysis (see Fig. 2). Discriminant variables are: Pb for Function 1; Al and Fe for Function 2.
- Fig. 4 Back-trajectories computed at 850 hPa level. Symbols mark every 12 hours along wind path.
- Fig. 5 Dendrogram produced by clustering 69 samples. Cluster analysis was performed on Q-FA scores using Ward's hierarchical agglomerative method and squared Euclidean distance measure.
- Fig. 6 Discriminant score scatterplot for three discriminant functions: classification of sample groups obtained by cluster analysis (see Fig. 5). Discriminant variables are Pb, Cd, Zn for Function 1; Na for Function 2.
- Fig. 7 Cumulative volume grain-size distribution of Saharan dust transported over Sardinia during autumn 1990 (solid line) and apring 1991 (broken dashed line).
- Fig. 8 Scanning electron micrographs of marker minerals. (a) Perfectly rounded palygorskite grain with felt-like surface texture, of desert type. (b) Talc grain typical of Local mining area. (c) Na-rich plagioclase grain typical of Local outcrops.











FIG. 3



FIG. 4













FIG. 8

	Saharan	Background	All
Aerosol	(n=10)	(n=45)	(n=55)
$(\mu g m^{-3})$	21 (17)	3 (2)	6 (4)
Rain	(n=12)	(n=19)	(n=31)
$(mg l^{-1})$	54 (12)	8 (3)	26 (18)

Table 1. Insoluble fractions in Saharan and Background aerosol and precipitation. Arithmetic mean (Geometric mean).

 Table 2. Wind component of particle concentration in air for Saharan and Background aerosols.

Wind	Saharan	Background
direction	(n = 10)	(n = 45)
	$\mu g m^{-3}$	$\mu g m^{-3}$
NE	0.1	0.1
E	3.2	0.4
SE	10.0	0.5
S	2.3	0.1
SW	1.7	0.1
W	1.4	0.3
NW	2.2	1.4
Ν	0.1	0.2
Total	21.0	3.1

Table 3. Chemical data of insoluble fraction of aerosols and precipitation. Averages with geometric means in brackets.

Element	Aerosol	Rain	Aerosol
	(%)	(%)	$(\mu g m^{-3})$
Si	16.5 (14.0)	17.5 (15.6)	1.182 (0.459)
Al	5.5 (4.4)	5.2 (3.9)	0.526 (0.143)
Ca	0.87 (0.51)	1.1 (0.7)	0.110 (0.017)
Na	5.0 (2.4)	2.0 (0.9)	0.241 (0.080)
Κ	2.1 (1.0)	1.8 (1.5)	0.095 (0.034)
Mg	1.0 (0.8)	1.3 (1.1.)	0.092 (0.026)
Fe	2.9 (2.6)	3.1 (2.8)	0.228 (0.085)
	$(mg kg^{-1})$	$(mg kg^{-1})$	$(\mu g m^{-3})$
Cd	6.5 (4.4)	1.4 (1.2)	0.000026 (0.016)
Zn	1216 (1045)	552 (444)	0.0066 (0.0034)
Pb	3324 (2213)	191 (153)	0.0112 (0.0073)
Cr	82 (65)	32 (27)	0.0006 (0.0003)

Variable	Fact. 1	Fact. 2	Fact. 3
Si	1.097*	0.44	0.00
Al	1.145*	0.11	0.41
Ca	1.162*	0.46	-0.08
Na	0.34	2.077*	-1.15
Κ	1.626*	-0.39	0.51
Mg	1.591*	-0.31	0.42
Fe	1.118*	0.20	0.32
Cd	-0.41	0.75	1.807*
Zn	-0.33	0.32	2.303*
Pb	-0.52	1.740*	0.56
Cr	0.24	1.514*	-0.28
% of variance	e explained by e	each factor	
-	36.4	36.0	18.1
Cumulative %	6 variance		
	36.4	72.4	90.5

Table 4. Q-Mode Factor score matrix. Elements with asterisks are those characterising each factor.