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Iron Speciation in Different Saharan Dust Advections and Effect of the Procedural Blank on the Results From X-ray Absorption Spectroscopy and Selective Leaching Experiments

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Abstract: In this work, we applied X-ray Absorption Spectroscopy (XAS) and selective leaching experiments for investigating iron speciation in different dust advections collected on different unwashed quartz fiber filters. XAS analysis evidenced a predominance of Fe(III) in 6-fold coordination for Saharan dust and a trend towards Fe(II) and 4-fold coordination in the order: Saharan dust, mixed Saharan, and non-Saharan aerosol samples. The role of the sampling substrate was evaluated explicitly, including in the analysis a set of blank filters. We were able to pinpoint the possible contribution to the overall XAS spectrum of the residual Fe on quartz as the concentration decrease towards the blank value. In particular, the filter substrate showed a negligible effect on the structural trend mentioned above. Furthermore, selective leaching experiments evidenced a predominance of the residual fraction on Fe speciation and indicated the lowest Fe concentrations for which the blank contribution is <20% are 1 µg for the first three steps of the procedure (releasing the acid-labile, reducible and oxidizable phases, respectively) and 10 µg for the last step (dissolving the insoluble residuals).

Keywords: iron speciation; Saharan dust; filter blank; XANES; EXAFS; selective leaching experiments; Monte Martano (Central Italy)

1. Introduction

The Sahara desert is a significant dust source, emitting 50% of global mineral dust [1]. Mineral dust is rich in iron compounds, and therefore it plays an essential role in the iron cycle worldwide. In fact, mineral dust sources supplies approximately 95% of the globally averaged iron budget [2,3] and North African deserts are the main emitting areas. Due to their proximity and to the preferred transport routes, Saharan dust is deposited in the Mediterranean basin [4] and in the North Atlantic ocean [5,6]. Oceans' iron fertilization promotes phytoplankton activity impacting on the sulfur cycle via increased DMS (dimethyl sulfoxide) production [7] and on the CO_2 cycle by enhancing its sequestration from the



atmosphere [8]. It is worth noting that dust ability to impact on phytoplankton activity is mainly linked to iron solubility and, in turn, to its bioaccessibility (see [9] for definition). Although being the most abundant transition metal in atmospheric aerosols [10], iron solubility in aerosols is highly variable (0.05 to 80% [11]) and depends not only on the aerosol source but also on atmospheric and cloud processing [12]. In fact, complex atmospheric processing including reductive processes, acid dissolution and photochemistry, can take place during transport. Longo et al. [13] found a correlation between the oxidation state and the estimated travel time, suggesting the extent of atmospheric processing is proportional to the transport time. Photo-reductive processes are mainly active at the aerosol particles' surface [14], however, these processes are reversible and produce a small effect on the overall iron chemical speciation [10,15]. Cloud processing, on the contrary, can lead to a significant increase in the soluble Fe(II) proportion due to the pH of cloud water [16,17]. The majority of iron in mineral dust is in the Fe(III) oxidation state ([10,18]), in the form of iron oxides like hematite and goethite [19] and accounting for 38% to 72% of total elemental iron by mass in Saharan dust [20]. Despite the lower solubility of Fe(III) compared to Fe(II), Fe(III) is known to contribute largely to the water soluble iron fraction [21]. Additionally to its impact on climate, soluble iron in fine aerosols can take part in reactions generating reactive oxygen species, therefore being identified as a harmful component of aerosol to human health. This characteristic could be relevant to the relationship between high particulate matter concentrations recorded in Europe during severe Saharan dust advections and mortality [22]. Iron speciation can be determined using a variety of techniques, as summarized by Majestic et al. [21], most of them being wet chemical methods that require an extraction step, hence being destructive and time-consuming. Amongst the non-destructive methods, Mössbauer spectroscopy is widely used; however, it requires high amounts of sample to be analysed [23]. X-ray absorption spectroscopy (XAS) allows for direct determination of metals speciation on smaller samples like aerosol filters. Formenti et al. [20] applied XAS to western Sahara mineral dust, Longo et al. [13] applied a combined XANES and wet chemical techniques approach for studying the effect of atmospheric processing on iron speciation after long-range transport and Petroselli et al. [24] compared iron speciation in Saharan dust and steel production emissions. Recently, McDaniel et al. [25] applied XANES for quantifying Fe(II) in aerosol sampled shipboard in the Atlantic Ocean and investigated the differences between Saharan dust and other provenance samples. XAS is non-destructive and the sampling filter substrate can be analysed directly without any sample preparation step. However, filter substrates can contain high levels of impurities that can affect the sample measurement. Iron, in particular, is present in filter substrates in non-negligible concentrations. Previous literature studies reported XAS data on Nuclepore [26] Teflon ([13,21,27] and cellulose ester filters [25]. Moreover, acid-cleaning procedures are usually implemented on filters to be used for XAS analysis in order to minimize the impact of the substrate's trace metals contamination on the analysis [21]. However, filter acid cleaning procedures are expensive and time-consuming and cannot be routinely applied to all the filters used for automated aerosol sampling in air quality monitoring sites. In order to investigate iron speciation in episodic events such as Saharan dust advections on routinely collected aerosol samples, the filter impurities contribution to the XAS spectrum needs to be assessed. The aim of this work is to perform XAS analyses and selective leaching experiments on samples collected during different long-range dust advections on different unwashed quartz fiber filter substrates. To this aim, iron speciation in two severe Saharan dust advections, two mixed Saharan dust advections and two non-Saharan samples will be compared. The filter contamination will be quantitatively evaluated and the effect of the blank filter on the XAS spectrum will also be investigated.

2. Results

2.1. Aerosol Samples Characterization

In this work, samples from a severe Saharan dust outbreak (SH_Dec), two mixed-Saharan advections (mixSH_Feb and mixSH_Apr) and two different, non-Saharan, air masses (nonSH_Oct and

nonSH_Jan) are considered. Aerosol mass (PM_{10} and $PM_{2.5}$) values are reported in Table 1. PM values temporal evolution for the five advections and the corresponding air masses back-trajectories (BTs) are reported in the Supplementary Material, Figures S1 and S2.

Table 1. Samples description: sampling date, filter code, aerosol mass concentration (PM_{10} and $PM_{2.5}$ in $\mu g m^{-3}$) and iron concentrations determined by ICP-OES for HVS (high volume sampler) PM_{10} and SWAM $PM_{2.5}$ samples expressed in $\mu g m^{-3}$ and converted to atoms cm⁻². Errors on the last digit are indicated in parentheses.

Date	Code	PM ₁₀	PM _{2.5}	PM _{2.5} /PM ₁₀	$Fe[\mu gm^{-3}]$	Fe [atoms cm ⁻²]
30 November 2014	SH_Dec_PM ₁₀	83.9		0.38	2.36(1)	$7.64(2) imes 10^{16}$
	SH_Dec_PM _{2.5}		32.3	0.38	0.39(2)	$3.2(1) \times 10^{16}$
01 December 2014	SH_Dec_PM ₁₀	86.9		0.36	7.81(1)	$2.564(3) imes 10^{17}$
	SH_Dec_PM _{2.5}		31.2	0.36	0.39(1)	$3.30(5) imes 10^{16}$
19 February 2014	mix-SH_Feb_PM ₁₀	18.9	7.7	0.41	1.174(1)	$2.701(3) \times 10^{16}$
23 April 2014	mix-SH_Apr_PM ₁₀	12.4	8.5	0.68	0.990(1)	$1.622(2) \times 10^{16}$
31 October 2014	non-SH_Oct_PM _{2.5}	18.9	16.9	0.89	0.099(3)	$8.2(2) \times 10^{15}$
07 January 2015	non-SH_Jan_PM _{2.5}	22.0	18.5	0.84	0.071(1)	$5.91(4) imes 10^{15}$
blank	blank_SWAM	-	-	-	0.026(4)	$2.2(3) \times 10^{15}$
blank	blank_HVS	-	-	-	0.055(1)	$1.44(3) \times 10^{15}$

The pure Saharan dust outbreak was sampled on 30 November and 1 December 2014. The transport came from Algeria and Tunisia, with a contribution from Libya on 30 November only. Both PM_{10} and $PM_{2.5}$ values recorded during the advection were extremely high compared to the annual mean value for the year 2014 (10 µg m⁻³ and 7 µg m⁻³, respectively). The calculated dust load [28], resulted in 80.0 µg m⁻³ and 83.0 µg m⁻³, accounting for 95% and 96% of the total PM_{10} , and 29.7 µg m⁻³ and 28.6 µg m⁻³, accounting for 92% of the total $PM_{2.5}$ on both days.

The two mixed-Saharan dust events occurred in late winter and early spring. On 19 February 2014 (mix-SH_Feb), Saharan dust from Libya mixed with a significant contribution from the Mediterranean sea was detected (Figure S2c). PM_{10} and $PM_{2.5}$ concentrations were 18.9 µg m⁻³ and 7.7 µg m⁻³, respectively and the dust load was 14.9 µg m⁻³, accounting for 75% of the total PM_{10} . The dust sampled on 23 April 2014 (mix-SH_Apr) was mostly coming from the Iberian Peninsula with a minor contribution from Northern Africa carried by high altitude air masses (Figure S2d). PM_{10} and $PM_{2.5}$ values read 12.4 µg m⁻³ and 8.5 µg m⁻³, respectively, and the dust load resulted in 3.0 µg m⁻³, accounting for 24% of the total PM_{10} mass concentration. The different $PM_{2.5}/PM_{10}$ ratios, together with the different dust contributions, reflect the different extent of the Saharan dust contribution in the two samples. Since high dust load and low $PM_{2.5}/PM_{10}$ ratios are typical of Saharan dust advections, the contribution of Saharan dust to mix-SH_Feb resulted higher than in mix-SH_Apr.

The first non-Saharan advection occurred on 31 October 2014 (non-SH_Oct). The sampled air mass came from Eastern Europe and the Balkans where it passed over a very polluted industrialized area in proximity of ground level (Figure S2e). Anthropogenic sources would explain the high PM air mass concentration (18.9 μ g m⁻³) and the extremely high PM_{2.5}/PM₁₀ ratio (0.9) of the sample.

The second non-Saharan advection occurred on 7 January 2015 (non-SH_Jan) and came from Northern Europe. In this case, the air masses spent most of the time in the high troposphere at a great distance from continental Europe (Figure S2f). The long-range advection combined with a regional contribution by anthropogenic aerosols could justify the high $PM_{2.5}$ mass concentrations (18.9 µg m⁻³) and the high $PM_{2.5}/PM_{10}$ ratio (0.8) of this sample.

Total iron concentrations for all the samples and blank filters, are reported in Table 1. Amongst the HVS (high volume sampler) PM_{10} samples, the highest Fe concentrations are recorded for the pure Saharan dust outbreak, reading 2.36 µg m⁻³ and 7.81 µg m⁻³ for 30 November and 1 December 2014, respectively. Lower Fe concentrations were recorded for the mixed Saharan advections, i.e., 1.17 µg m⁻³ for mix-SH_Feb and 0.99 µg m⁻³ for mix-SH_Apr. The high iron concentrations recorded during the Saharan dust advections, together with the high PM_{10} mass concentrations and calculated dust loads, suggest the Saharan dust contribution to Fe concentration to be dominating in these samples series. As regards the $PM_{2.5}$ samples series, Fe concentrations in Saharan dust samples (0.39 µg m⁻³) are about five times higher than in non-Saharan samples (0.099 µg m⁻³ for non-SH_Oct and 0.071 µg m⁻³ for non-SH_Jan). This evidence suggests that the finer particle size fraction plays a role in transporting iron in the atmosphere alongside the coarser one which is typically associated with mineral dust emissions.

2.3. Iron Speciation

2.3.1. X-ray Absorption Spectroscopy

Figure 1 shows the XANES spectra for PM₁₀ (Figure 1a) and PM_{2.5} samples (Figure 1b), respectively.



Figure 1. Lefthand panels show the XANES spectra of (**a**) Saharan dust PM_{10} samples (SH_Dec_3011 and SH_Dec_0112) and mixed Saharan PM_{10} samples (mix-SH_Feb and mix-SH_Apr); (**b**) Saharan dust $PM_{2.5}$ samples (SH_Dec_3011 and SH_Dec_0112), non Saharan $PM_{2.5}$ samples (non-SH_Oct and non-SH_Jan) and SWAM blank filter. A blow-up of the pre-edge features of the same samples is reported in the corresponding righthand panels. Herein the open circles are the experimental points and the lines the model best fits.

The edge energies for the four samples collected during the SH_Dec advection are nearly coincident at 7125.8 eV, compatible with Fe(III). For the mixed-Saharan and the non-Saharan samples, we can observe a gradual shift towards lower energies (from 7124.2 eV to 7122.2 eV), in an order that reflects the decrease in Saharan dust contribution, aerosol loading and iron concentration (mix-SH_Feb>mix-SH_Apr>non-SH_Oct>non-SH_Jan). Moreover, all the spectra of the Saharan and mixed-Saharan samples result similar, with a peculiar feature around 7149 eV, which is not present in the non-Saharan samples. This feature is more pronounced in more ordinate Fe(III) compounds such as hematite, rather than in more disordered compounds like ferrihydrite.

The pre-edge features were fitted in all the samples, and the results are reported in Table 2. The centroid position ranges between 7115.04 eV in Saharan dust samples (HVS PM_{10} for SH_Dec-3011) to 7113.93 eV in the non-SH_Jan sample.

Sample (PM ₁₀)	Centroid (±0.2)	Intensity (±0.02)
SH_Dec-3011	7114.9	0.05
SH_Dec-0112	7115.0	0.05
mix-SH_Feb	7114.7	0.11
mix-SH_Apr	7114.5	0.17
Sample (PM _{2.5})	Centroid (±0.2)	Intensity (±0.02)
SH_Dec-3011	7115.0	0.05
SH_Dec-3011 SH_Dec-0112	7115.0 7114.9	0.05 0.05
SH_Dec-3011 SH_Dec-0112 non-SH_Oct	7115.0 7114.9 7113.9	0.05 0.05 0.16
SH_Dec-3011 SH_Dec-0112 non-SH_Oct non-SH_Jan	7115.0 7114.9 7113.9 7114.4	0.05 0.05 0.16 0.20

Table 2. XANES pre-edge features fit results for HVS PM_{10} (upper panel) and $PM_{2.5}$ (lower panel) samples.

A comparison of the pre-edge parameters with literature reference data ([29], rescaled according to [30]) is shown in Figure 2. A gradual shift is observed, from high-energies/low-intensities for the pure Saharan dust samples (a–d) towards lower-energies/higher-intensities for non-Saharan samples. Moreover, the coordination geometry shows a decreasing trend from 6-coordinated in pure Saharan dust and mix-SH_Feb samples to lower coordination (5 and 4) for the other mix-SH and non-SH samples suggesting a less symmetrical Fe(III) coordination geometry with decreasing aerosol loading.



Figure 2. XANES pre-edge features fit parameters scatterplot: comparison between samples and standard Fe(II) and Fe(III) compounds studied in previous literature ([29,30]). Legend for SH_Dec 2014 samples: (a) HVS PM₁₀ 30.11.2014, (b) HVS PM₁₀ 01.12.2014, (c) SWAM PM_{2.5} 30.11.2014, (d) SWAM PM_{2.5} 01.12.2014

Regarding EXAFS analyses, Figure 3 shows the k^2 -weighted EXAFS spectra in their relevant fit window and the relative fitted Fourier transforms for HVS PM_{10} samples (Figure 3a), $PM_{2.5}$ and the SWAM blank filter (Figure 3b).



Figure 3. Lefthand panels show EXAFS spectra of (**a**) Saharan dust PM_{10} samples (SH_Dec_3011 and SH_Dec_0112) and mixed Saharan PM_{10} samples (mix-SH_Feb and mix-SH_Apr); (**b**) Saharan dust $PM_{2.5}$ samples (SH_Dec_3011 and SH_Dec_0112), non Saharan $PM_{2.5}$ samples (non-SH_Oct and non-SH_Jan) and SWAM blank filter. Fourier transformed spectra fits are reported in the corresponding righthand panels. Herein the open circles are the experimental points and the lines the model best fits (see text).

The EXAFS spectrum was reproduced with a two-shell ferrihydrite model, and the fit parameters are shown in Table 3. The coordination number (N) was strongly correlated with the Debye–Waller factors, and was thus obtained by a different first shell fit, linking N and r(Fe-O) using the results of Bond Valence Method calculations ([31,32]).

For the HVS PM_{10} samples, we observe a decrease in the Fe-O bond distance from 1.98 Å for SH_Dec to 1.95 Å for mix-SH_Apr, which agrees with those calculated using the bond valence method (BVM, [32]) for 6 and 5-coordinated Fe(III) (Table S1), and coordination number shows a similar decrease (from 5.5 to 5.2). A similar decreasing trend in both Fe-O bond distances (1.98 Å to 1.92 Å) and coordination number (5.5 to 4.9) are observed for $PM_{2.5}$ samples of SH_Dec to non-SH_Jan. Therefore, the overall trend follows the order: SH_Dec>mix-SH_Feb>mix-SH_Apr>non-SH_Oct>non-SH_Jan. It is worth noting that both SWAM $PM_{2.5}$ and HVS PM_{10} samples from SH_Dec show the same spectral

features and fit parameters. We can, therefore, exclude any influence of the sampling system (sampler and substrate) for this event, which is, however, the most intense amongst the presented ones.

Table 3. EXAFS data analysis results. Bond distances (r), coordination numbers (N) and amplitudes

 Sample Fe-O Fe-Fe

 r [Å]
 N
 σ^2 [Ų]
 r [Å]
 N
 σ^2 [Ų]

PM ₁₀						
SH_Dec-3011	1.98 ± 0.01	5.4 ± 0.5	0.009 ± 0.005	2.96 ± 0.02	0.8 ± 0.5	0.02 ± 0.02
SH_Dec-0112	1.98 ± 0.01	5.5 ± 0.4	0.009 ± 0.003	2.95 ± 0.01	0.8 ± 0.3	0.01 ± 0.01
mix-SH_Feb	1.96 ± 0.01	5.2 ± 0.4	0.010 ± 0.004	2.93 ± 0.01	0.6 ± 0.3	0.02 ± 0.02
mix-SH_Apr	1.95 ± 0.01	5.2 ± 0.2	0.006 ± 0.005	3.03 ± 0.01	2.2 ± 0.9	0.02 ± 0.02
PM _{2.5}						
SH_Dec-3011	1.98 ± 0.01	5.4 ± 0.3	0.009 ± 0.003	2.95 ± 0.01	0.9 ± 0.3	0.01 ± 0.01
SH_Dec-0112	1.98 ± 0.01	5.5 ± 0.6	0.010 ± 0.005	2.95 ± 0.02	0.8 ± 0.5	0.02 ± 0.02
non-SH_Oct	1.93 ± 0.01	4.8 ± 0.3	0.009 ± 0.004	2.94 ± 0.01	0.9 ± 0.3	0.01 ± 0.01
non-SH_Jan	1.92 ± 0.01	4.9 ± 0.1	0.004 ± 0.002	2.98 ± 0.01	4.8 ± 0.6	0.02 ± 0.01
SWAM-blank	1.84 ± 0.01	4.4 ± 0.1	0.008 ± 0.004	2.84 ± 0.01	1.1 ± 0.1	0.01 ± 0.01

The second shell Fe-Fe distance is about 2.95 Å in the majority of the samples. The slight variations observed between the samples can be ascribed to structural defects and partial substitution of Fe(II) and Fe(III) on the same site, while the much shorter distance in the blank filter is in line with a different coordination geometry.

In order to get an insight on blank filter contribution to the XAS spectrum, a SWAM quartz fiber filter was analysed. Despite the spectrum of a blank HVS filter is not available, we assume the iron local structure to be similar in both cases because the filter material and manufacturer are the same. The XANES spectrum of the blank filter is shown in Figure 1b and shows an edge energy of 7121.5 eV, compatible with Fe(II). The EXAFS fit of the blank filter (Figure 3b, Table 3) shows a Fe-O bond distance of 1.84 Å, compatible with 4-coordinated Fe(III) (1.87 Å, [32], Table S1), and a 4.4 coordination. Considering that XANES data are not quantitative, they are in good agreement with EXAFS data. In fact, in the blank filter iron results low coordinated as expected for being an impurity ion in the quartz matrix.

2.3.2. Selective Leaching Experiments

Leaching test results are reported in Table 4. Fe concentrations in non-SH_Oct and non-SH_Jan samples were too low to obtain reliable Fe fractionation data after blank subtraction, and they are hence excluded from further analyses.

Date	Sample	I (%)	II (%)	III (%)	IV (%)	tot Fe (µg)
24 April 2014	mix-SH_Apr_PM ₁₀	2	5	23	69	2.0
19 February 2014	mix-SH_Feb_PM ₁₀	10	11	8	70	11.1
30 November 2014	SH_Dec_PM ₁₀	14	6	8	72	29.6
01 December 2014	SH_Dec_PM ₁₀	12	7	8	73	100.6
30 November 2014	SH_Dec_PM _{2.5}	16	7	8	69	9.6
01 December 2014	SH_Dec_PM _{2.5}	18	8	13	61	11.0

Table 4. Leaching test results after blank subtraction.

Legend: I step: acid-labile fraction, II step: reducible Fe oxides and hydroxides, III step: oxidizable phase, IV step: insoluble residuals.

Saharan dust HVS PM_{10} samples collected on 30 November and 1 December 2014 show a consistent fractionation, with the iron oxides fraction (step II) accounting for 7% of total iron. In the mix-SH_Feb sample, fraction II is enriched to 11% of total Fe, while the mix-SH_Apr sample shows

lower percentages in the acid-labile and the reducible fractions (step I and II) and higher values in the oxidizable phase, which accounts for 23% of total Fe.

Comparing PM_{10} (i.e., HVS samples) and $PM_{2.5}$ samples for the SH_Dec advection event, the iron fractionation for the two different size fractions is consistent on 30 November 2014, but different in the 1 December case. In fact, the $PM_{2.5}$ fraction shows slightly higher percentages in step I and III and lower residual (step IV) with respect to the coarse fraction.

The blank contribution in the sequential extraction solutions is evaluated for ten atmospheric aerosol samples (listed in Supplementary Material, Table S2) and reported as a function of the total Fe amount (Figure 4). The influence of the procedural blank on the first three steps of the procedure is quite similar and increases as the total Fe amount decreases. The fourth step is characterized by higher blank values than the previous ones, thus affecting the analytical results more severely. We estimate that the reagents account for about 30% of the total blank values, whereas the remaining 70% is due to the filter contribution. Finally, Figure 4 shows that the lowest Fe content for which the blank contribution is <20%, which can be regarded as a good target, resulted to be 1 μ g for the steps I-II-III and 10 μ g for step IV.



Figure 4. Blank contribution on the results from selective leaching experiments as a function of the Fe amount.

3. Discussion

In this work, iron speciation is investigated in different dust advections collected at the Monte Martano site, Central Italy, on different unwashed quartz fiber filters. Five advection events are considered, namely one pure Saharan dust advection, two mixed Saharan advections and two non-Saharan advections. The aerosol mass concentrations and the total iron concentrations decrease from pure Saharan dust to non-Saharan samples reflecting the decrease in Saharan dust contribution to the sampled aerosol. The iron concentrations observed for the Saharan and mix-Saharan dust samples are comparable to the ones recorded by Mc Daniel et al. [25], who reported a range between 1547 ng m⁻³ and 2491 ng m⁻³ for Saharan dust collected in the Atlantic Ocean. Mixed Saharan dust advections results are also within the range determined by Longo et al. [13] for long-range transported Saharan dust sampled in Greece and in the Atlantic Ocean (735–1580 ng m⁻³). These observations confirm Monte Martano is a representative site for studying long-range Saharan dust advections in the Central Mediterranean Basin.

3.1. Iron Speciation in the Different Advections

Iron speciation is investigated by both X-ray Absorption Spectroscopy and selective leaching experiments. XANES results evidence Fe(III) is predominant in all the Saharan dust samples, which spectra show common features such as the same edge energy (7125.8 eV), similar spectral shape and pre-edge features parameters. These results are in accordance with the results from McDaniel et al. [25],

who observed nearly coincident spectra for Saharan dust samples, with an edge energy of 7126 eV. Moreover, the shape of the pre-edge features in Saharan dust samples is compatible with the work by Formenti et al. [20], who observed a double peak structure with centroid positioned between 7114 eV and 7116 eV as a signature for iron oxides in Saharan dust. However, the pre-edge energies obtained for Saharan dust are slightly higher than what reported by Longo et al. [13], i.e., pre-edge centroid between 7114.4 eV and 7113.6 eV for dust sampled in Greece, suggesting a less effective atmospheric processing in our samples, that might be attributed to the high velocity of the air masses in the studied Saharan dust advection. Regarding non-Saharan advections, lower edge energies (7124–7122 eV) are compatible with the results of McDaniel et al. [25] for samples with North American and marine provenances. However, unlike Saharan dust which is a well defined and well studied type of aerosol, the few non-Saharan samples presented in this study do not permit a thorough characterization of aerosol transported from different sources to the Monte Martano site.

Both the XANES and the EXAFS regions of the spectrum evidence a shift from mainly Fe(III) in 6-fold coordination towards a mixture of Fe(II) and Fe(III), where the latter is still predominant, and lower coordination going from pure Saharan dust to non-Saharan samples in the order: SH_Dec>mix-SH_Feb>mix-SH_Apr>non-SH_Oct>non-SH_Jan. This trend is in accordance with the observations on dust load and total iron concentrations, suggesting it is mainly driven by the decreasing contribution of Saharan dust to the overall dust sampled.

Selective leaching experiments evidence the dominance of the insoluble residual fraction in all the samples, while the iron oxides fraction accounts for about 7% of total iron on Saharan and mixed-Saharan samples. This percentage is much lower than what reported by Formenti et al. [20], who observed the iron oxides contribution in Saharan dust being in the range 38–72%, and apparently not in accordance with XAS results, which point towards the presence of Fe(III) oxides in determining the observed spectral features. However, it has to be considered that some iron oxides identified in Saharan dust by XAS speciation analyses (e.g., hematite, goethite, ferrihydrite) could not be completely extracted in step II, hence contributing to the residual insoluble fraction ([33]; [34]; [35]), which accounts for about 70% for all the Saharan dust samples considered in this study.

The leaching results for the mixed Saharan samples give us an insight on the mixing and ageing of the dust during its permanence in the atmosphere. The mix-SH_Apr sample, in particular, shows higher values in the oxidizable phase (step III) which suggests a higher level of atmospheric processing with respect to the other advections, compatible with the lower velocity of the air mass. This hypothesis is supported by XAS results as well, which show a higher contribution from Fe(II) in the mix-SH_Apr sample with respect to SH_Dec and mix-SH_Feb samples. Finally, the comparison between the two particle size fractions for the SH_Dec event evidence a different fractionation in the PM_{2.5} sample of 1 December 2014, with respect to the PM₁₀ collected on the same day. This evidence suggests a cleaner Saharan dust advection in the fine fraction compared to the previous day. This observation is supported by previous studies that evidenced a higher degree of contamination by anthropogenic emissions on 30 November compared to 1 December [36].

In summary, iron speciation measurements show a very clear and consistent trend which reflects the contribution of Saharan dust to the overall aerosol composition and give insights on the atmospheric processing of the samples during their long-range transport to Central Italy.

3.2. Effect of the Procedural Blank on Fe Speciation Measurements

In order to evaluate the effect of the procedural blank on iron concentrations and speciation, blank HVS and SWAM quartz fiber filters were analysed along with the samples, and their total iron concentrations are reported in Table 1. Comparing the Fe concentrations in blanks and samples, we observed that for the HVS PM_{10} samples, the blank contribution is always below 6% of the total iron concentration, due to the high aerosol loading of the filters. For the SWAM samples, on the other hand, the filter contribution results no more negligible and reaches values of 26 and 37% of the total Fe content in the two non-Saharan samples (non-SH_Jan, respectively).

The XANES and EXAFS parameters obtained for the blank filters suggest that the contribution of the blank filter becomes gradually more important in determining the XAS spectral features in the XANES region in a trend from the most loaded to the least loaded filter samples (SH_Dec>mix-SH_Feb>mix-SH_Apr>non-SH_Oct>non-SH_Jan). However, the shape of the pre-edge features for the blank filter shows two components, while the shape of the peaks for the non-SH_Jan is mono-modal. Therefore, in the case of XAS analyses, it is not possible to disentangle the effect of the increased blank contribution from the decreased Saharan dust contribution to the observed trend.

Leaching test results show that the blank values are much higher in the last step of the procedure, which dissolves the residual fraction. This can be ascribed both to the use of concentrated acids (HNO₃, HCl and HF, see experimental method), which could increase the reagent blank, and to the use of a more aggressive extracting solution, which enhances the Fe released from the filter matrix. Leaching data allowed to quantify the blank contribution to the samples, and it was possible to calculate the minimum Fe concentration for which the blank contribution is lower than 20%, which can be regarded as a good target for unbiased Fe fractionation results.

4. Materials and Methods

4.1. Sampling

In this study, we considered the most significant dust advections recorded at the rural regional background station of Monte Martano, Central Italy (MM, 1100m a.s.l.; [37]), between February 2014 and January 2015. The site is part of the WMO SDS-WAS (World Meteorological Organization Sand and Dust Storms—Warning Advisory and Assessment System, https://sds-was.aemet.es) network for Saharan dust transport monitoring since 2013 and of the EMEP (European Monitoring and Evaluation Programme, https://www.emep.int) network since 2017. Saharan dust advections are forecasted and identified on the basis of air mass back-trajectories. Back-trajectories (BTs) are calculated using Hysplit v4 [38] using NCEP-GFS (National Centers for Environmental Prediction - Global Forecast System) model data with $1 \times 1^{\circ}$ resolution, and applying four ending altitudes (0, 500, 1000 and 3000 m a.g.l.) and a 6-hours time resolution [24].

The samples were collected by means of a SWAM dual-channel low volume sampler (FAI instruments, 38.33 Lmin^{-1}) equipped with PM₁₀ and PM_{2.5} impactors and a high volume sampler (Air HVS flow, Analitica Strumenti, 1140 Lmin^{-1}) equipped with a PM₁₀ impactor. The filters used on the two samplers are quartz fiber filters, namely Whatman QM 47 mm and Whatman QMA for SWAM and HVS samplers, respectively. The sampling time for the SWAM sampler is 24 h, while the sampling interval for the high volume sampler was set between 6 and 18 h according to the duration of the intrusion. Filter samples were kept frozen until analysis, as this preservation method was proven to minimize the loss of Fe(II) due to oxidating processes [14,39]. Filters were then defrost and cut into pieces of suitable size for the different analyses described in the following paragraphs.

4.2. Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES)

Samples were acid digested using a Microwave Assisted Reaction System (MARS, CEM Corporation), using HNO₃ (Millipore Suprapur, 65%) and H₂O₂ (Carlo Erba Reagents, 30–32%) in 4:1 (v/v) proportion. Analyses were performed using an Ultima II ICP-OES (Horiba Scientific) equipped with an ultrasonic nebulizer. Iron was quantified choosing the 259.94 nm wavelength and performing a calibration with five multi-element standard solutions prepared from Précis Quality Control Standard 21 (100 mg L⁻¹, 5% HNO₃, Horiba-Jobin Yvon) in a concentration range between 0.01 and 1 ppm. The limits of detection (LOD) and of quantification (LOQ) of Fe were estimated from the calibration curve, resulting 0.04 and 0.14 mg L⁻¹, respectively.

In order to determine iron fractionation (i.e relative distribution among the various mineralogical phases), the sequential extraction method proposed by Pueyo et al. [40] and adopted by the European Community Bureau of Reference (BCR) was used. The method, originally developed for soil and sediment samples, was also applied to the sequential chemical fractionation of atmospheric aerosols, as reviewed by Smichowski et al. [41]. The BCR procedure consists of three sequential extraction steps plus a fourth step which allows to quantify Fe concentration in the residual fraction as well. The first step consists of treating the sample with acetic acid for releasing the acid-labile fraction (carbonates and labile organic matter), while in the second step hydroxylamine hydrochloride is used to release the reducible compounds (Mn and Fe oxides and hydroxides). The third step consists of a microwave-assisted digestion with hydrogen peroxide and allows to release the oxidizable phases (organic matter and sulfides). The residues from third step are then acid-digested in the microwave system using a HNO₃:HCl:HF 1:3:2 (v/v) mixture for dissolving the residual fraction (silicates and alluminosilicates). A more detailed description of the procedure used is reported in Petroselli et al. [35]. All the samples obtained from the various steps were then analysed by inductively coupled plasma atomic emission spectrometry (Vista PRO by Agilent Technologies, Santa Clara, CA, USA), using on-line internal standardization (Lu 291.139 nm) and measuring the emission intensity at selected wavelengths: (Fe 234.350, 240.489, 259.837 nm).

4.4. X-ray Absorption Spectroscopy (XAS)

X-ray Absorption (XAS) spectra at the Fe K-edge (7112 eV) were collected at the European Synchrotron Radiation Facility (ESRF) of Grenoble, France at the LISA (formerly GILDA) CRG beamline (BM08, [42]). The beamline is equipped with a sagittally focusing monochromator using Si(311) crystals and a pair of Pd-coated mirrors (Ecutoff = 18 keV) for harmonic rejection and beam focusing. The fluorescence emission from the samples was collected using a 12 elements Ge detector, and at least three scans per sample were averaged in order to improve the signal-to-noise ratio. The XAS spectra were recorded between about 6900 and 7700 eV in order to include both the XANES (X-ray Absorption Near Edge Spectroscopy) and the EXAFS (Extended X-ray Absorption Fine Structure) regions. A metallic Fe foil was placed in the second vacuum chamber and its spectrum was collected in transmission mode during every sample scan in order to provide a reliable energy calibration. The beam size ensures the acquisition of spectra over a macroscopic portion of the sample, and the tests for sample beam damage exclude this eventuality.

The obtained spectra have been then analysed by means of appropriate codes (FEFF8, Athena, Artemis; [43]) to obtain information from both the XANES and the EXAFS regions, separately. The pre-edge features of the XANES part of the spectrum were fitted keeping the number of components at a minimum in all cases in order to determine the centroid position and the intensity of the peaks using Origin(Pro)8 (OriginLab Corporation, Northampton, MA, USA).

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Abbreviations

The following abbreviations are used in this manuscript:

BCR	European Community Bureau of Reference
BTs	Back Trajectories
BVM	Bond Valence Method
DMS	Dimethyl Sulfate
EMEP	European Monitoring and Evaluation Programme
ESRF	European Synchrotron Radiation Facility
EXAFS	Extended X-ray Absorption Fine Structure
HNLC	High Nutrient Low Chlorophyll
HVS	High Volume Sampler
ICP-OES	Inductively Coupled Plasma - Optical Emission Spectroscopy
LISA	Linea Italiana per la Spettroscopia d'Assorbimento x
LOD	Limit of Detection
LOQ	Limit of Quantification
MM	Monte Martano
NCEP-GFS	National Centers for Environmental Prediction - Global Forecast System
SWAM	Model of particulate matter sampling instrument (FAI instruments)
WMO SDS-WAS	World Meteorological Organization Sand and Dust Storms-Warning Advisory and Assessment System
XAS	X-ray Absorption Spectroscopy
XANES	X-ray Absorption Near Edge Structure

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