Evidence for ships emissions in the Central Mediterranean Sea from aerosol chemical analyses at the island of Lampedusa

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Abstract

Measurements of aerosol chemical composition made on the island of Lampedusa, south of the Sicily channel, during years 2004–2008, are used to identify the influence of ship emissions on aerosol particles in the Central Mediterranean. Evidence of ship emissions influence is found in 17% of the daily samples. Aerosol samples influenced by ships are characterized by elevated Ni and V soluble fraction (about 80% for aerosol from ships, versus about 40% for crustal particles), high V and Ni to Si ratios, and values of $V_{\text{sol}} > 6 \text{ ng m}^{-3}$. Back trajectories analysis on the selected events show that air masses prevalently come from the Sicily channel, where an intense ship traffic occurs.

$V_{\text{sol}}$, $Ni_{\text{sol}}$, and non-sea salt $SO_{4}^{2-}$ (nss$SO_{4}^{2-}$) show a marked seasonal behaviour, with an evident summer maximum. Such a pattern can be explained by several processes: (i) increased photochemical activity in summer, leading to a faster production of secondary aerosols, mainly nss$SO_{4}^{2-}$, from the oxidation of $SO_{2}$ in the ship plume; (ii) stronger marine boundary layer (MBL) stability in summer, leading to higher concentration of emitted compounds in the lowest atmospheric layers; (iii) more frequent meteorological conditions leading to consecutive days with trajectories from the Sicily channel in summer.

A very intense event in spring 2008 was studied in detail, also using size segregated chemical measurements. These data show that elements arising from heavy oil combustion (V, Ni, Al, Fe) are distributed in the sub-micrometric fraction of the aerosol, and the metals are present as free metals, carbonates, oxides hydrates or labile complex with organic ligands, so that they are dissolved in mild condition (HNO$_3$, pH1.5).

Data suggest a characteristic nss$SO_{4}^{2-}$/V ratio in the range 200–400 for ship emission aerosols in summer at Lampedusa. By using the value of 200 a lower limit for the ship contribution to total sulphates is estimated. Ship emissions account, as a summer average, at least for 1.2 µg m$^{-3}$, representing about 30% of the total nss$SO_{4}^{2-}$, 3.9% of PM$_{10}$, 8% of PM$_{2.5}$, and 11% of PM$_{1}$. Within the used dataset, sulphate from ship
emissions reached the peak value of 6.1 µg m\(^{-3}\) on 26 June 2008, when it contributed by 47 % to nssSO\(_4^{2-}\), and by 15 % to PM\(_{10}\).

1 Introduction

Gas and particles emitted from ships intervene in many processes. Recent studies investigate the impact of gases and particles emitted by ship on human health (Corbett et al., 2007), on acidification and eutrophization of water and soil in coastal regions due to deposition of sulphur and nitrogen compounds (Derwent at al., 2005), and on climate through sulphur containing particles (Devasthale et al., 2006; Lauer et al., 2007), greenhouse gases (Stern, 2007), and absorbing black carbon aerosols (Lack et al., 2009). Moreover, sulphate aerosol has an indirect climate effect influencing cloud structure and properties (e.g., Conover, 1966; Coakley and Walsh, 1987).

Although shipping contributes significantly to the international transportation sector, its emissions are not well quantified and are one of the least regulated anthropogenic sources (IMO, 2008). Several studies point towards the need of international regulations on ship emissions, as those active in Europe, where the land based emissions of sulphur have been successfully reduced since 1980’s.

In this context, it is necessary to investigate the current impact of the ship emissions on the formations of primary and secondary aerosols, and how the predicted future growth of ship traffic and the geographical expansion of waterways and ports, possibly combined with international regulations, are going to affect the atmospheric composition.

Studies on ship aerosol contribution are so far based on inventories and model analyses (e.g., Marmer et al., 2009; Olivier et al., 2005; Eyring et al., 2005; Vestreng et al., 2007) and show that the contribution of ships to air pollution in the Mediterranean atmosphere is significant, although its quantification strongly dependent on the used emission inventory. The verification of ship emission inventories, and particularly those in the open sea, against observations is a difficult task due to lack of continuous
measurements over the open sea and to the complex involved atmospheric processes. Recently, several studies (Petzold et al., 2007; Moldanova et al., 2009; Lack et al., 2009; Agrawal et al., 2008a,b, 2010) reported chemical, physical, and optical properties of emitted particles and gases by analyzing plumes of a large number of commercial shipping vessel, also considering different engines load condition. These studies provide the emission factors of various gases (carbon monoxide, nitrogen oxides, sulphur dioxide, carbon dioxide), chemical compounds in the particulates (S, metals), and aerosol mass emitted from marine engines fed with heavy fuel oil.

Experimental studies of ship aerosol contribution based on aerosol chemical composition are scarce in particular for the Central Mediterranean Sea, whose atmosphere appears to be one of the most polluted in the world due to the high population density (Kouvarakis et al., 2000; Lelieveld and Dentener, 2000; Marmer and Langmann, 2005) and where a very intense ship traffic, connecting the Atlantic and the Indian Oceans, occurs.

The present paper presents the first experimental identification and quantification of ship aerosols based on chemical analyses of PM$_{10}$ samples collected at the island of Lampedusa in the Central Mediterranean Sea.

2 Measurements and methods

The measurements were carried out at the Station for Climate Observations, maintained by ENEA (the national Agency for New Technologies, Energy, and Sustainable Economic Development of Italy) at Lampedusa (35.5° N, 12.6° E). Lampedusa is a small island in Central Mediterranean sea far at least 100 km from the nearest Tunisian coast. At the Station for Climate Observations, which is located on a 45 m a.s.l. plateau on the north-eastern coast of Lampedusa, continuous observations of greenhouse gases concentration (Artuso et al., 2009, 2010), aerosol properties (Pace et al., 2006; Meloni et al., 2006), total ozone, ultraviolet irradiance (di Sarra et al., 2002; Meloni et al., 2004; Di Biagio et al., 2009), and other climatic parameters are carried
In this study we will use aerosol optical properties measured with a multi-filter rotating shadowband radiometer (MFRSR), and chemical analyses of sampled aerosols on filters.

The MFRSR (Harrison et al., 1994) is a seven-channel radiometer which measures global and diffuse irradiances, and allows the determination of column aerosol optical depth at 5 wavelengths (416, 496, 615, 671, and 869 nm). The measurement details and data retrieval is described by Pace et al. (2006).

The aerosol sampling was started in June 2004 at daily resolution, alternating in sequence sampling of PM$_{10}$, PM$_{2.5}$, and PM$_{1.0}$ (particulate matter with aerodynamic diameter lower than 10, 2.5 and 1.0 µm, respectively). Only the PM$_{10}$ sampling head at daily resolution was used since 2007. During the sampling period some interruptions occurred due to technical failures. Here results on the chemical composition of PM$_{10}$ are reported. The sampler was loaded with 47 mm diameter 2 µm nominal porosity Teflon filters. The filters were weighted before and after sampling in order to obtain the mass of the collected atmospheric particulate. All filters were conditioned for at least 24 h prior to weighing at a relative humidity of 35–45 % and temperature of 25°C.

A quarter of each Teflon filter was extracted using MilliQ water (about 10 ml, accurately evaluated by weighing) in ultrasonic bath for 15 min, and the ionic content was evaluated by ion chromatography. Every sample was analysed for cations (Na$^+$, NH$_4^+$, K$^+$, Mg$^{2+}$, Ca$^{2+}$), inorganic anions (F$^-$, Cl$^-$, NO$_3^-$, SO$_4^{2-}$), and some organic anions (methanesulphonate – MSA, acetate, formate, glycolate, oxalate) as reported in Becagli et al. (2011).

Another quarter was extracted in ultrasonic bath for 15 min with MilliQ water acidified at pH 1.5–2 with ultrapure nitric acid obtained by sub-boiling distillation. This extract was used for determination of the soluble part of metals by Inductively Coupled Plasma Atomic Emission Spectrometer (ICP-AES, Varian 720-ES) equipped with an ultrasonic nebulizer (U5000 AT$^+$, Cetac Technologies Inc.). Samples have been spiked with 100 ppb of Ge used as internal standard ($\lambda$ = 209.426 nm), and calibration stan-
standards were prepared by gravimetric serial dilution from mono standards at 1000 mg l\(^{-1}\). The value of pH was chosen because it is the lowest values found in rainwater (Li and Aneja, 1992) and leads to the determination of the metals fraction more available for biological organisms. Filters field blancks show V and Ni soluble fractions (hereafter \(V_{\text{sol}}\) and \(Ni_{\text{sol}}\)) concentrations below the detection limits (0.04 and 0.09 ng m\(^{-3}\), respectively) in working conditions.

The remaining half filters were analysed for the total (soluble + insoluble) elemental composition by the proton induced X-ray emission (PIXE) technique (Calzolai et al., 2006; Chiari et al., 2005). The PIXE analysis was carried out for a reduced number of samples, therefore, the elemental composition is available for a restricted data set. In addition, the amount of V was in several cases below the minimum detection limit (MDL) of the PIXE technique.

Additional sampling with an 8 stage impactor equipped with Teflon back-up filter was performed in Lampedusa in the period 17 May–20 June 2008. Half of each filter was used to determine the ionic composition, while the other half was used to measure the metals soluble fraction content with the same methodology above described.

3 Results

3.1 Identification of ship aerosols: chemical composition

Sulphur is the dominant element in the exhausts of heavy fuel oil, followed by V and Ni (Agrawal et al., 2008a). The metals (V and Ni) are also present in the Earth crust as minor constituents. The mean upper continental crustal V/Ni ratio is 2.06 w/w (HENDERSON and HENDERSON, 2009). Heavy oil is enriched in V and Ni content with respect to the crust, and these metals are generally used as markers of heavy oil combustion in all size fractions of the atmospheric particulate. Mazzei et al. (2008) report a characteristic value of \(V/Ni = 3.2\) for ships, obtained by applying statistical approaches (Positive Matrix Factorization) to an extensive chemical data set of aerosol sampled near a har-
bour. A wide V/Ni ratio (2.3–4.5) was measured by direct sampling at the exhausts of different auxiliary ship engines fed by different fuels (Nigam et al., 2006), and from the main propulsor ship engine at different speed mode (Agrawal et al., 2008a,b).

In this study we used the Ni/Si and V/Si ratios to distinguish between V and Ni due to heavy oil combustion and to crustal sources. Silicon is considered the main marker for crust. Ni and V determined by PIXE (i.e. their total content) was used for Ni/Si and V/Si ratios calculation. The Ni/Si ratio is one order of magnitude higher than expected for crustal particles (dashed line in Fig. 1) in 79% of the measured samples.

An higher fraction of enriched samples (more than 10 times above the crustal ratio) is obtained for the V/Si ratio (88% of the samples are enriched in V). The V total content was determined in a minor number of samples than Ni because there are cases with V below the PIXE detection limit 2.6 ng m\(^{-3}\). As a consequence, the fraction of data with elevated ratio is higher for V/Si than for Ni/Si.

Most of the samples considered not enriched (V/Si < 10 \(\cdot\) (V/Si)\(_{\text{crust}}\) and Ni/Si < 10 \(\cdot\) (Ni/Si)\(_{\text{crust}}\)) belongs to days in which the crustal content is dominant (i.e. Si > 800 ng m\(^{-3}\), red dots in the plot) and back trajectories (not shown for brevity) confirm that the air masses originate from the Sahara desert or pass over the desert before reaching Lampedusa.

The measured concentrations of V\(_{\text{sol}}\) and Ni\(_{\text{sol}}\) for the non-enriched events are lower than for the enriched cases, and are below 8 and 2.6 ng m\(^{-3}\), respectively. Only 6 events (less than 5% of the dataset) are in the range 6–8 ng m\(^{-3}\) for V, and 2.3–2.6 ng m\(^{-3}\) for Ni. The threshold of 6 ng m\(^{-3}\) (hereafter V\(_{\text{st}}\)) for V\(_{\text{sol}}\) is chosen to identify enriched samples on the basis of measurements of V\(_{\text{sol}}\).

Soluble fractions of V and Ni, instead of the respective total amounts derived from the PIXE analyses, are used in the following discussion because the soluble fraction represents the anthropic sources better than total content. Indeed, the V and Ni solubility appear to depend on the originating source. By plotting V\(_{\text{sol}}\) and Ni\(_{\text{sol}}\) versus the total V and Ni, respectively, we obtain two different behaviours: the ship aerosol cases (V\(_{\text{sol}}\) > 6 ng m\(^{-3}\)) display a larger slope, thus a higher solubility than and Saharan dust
events (Si > 800 ng m\(^{-3}\) and \(V_{\text{sol}} < 6 \text{ ng m}^{-3}\)). Table 1 reports slopes and regression coefficients for the various plots. In the events classified as influences by anthropic source (i.e. high temperature processes) both V and Ni are easily dissolved (the soluble fraction is 80 % and 77 % for V and Ni, respectively) in the mild extraction condition (HNO\(_3\) – pH 1.5) because they are mainly present as free metals, oxides hydrates, or organo-metal compounds. On the contrary, the presence of V and Ni in the silica matrix or as oxides in samples with high dust content is expected to produce a lower solubility (40 % and 45 % for V and Ni, respectively). Consequently, samples influenced by ships have a higher content of \(V_{\text{sol}}\) and \(Ni_{\text{sol}}\), which are present in the environment in the available form for biological organisms, and are more capable to exert toxicity.

Figure 2 shows the temporal evolution of the daily average aerosol optical depth (AOD) at 500 nm, of the daily \(PM_{10}\) mass concentration, and of the main markers of heavy fuel oil: non sea salt sulphate (nssSO\(_4^{2-}\)), \(V_{\text{sol}}\) and \(Ni_{\text{sol}}\) from \(PM_{10}\) samples collected at Lampedusa in the period June 2004–December 2008. Three-month averages of the measured parameters are also shown to emphasize the seasonal evolution. Cases in which the measured AOD is strongly influenced by Saharan dust are also reported in the figure (red dots in Fig. 2a). The method by Pace et al. (2006), based on the measured values of AOD and Ångström exponent and on the analysis of the backward trajectoreis, was used for the identification of cases with large Saharan dust contribution to the AOD. Since the AOD provides information on the entire air column, the identification of a Saharan dust case does not necessarily imply that mineral particles are present close to the surface.

Figure 2 shows a marked seasonal pattern with spring/summer maxima of the AOD and of the tree chemical markers. As discussed by Di Iorio et al. (2009), the dust optical depth and vertical distribution show a large seasonal cycle, with elevated AOD and a wider vertical extension in spring and summer; the seasonal change is mainly controlled by dust transport occurring over the boundary layer. On the contrary, non dust cases and boundary layer aerosols display a very limited seasonal change (black dots and blue line in Fig. 2a), as it is also confirmed by the \(PM_{10}\) measurements (Fig. 2b).
The PM$_{10}$ concentration reaches elevated values, up to about 140 µg m$^{-3}$, mainly in spring; also the highest peaks in AOD in 2007 and 2008, close to 1, occur in spring. The highest peaks in PM$_{10}$ and AOD are due to Saharan dust events, and will not be discussed in detail here. Elevated values and isolated peaks of V$_{\text{sol}}$, Ni$_{\text{sol}}$, and nssSO$_4^{2-}$ are observed throughout spring/summer and especially in June/July. A scatter plot of Ni$_{\text{sol}}$ versus V$_{\text{sol}}$, and of nssSO$_4^{2-}$ versus V$_{\text{sol}}$ is shown in Fig. 3. V$_{\text{sol}}$ and Ni$_{\text{sol}}$ are closely related, suggesting that the two species originate from the same source. The behaviour of non-sea salt sulphate, conversely, does not appear to be closely linked to V$_{\text{sol}}$ and Ni$_{\text{sol}}$.

Beside heavy oil combustion, several sources (anthropic origin from long range transport, marine biogenic, crustal, volcanic) contribute to the non sea salt sulphate in Central Mediterranean Sea. For this reason V$_{\text{sol}}$ and nssSO$_4^{2-}$ are not directly coupled, and the quantification of SO$_4^{2-}$ contribution from ship emissions to the nssSO$_4^{2-}$ total budget is a difficult task. Looking in detail the V$_{\text{sol}}$–Ni$_{\text{sol}}$ correlation (reported in Fig. 3a for the whole data set) we found an higher correlation ($R^2 = 0.978$, $n = 131$) between V$_{\text{sol}}$ and Ni$_{\text{sol}}$ for the cases classified as influenced by anthropic emissions, with a slope of 2.98 ± 0.04, in accord with those reported in literature for this source (Nigam et al., 2006; Mazzei et al., 2008; Agrawal et al., 2008a,b). Anyway, very good correlation is also found between V$_{\text{sol}}$ and Ni$_{\text{sol}}$ ($R^2 = 0.945$, $n = 144$) for events classified as crustal; for these events the slope of the linear fit is lower (2.54 ± 0.05) and nearer to the mean V/Ni crustal ratio (2.06). Mixing between particles of crustal and ship origins may play a role also in events characterised by a high dust content.

3.2 Identification of ship aerosols: trajectory analysis

A progressive vector analysis based on wind measurements of the meteorological station at Lampedusa was carried out to determine the relationship between the air mass chemical composition and origin.
Back-trajectories were calculated using local wind data measured at the Station for Climate Observations on a 10 m tall mast with a time resolution of 10 min. The method is based on the assumption that at each time step the wind along the back trajectory is equal to the one measured at the same time at Lampedusa.

The geographic position of Lampedusa, i.e. far at least 100 km from the nearest Tunisian coast, makes this methodology applicable.

The validity of the method is however limited to the maritime area surrounding Lampedusa and to a limited time interval. In particular this kind of analysis is not reliable in proximity of the coasts or over land. On the other hand, the use of relatively frequent wind measurements can be advantageous with respect to standard back-trajectories based on meteorological analyses, which have a broader spatial and temporal resolution, especially considering the goal of this trajectory analysis: the determination of the air mass paths in the maritime region surrounding Lampedusa.

Figure 4 shows 18-h long trajectories arriving at Lampedusa at the middle of the PM sampling interval. Trajectories are grouped in three-month periods, and plotted separately for cases with $V_{sol} < 6 \text{ ng m}^{-3}$ and $V_{sol} > 6 \text{ ng m}^{-3}$. Similar results are found by using 12 or 24-h trajectories. Trajectories were also calculated for different arrival times, at the beginning or at the end of the PM sampling interval. Except than in few cases, no substantial changes in the trajectory pattern are found.

The origin of the air masses corresponding to $V_{sol} < V_{st}$ reflects the overall wind direction statistics at Lampedusa, without specific preferred directions (22% of cases wind originates from N–NW; all the other directions show frequencies of occurrence between 2% and 5%). The air masses showing evidence of ship influence display a strong dependency on the originating direction: they are mostly of northern origin, and passed over the Strait of Sicily, i.e. in correspondence with the main maritime route crossing the Mediterranean sea from the Strait of Gibraltar to the Suez Canal. 26.5% (35) of the cases with $V_{sol} > V_{st}$ correspond to air masses coming from a narrow direction ($345^\circ \pm 7.5^\circ$); 63.6% (84) of the cases correspond to air masses originating between $322.5^\circ$ and $37.5^\circ$. The same region also includes 80.8% of the cases with
V_{sol} > 10 (52 cases). Trajectories with V_{sol} > V_{st} do not show any significant dependency on the wind intensity (the mean wind intensity along the back trajectory ranges from 1.5 to 11 m s^{-1}).

The good correspondence between the air mass origin and the Mediterranean main route of large vessels confirms the validity of the selected V_{sol} threshold for identifying PM samples influenced by ship emissions. Figure 5 shows a statistic of the number of cases with V_{sol} < 6 and V_{sol} > 6 ng m^{-3} for different originating trajectory directions. Data are displayed for April-May-June, and July-August-September, separately; only two quarterly statistics are shown because the few ship events measured in the autumnal and winter quarter. Evidently, as also appears from Fig. 4, air masses coming from Sicily Channel during autumn and winter does not necessarily show elevated values of V_{sol}, suggesting that transport from north north-west is required, but is not sufficient to determine high concentration of ship aerosol; different processes probably play a role in the observed seasonal evolution.

### 3.3 Seasonal evolution

As shown in Fig. 2, V_{sol}, Ni_{sol}, and nssSO_{4}^{2-} display a large seasonal cycle, with a marked maximum in spring/summer. Three-month averages of the different parameters displayed in Fig. 2 have been added to highlight their seasonal evolution. The yearly evolution of the three-month averages of V_{sol} and nssSO_{4}^{2-} appear remarkably similar, suggesting that, although the sources may not be coupled, they may partly respond to similar mechanisms. As discussed above, the seasonal behaviour is not due to a significant change of the dynamical patterns lading to more frequent trajectories from the ship route. Different processes may contribute to produce this seasonal cycle.

Production of secondary aerosol is influenced by solar radiation. This effect is well known for the formation of nssSO_{4}^{2-}, which is derived from oxidation of SO_{2} by OH (see, e.g., Barbu et al., 2009). In its turn, the OH production is linked to elevated levels of ultraviolet radiation. The Central Mediterranean is characterized by elevated levels of UV radiation in summer, and high ozone photolysis and OH production (e.g.,
Consequently, SO₂ conversion into SO₄²⁻ is expected to be faster in summer. Ault et al. (2010), in a recent study found V and SO₄²⁻ in single aerosol particles from fuel combustion; they found that SO₄²⁻ is in higher concentration than in particles not containing V and suggest a catalytic effect of V on the oxidation of SO₂ to SO₄²⁻. In this process the metals would be involved in the secondary aerosol formation, favoured by high level of solar radiation.

The structure of the planetary boundary layer is expected to play a role in producing the observed seasonal cycle. Song at al. (2003) used a Lagrangian photochemical box model for an air parcel emitted from ship in the marine boundary layer. They found that conditions favouring the stability of the marine boundary layer produce a larger increase in the SO₂ and sulphate concentrations than an increased emission of SO₂. The marine boundary layer is generally more stable in summer than in winter (e.g., Dayan et al., 1989), and this effect may produce elevated concentrations of sulphate and metals during this season.

Also the variability of the traffic ship can play a role in determining the observed annual cycle. Nevertheless Marmer and Langmann (2005) found summer mean concentration of sulphate aerosol in the lowest level of their model almost four times higher than in winter, using constant emissions throughout the year. This result suggests that a possible annual ship emission variability is not the dominant factor for explain the observed seasonal cycle.

The number of ship aerosol events with duration of more than one day is also largest in summer.

Table 2 presents some statistics on V_{sol} measurements and the cases of consecutive days with V_{sol} > V_{st}. The whole database extends over 5 yr, but only the last 2 yr are characterised by regular daily samples. As shown in Table 2, from 2004 to 2006 there are a total of 235 V_{sol} measurements (i.e. yearly number of measurements varying between 66 and 91), and only about 12% of the cases presents V_{sol} > V_{st}.

The representativeness of 2007 and 2008 is definitely larger than in previous years, with a total of 549 daily measurements of V_{sol}. Thus we believe that results for 2007–
2008, with about 19% of cases with $V_{\text{sol}} > V_{\text{st}}$ is more representative for the normal situation at the Lampedusa.

A simple statistics on ship events lasting more than one day is reported in Table 2. The number of cases lasting more than one day is larger in 2007–2008 (70.2% of the cases last between 2 and 5 consecutive days) than in the 2004–2005 period (28.6%); however, results for 2004–2006 are biased, since PM$_{10}$ samples were not taken continuously. This effect tends to produce an underestimate of the influence of ship events in 2004–2006. Moreover, most of the episodes lasting more than one day show mean $V_{\text{sol}}$ values larger than the mean $V_{\text{sol}}$ of the single day cases, suggesting a progressive increase of the ship aerosol load in specific situations, leading to a strong intensity of episodes lasting for more than one day.

### 3.4 June/July 2008 event: trajectory analysis

Figure 6 shows the back trajectories of a particular long lasting (20 days) and interesting event in which the ship influence on the aerosol chemical composition was observed to occur almost continuously from 20 June to 9 July, 2008. Elevated values of $V_{\text{sol}}$ were measured throughout the period; $V_{\text{sol}}$ decreased below $V_{\text{st}}$ during only 3 days.

At the beginning of the event (20 and 21 June) the wind intensity decreased and advected air from the middle of the Sicily channel. From 22 to 24 June there is a notable reduction of the wind intensity, and easterly winds. On 25 June the wind was again coming from north-west, and values of $V_{\text{sol}} < V_{\text{st}}$ were measured. From 26 June to 2 July $V_{\text{sol}}$ reached very high values, including the maximum value of the whole dataset (30.6 ng m$^{-3}$) on 26 June; 4 out of the 7 values of $V_{\text{sol}} > 20$ ng m$^{-3}$ of the dataset occurred between 26 June and 1 July. In this period the air mass came from north, progressively shifting from northwest to northeast. A change of the wind direction led to southern local air masses and $V_{\text{sol}} < V_{\text{st}}$ on 3 July. With the exception of the back trajectory of 7 July, when southern air mass induced a low value of $V_{\text{sol}}$, the period ends with a recovery of the northerly air mass flow, still inducing high values of $V_{\text{sol}}$. 
This long extended period with influence from ship emissions confirms that the origin of the air masses plays a central role. In particular, the strait of Sicily seems to be the main source region. Moreover, very low winds connected with stagnant conditions are not responsible for elevated values of $V_{\text{sol}}$, confirming that high $V_{\text{sol}}$ values are not of local origin (i.e. from harbour activities).

3.5 June/July 2008 event: size segregated chemical composition

Figure 7 shows the particle size distribution derived from multi-stage impactor sampling. The size distribution of the main markers of ship emissions and crustal elements for the first day of the described event (20 June 2008) and for 17 May 2008, characterized by a high crustal content, are shown. The backward trajectories of 17 May, computed over 24 h, shows that the airmass originated from North Africa.

As expected, V and Ni in the ship aerosol event display a maximum in the finest mode (diameter < 0.4 µm). Conversely, their concentrations peak at larger size (1.1–2.1 µm for Ni, and 0.4–0.7 µm for V) during the Saharan dust event.

The sulphate distribution shows a monomodal distribution peaked in the 0.4–0.7 µm fraction during the ship event, due to the main secondary source. On 17 May sulphate displays a bimodal distribution with a second mode (1.1–2.1 µm) related to its primary sources from Saharan dust and sea spray.

Al and Fe are considered typical markers of the crustal source and are mainly present in the coarse mode on 20 June. During the ship event their distributions are shifted towards fine particles, suggesting that also the anthropic source may contribute to the occurrence of these elements. The presence of a fine mode also on 17 May suggest the presence of mixed source in the selected event.

It must be noticed that even if for Ni and V we report here the pH 1.5 soluble fraction, and the solubility is different for the two aerosol sources, their concentrations are higher in the ship event than during the Saharan dust case. The situation is different for the crustal markers. Al shows higher concentrations in the Saharan dust event, while Fe concentrations are higher in the ship case. Part of this behaviour is due to the
different solubility of Fe of different origin. In Saharan dust aerosol Fe is associated with the silicate matrix, or present as oxides, and these compounds are not soluble in the applied conditions (HNO₃ – pH 1.5). In PM₁₀ samples collected at Lampedusa Fe_{sol} is generally less than 10% of the total Fe. On the contrary, the Fe solubility is high for anthropic sources, due to the presence of metallic iron or hydrated iron oxide, both soluble in HNO₃.

### 3.6 Ship contribution to the total aerosol load

By using specific markers of different sources and applying receptor models it is possible to quantify the mass contribution of a specific source to the total PM (Gordon, 1988). Emission factors from different marine vessels and engines can provide insightful information to aid source apportionment studies even if the ratios between the selected markers change from the source to the receptor site for secondary specie such as nssSO₄²⁻. The main species emitted in the ship plume in gas phase and precursor of particulate are NOₓ and SO₂ (Agrawal et al., 2008a,b). The photochemistry of NOₓ leading to NO₃⁻ in the particulate phase is complex, especially in summer (see e.g., Chen et al., 2005), and their contribution to the particulate phase is not here quantified. In this section the ship emissions contribution of SO₄²⁻ to the total amount of nssSO₄²⁻ and to the PM mass is estimated, considering the ratios between the precursor of particulate in the exhausts of the ship engine and V, which is taken as a representative and conservative marker of ship emission. Agrawal et al. (2008a,b) report average SO₄²⁻/V emission ratio of 11–27 in the particulate exhaust (PM₂.₅) of the main engine of different ocean going container vessels operating at different load. The same authors report higher values (up to about 500) for SO₂(gas)/V(particulate). However, the amount of SO₄²⁻ in the airmass is expected to grow fast due to SO₂ conversion into sulphate; this conversion is faster in high UV radiation and high humidity conditions.

If we assume an e-folding time of about 2 days for the SO₂ to SO₄²⁻ conversion (Restad et al., 1998), the SO₄²⁻ mass is expected to increase by a factor of 10 within
10 h with respect to that contained in the air mass 1 h after the emission. Since the starting \(\text{SO}_4^{2-}/V\) ratio is in the range 11–27, after 10 h we expect to have a \(\text{SO}_4^{2-}/V\) ratio of about 110–270. This ratio is expected to vary still more largely, depending on the efficiency of the dry and wet \(\text{SO}_2\) oxidation, and the elapsed time between emission and aerosol sampling. Sulphates originating from processes other than ships contribute to the total \(\text{SO}_4^{2-}\) load in the collected aerosol, and larger \(\text{SO}_4^{2-}/V\) ratios may be expected. As shown by Fig. 3, all data display nss\(\text{SO}_4^{2-}/V \geq 200\), and we take \(\text{SO}_4^{2-}/V = 200\) as a lower limit for particles originating by ships at Lampedusa. In eighteen cases, highlighted in red in Fig. 3, the ratio is close to this limit. Except for two days in March, all cases with nss\(\text{SO}_4^{2-}/V\) close to 200 occur between May and August. During the special event of June/July 2008 the nss\(\text{SO}_4^{2-}/V\) ratio values were all in the range 250–400, except for few spikes up to 1000, possibly due to the contribution from additional sources of \(\text{SO}_4^{2-}\). A value of \(\text{SO}_4^{2-}/V\) in the same range (285) is derived for the finest particles stage (0.1–0.4 µm) of the impactor data for the ship event discussed in Sect. 3.5. Newly formed secondary particles, in this particular event mainly arising from ship emissions, are predominantly present in this size range. Conversely, larger particles show values of \(\text{SO}_4^{2-}/V > 400\), due to an expected stronger contribution of sulphate particles arising from other sources, and the role of coagulation and accretion processes during transport.

By using \(\text{SO}_4^{2-}/V = 200\) as a limit value for ship emission in summer and using the measured values of \(V_{\text{sol}}\) determined at Lampedusa, it is possible to obtain a rough estimation of the minimum contribution of sulphate from the ship source to the total sulphate and total aerosol mass. Due to different meteorological conditions and photochemical activity such a value can not be applied to samples collected in winter.

For the recognised ship aerosol events the average \(\text{SO}_4^{2-}\) contribution is \(2.12 \mu g \text{ m}^{-3}\), corresponding to 43.6 % of total nss\(\text{SO}_4^{2-}\) mass. This value is higher than the value modelled by Endersen et al. (2003; 2.9 %) as which is however an annual global average. The high values we found are likely due the regional (highly affected by ship
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4 Conclusions

Daily aerosol samples collected at Lampedusa from 2004 to 2008 were analyzed to determine their chemical composition and to identify the possible influence of emissions from ships in the Central Mediterranean.

Ship emissions events are identified using V and Ni, specific markers of this source. Their ratio to Si is used to discriminate between crustal and ship sources. The Ni and traffic) and seasonal (late spring/summer) scale here considered. From regional model study Marmer and Langmann (2005) estimated an average sulphate concentration of 3.3 µg m\(^{-3}\) over the Mediterranean basin in summer 2002, with higher values in the Sicily channel. The modelled contribution of ship emission respect to land source, considering negligible the natural contribution is as average higher than 50\%. Total nssSO\(_4^{2-}\) values modelled by Marmer and Langmann (2005) in the summer months (JJA) are in the range 4.5–5.2 µg m\(^{-3}\) over the 2002–2003 period in JJA in the Strait of Sicily, consistent with the measured nssSO\(_4^{2-}\) average concentration at Lampedusa: 4.0 µg m\(^{-3}\) over the 2004–2008 period in JJA. In the same time period the mean SO\(_2^{−}\) from ship emission computed using the V\(_{\text{sol}}\) content is 1.2 µg m\(^{-3}\), corresponding to about 30\% of the total nssSO\(_4^{2-}\) content. This percentage is lower than the one modelled by Marmer and Langmann (2005), possibly due to an underestimation of the true SO\(_2^{−}/V\) ratio and/or the underestimation of long range and especially marine biogenic contribution in the model.

During the June/July 2008 special event shown in Fig. 7 (26 June 2008) the nssSO\(_4^{2-}\) from ship emission reached 6.1 µg m\(^{-3}\), contributing by 15\% to the total PM\(_{10}\). As shown in Sect. 3.5, the ship aerosols belong mainly the fine particles fraction, and a by using the average summer V content and mass data obtained at Lampedusa for PM\(_{2.5}\) and PM\(_{1}\) we obtain a contribution of SO\(_2^{−}\) from ship aerosol of about 8\% and 11\%, respectively, in the two aerosol fractions in summer.
V soluble fractions are consistently higher for the cases influenced by ship emissions than for mineral particles.

The V soluble fraction was chosen to identify days affected by high ship aerosol content. A threshold of 6 ng m$^{-3}$ was established to select ship aerosol events.

A progressive vector analysis based on wind measurements confirms that the selected events are affected by sea going ship, mainly transiting in the Sicily Chanel, and not from local pollution (i.e. Lampedusa harbour activities).

A marked seasonal behaviour, with summer maxima, is observed for all the main ship aerosol markers ($V_{sol}$, $Ni_{sol}$ and nssSO$^{2-}$). The following processes occurring in summer are believed to play an important role in determining the observed seasonal evolution:

1 – Increased photochemical activity of atmosphere in summer, leading to a larger production of secondary aerosols, mainly nssSO$^{2-}$.

2 – MBL stability, higher in summer than winter, that prevents the accumulation of ship emission compounds in the lowest atmospheric levels in winter.

The higher occurrence of late spring/summer ship episodes lasting more than one day in correspondence with dominant wind direction from the Sicily Channel, indicates an increased contribution of ship emission to the aerosol load in Central Mediterranean, modulated by the atmospheric dispersion and transport processes.

A very intense event in spring 2008 was chemically and size characterised showing that elements arising from heavy oil combustion (V, Ni but also Al, Fe) are distributed in the sub-micrometric fraction of the aerosol. The metals are present as free metals, carbonates, oxides hydrates or labile complex with organic ligands, so that they are dissolved in mild condition (HNO$_3$, pH1.5).

The SO$_{4}^{2-}$/V ratio of 200 is proposed as lower limit characteristic ratio for ship emission in summer at Lampedusa. The experimental determination of characteristic ratio of ship markers in a receptor site is particularly useful for understanding and quantifying the profile emission.
Between the components emitted by ship, $\text{SO}_4^{2-}$ contributes significantly to the total PM mass. In summer the estimated average mass is 1.2 $\mu\text{g m}^{-3}$, about 30% of the total nss$\text{SO}_4^{2-}$, 3.9% of PM$_{10}$, 8% PM$_{2.5}$, and 11% PM$_1$. Sulphate from ship emissions reach a peak value of 6.1 $\mu\text{g m}^{-3}$ on 26 June 2008, corresponding to 47% of nss$\text{SO}_4^{2-}$, and 15% of PM$_{10}$. It has to be emphasized we used a conservative value for the $\text{SO}_4^{2-}/V$ ratio which is expected to produce a lower limit of the ship-derived sulphates, and the contribution of ship emissions may be higher than derived here.

For future studies, more data on size resolved chemical composition (including other pollutants emitted in the ship plume, e.g. organic and elemental carbon) from sites located in the open sea can provide a better evaluation of the regional and global impact of ship aerosols on the total aerosol budget and on climate.

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**References**

Artuso, F., Chamard, P., Chiavarini, S., di Sarra, A., Meloni, D., Piacentino, S., and Sfer-
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**Table 1.** Correlation parameters between V and Ni soluble and total fractions for ship aerosol cases ($V_{\text{sol}} > 6 \text{ ng m}^{-3}$) and for Saharan dust cases ($\text{Si} > 800 \text{ ng m}^{-3}$ and $V_{\text{sol}} < 6 \text{ ng m}^{-3}$).

<table>
<thead>
<tr>
<th></th>
<th>Slope ($\pm$ error)</th>
<th>$R^2$</th>
<th>n.</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_{\text{sol}}/V_{\text{tot}}$ ship events</td>
<td>0.80 ($\pm$0.02)</td>
<td>0.932</td>
<td>113</td>
</tr>
<tr>
<td>$V_{\text{sol}}/V_{\text{tot}}$ pure Saharan dust events</td>
<td>0.40 ($\pm$0.02)</td>
<td>0.824</td>
<td>91</td>
</tr>
<tr>
<td>$\text{Ni}<em>{\text{sol}}/\text{Ni}</em>{\text{tot}}$ ship events</td>
<td>0.77 ($\pm$0.02)</td>
<td>0.939</td>
<td>113</td>
</tr>
<tr>
<td>$\text{Ni}<em>{\text{sol}}/\text{Ni}</em>{\text{tot}}$ pure Saharan dust events</td>
<td>0.45 ($\pm$0.02)</td>
<td>0.785</td>
<td>127</td>
</tr>
</tbody>
</table>
Table 2. For the two periods having different resolution time (2004–2006 and 2007–2008) are reported: the number of PM samples (measurements of $V_{\text{sol}}$), the number (frequency occurrence on the total number of annual $V_{\text{sol}}$ measurements) of measurements with $V_{\text{sol}} > 6 \text{ ng m}^{-3}$, the mean and standard deviation values. The number of episodes of single and consecutive days (frequency occurrence on the number of annual $V_{\text{sol}}$ measurements affected by ship emission) of $V_{\text{sol}} > 6 \text{ ng m}^{-3}$, the correspondent periods mean and standard deviation values of the single and multiple day episodes are also presented.

<table>
<thead>
<tr>
<th>Year</th>
<th>Number of PM ($V_{\text{sol}}$) samples</th>
<th>Number of $V_{\text{sol}} &gt; 6 \text{ ng m}^{-3}$ measurements, annual occurrence, mean and std values</th>
<th>Number of episodes single or consecutive days with $V_{\text{sol}} &gt; 6 \text{ ng m}^{-3}$, daily annual occurrence, mean and std values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>single day</td>
<td>2 days</td>
</tr>
<tr>
<td>2004–2006</td>
<td>241 (235)</td>
<td>28</td>
<td>20 episodes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>11.9 %</td>
<td>71.4 %</td>
</tr>
<tr>
<td></td>
<td></td>
<td>9.7 ± 2.8</td>
<td>8.1 ± 2.8</td>
</tr>
<tr>
<td>2007–2008</td>
<td>594 (549)</td>
<td>104</td>
<td>31 episodes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>18.9 %</td>
<td>29.8 %</td>
</tr>
<tr>
<td></td>
<td></td>
<td>11.0 ± 5.3</td>
<td>8.2 ± 2.5</td>
</tr>
</tbody>
</table>
Fig. 1. Temporal evolution of $\text{Ni}_{\text{tot}}/\text{Si}_{\text{tot}}$ in PM$_{10}$ samples collected at Lampedusa island. Black line represent the Ni/Si mean ratio in upper continental crust (Henderson and Henderson, 2009), dashed line represent the threshold values for sharing samples enriched (one order of magnitude higher than black line). Red dots are samples with high dust content ($\text{Si} > 800 \text{ ng m}^{-3}$).
Fig. 2. Temporal evolution of AOD, PM$_{10}$, nssSO$_4^{2-}$, $V_{\text{sol}}$, and Ni$_{\text{sol}}$, at Lampedusa island. Blue lines in plot (b,c,d,e) represent the three months mean. Red dots in the plot a are related to Saharan dust events chosen as AOD > 0.15 and $a < 0.5$ (Pace et al., 2006). Black dots represent the remaining days. In this plot the dark line represent the AOD three month average, the blue one the three month mean excluding the Saharan dust event (i.e. calculated over the black dot).
Fig. 3. Scatter plots $N_{i\text{sol}}$ (a) and nssSO$_4^{2-}$ (b) vs. $V_{\text{sol}}$. Line in plot a represents the linear correlation, dashed line in the plot b represents the ratio nssSO$_4^{2-}/V_{\text{sol}}=200$. Red dots in plot (b) represent the eighteen events for which the nssSO$_4^{2-}/V$ ratio is about 200.
Fig. 4. Figures (a, c, e, g) and (b, d, f, h) show, respectively the 18 h back trajectory relatives to the measurements of $V_{sol}$ less than or great/equal to 6 ng m$^{-3}$ (i.e. $V_{st}$). As reported in figure the trajectories are grouped in quarters, starting from January and ending with December.
Fig. 5. Number of cases with $V_{\text{sol}} < 6 \, \text{ng m}^{-3}$ (blue regions) and $V_{\text{sol}} > 6 \, \text{ng m}^{-3}$ (red regions) for different originating directions of the corresponding airmass trajectories for April-May-June (left), and July-August-September (right).
Fig. 6. 18-h back trajectories for the 20 June–9 July 2008 period, when only 3 daily $V_{\text{sol}}$ measurements (25 June, 3 July and 7 July) were below $V_{\text{st}}$. The 24-h back trajectory of 17 May is also presented. The number at the end of each back-trajectories indicates the corresponding day of June and July.
Fig. 7. $V_{\text{sol}}$, $\text{Ni}_{\text{sol}}$, $\text{SO}_4^{2-}$, $\text{Al}_{\text{sol}}$, $\text{Fe}_{\text{sol}}$ size distributions during a ship event (plots a, c, e, g, i, 20 June 2008) and during a Saharan dust event (plots b, d, f, h, l, 17 May 2008).