Vertical distribution of aerosol optical properties in the Po Valley during the 2012 summer campaigns

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Abstract. Studying the vertical distribution of aerosol particle physical and chemical properties in the troposphere is essential to understand the relative importance of local emission processes vs. long-range transport on column-integrated aerosol properties (e.g. the aerosol optical depth (AOD), affecting regional climate) as well as on the aerosol burden and its impacts on air quality at the ground. The main objective of this paper is to investigate the transport of desert dust in the middle troposphere and its intrusion into the planetary boundary layer (PBL) over the Po Valley (Italy), a region considered one of the major European pollution hot-spots for the frequency of particulate matter (PM) limit values exceedances. Events of mineral aerosol uplift from local (soil) sources and phenomena of hygroscopic growth at the ground are also investigated, possibly affecting the PM concentration in the region as well. During the PEGASOS 2012 field campaign, an integrated observing-modeling system was set up based on near-surface measurements (particle concentration and chemistry), vertical profiling (backscatter coefficient profiles from LiDAR and radiosoundings) and Lagrangian air masses transport simulations by FLEXPART model. Measurements were taken at the San Pietro Capofiume supersite (44°39’ N, 11°37’ 0 E, 11 m a.s.l), located in a rural area relatively close to some major urban and industrial emissive areas in the Po Valley. Mt. Cimone (44°12’ N, 10°42’ E, 2165 m a.s.l.) WMO/GAW station observations are also included in the study to characterize regional-scale variability. Results show that, in the Po Valley, aerosol is detected mainly below 2000 m a.s.l. with a prevalent occurrence of non-depolarizing particles (> 50 % throughout the campaign) and a vertical distribution modulated by the PBL daily evolution. Two intense events of mineral dust transport from Northern Africa (19–21 June and 29 June to 2 July) are observed, with layers advected mainly above 2000 m height, but lately sinking and mixing in the PBL. As a consequence, a non-negligible occurrence of mineral dust is observed close to the ground (~ 7 % of occurrence during a 1-month campaign). The observations unambiguously show Saharan dust layers intruding the Po Valley mixing layer and directly affecting the aerosol concentrations near the surface. Finally, LiDAR observations indicate also strong variability in aerosol on shorter timescales (hourly) highlighting: a) events of hygroscopic growth of anthropogenic aerosol, visible as shallow layers of low depolarization near the ground. Such events are identified during early morning hours at high relative humidity (RH) conditions (RH > 80 %). The process is observed
concurrently with high PM1 nitrate concentration (up to 15 µg cm$^{-3}$), hence mainly explicable by deliquescence of fine anthropogenic particles, and during mineral dust intrusion episodes, when water condensation on dust particles could instead represent the dominant contribution; b) frequent events (mean diurnal occurrence of $\sim$22% during the whole campaign) of rapid uplift of mineral depolarizing particles in afternoon-evening hours up to 2000 m a.s.l. height. The origin of such particles cannot be directly related to long-range transport events, being instead likely linked to processes of soil particles resuspension from agricultural lands.

1 Introduction

The Po river basin in Northern Italy is one of the most important emissive area in Europe, characterized by high concentration of both natural and anthropogenic aerosol and trace gases (Monks et al., 2009). The geographical location of this region, surrounded by two mountain ranges, promotes frequent occurrence of stagnant meteorological conditions (Rossa et al., 2012), with accumulation of local pollution (Hanke et al., 2003; Crosier et al., 2007; Saarikoski et al., 2012; Decesari et al., 2014) from industrial, urban and agricultural emissions, and complex processes of aerosol-chemicals transformation. At the same time, the relative proximity to the Sahara desert, that represents the major mineral dust source of the planet (Prospero et al., 2002; Washington et al., 2003), makes this region subject to long range mineral dust transport, especially during the summer season (Kalivitis et al., 2007; Marinoni et al., 2008; Pederzoli et al., 2010; Carnevale et al., 2012). Such combination leads to unusually high concentrations of atmospheric pollutants and particulate matter, with frequent and prolonged periods of intense pollution. The large amount of people living in the region (more than 20M potentially exposed to high pollution levels) accentuates the need of accurate studies on particulate variability over the Po Valley.

- Local anthropogenic sources. The co-emission of particles and NOx from combustion emissions and the widespread sources of ammonia from agricultural activities over the region (Clarisse et al., 2009) lead to the accumulation of primary carbonaceous particles and secondary inorganic aerosols (ammonium nitrate) in the lower layers of the atmosphere. Crosier et al. (2007), during a campaign in summer 2004, observed that, under easterly flow, ammonium sulphate and organics dominated the sub-micron aerosol particle fraction while, under westerly anticyclonic flow, the large NOx and ammonium emissions at the surface resulted in a large ammonium nitrate concentration in air masses recirculating over the Po Valley. During summer 2012, under similar meteorological features (anticyclonic conditions and PBL air recirculation), Sandrini et al. (2016) observed a significant enhancement of secondary organic and inorganic aerosol particle mass. They also pointed out differences in rural and urban area aerosol behaviour: rural areas, during night, were characterized by higher relative humidity and lower temperature compared to the urban one, and showed higher fine nitrate nocturnal concentration and formation of ammonium nitrate in the large accumulation mode (0.42-1.2 µm). This large amount of highly hydrophilic compounds significantly increase aerosol particle light scattering due to additional water uptake: ammonium nitrate-rich particles are very hydrophilic and, in conjunction with the high relative humidity conditions often encountered in the Po Valley floor, contribute to the build-up of hazes in the region, also in the summer season (Hodas et al., 2014).
Long range natural sources: Saharan dust transport. Mineral dust intrusion episodes can significantly affect public health: epidemiological studies on Italy (Sajani et al. (2010)) revealed increased respiratory mortality in correspondence of such events. During summer months, saharan dust particles are uplifted to the mid-troposphere levels (up to 5 km) by the strong surface winds and the large-scale convection that typically involves north Sahara in this season (Querol et al., 2009a). Mineral dust is then transported over the whole Mediterranean basin, following usually anticyclonic pattern of circulation (Gkikas et al., 2013) triggered by the extended subtropical anticyclone of the Atlantic Azores. Mineral dust over the Mediterranean is found to be usually transported in the lower free troposphere between 2 and 5 km of altitude (Papayannis et al., 2005; Kalivitis et al., 2007); nevertheless, there is indication of episodes of mixing with PBL air (Bonasoni et al., 2004; Perrino et al., 2008; Pederzoli et al., 2010), with a contribution to the surface particulate mass concentration estimated to be of the order of 10 µg m\(^{-3}\). While the above mentioned studies rely on in situ measurements, a direct evidence of mineral dust entrainment in the PBL from continuous vertically resolved observations over the Po Valley is still lacking.

As accumulation of pollution can affect both regional climate and public health (Lelieveld et al., 2002; Kanakidou et al., 2011), a better understanding on the processes contributing to the high concentration of PM on the region is needed. The present paper offers more details on aerosol processes related to both local and long-range sources, and their interaction, in the Po Valley region. The study is based on the analyses of continuous and vertically resolved profiles of particle light scattering and depolarization: in particular, it exploits Light Detection and Ranging o Laser Imaging Detection and Ranging (LiDAR) aerosol characterization, assisted by ground observations and transport models, to identify the origin of the particulate entering and mixing in the PBL during summer season. The LiDAR represents a widely used technique for studying the vertical and temporal distribution of particulate matter optical properties (e.g., Hamonou et al., 1999; Matthias et al., 2002; Dulac and Chazette, 2013; Pappalardo et al., 2004; Amiridis et al., 2005; Papayannis et al., 2005; Flentje et al., 2010b). Coupled with in situ observations and model analysis, LiDAR observations can also be used to derive transport pathways and physical-chemical processes (e.g., Papayannis et al., 2005; Größ et al., 2013). In this work, LiDAR observations are compared with near surface measurement techniques from sites at different altitudes, and supported by Lagrangian model simulations, to provide new insights on the processes that affect aerosol particle variability in the Po Valley and improve the understanding on the possible role of different local and remote sources on the PM level over the region. The focus is on the observation of mineral aerosol in the PBL, on the analysis of its origins and on its mixing and interaction with local aerosol particles and anthropogenic pollution. Please note that, from here on, by "aerosol" we will refer to just the aerosol particle phase, excluding the carrier gas.

2 Observations and methodology

Observations were performed in the framework of the SuperSito project, coordinated by the Regional Agency of Prevention and Environment and funded by Region Emilia-Romagna (ARPA-ER, Italy www.supersito-er.it), and of the FP7 European project PEGASOS (Pan-European Gas-AeroSOl-climate interaction Study, pegasos.iceht.forth.gr).
2.1 Measurement stations

The San Pietro Capofiume (hereafter SPC) station (44°39’ N, 11°37’ 0 E, 11 m a.s.l) is located in the South-Eastern part of the Po Valley, at a flat rural background site relatively close to densely populated cities and industrial sites (i.e. 30 km NE from Bologna urban area, with about 0.5 M inhabitants, and 20 km S from Ferrara, with 0.15 M inhabitants). SPC is included in the list of WMO/GAW regional stations, being representative of the surrounding wider region. The Mt. Cimone (44°12’ N, 10°42’ E, 2165 m) WMO/GAW global station (hereafter CMN), located 100 km south-west from SPC, is instead situated at the top of the highest peak of the Northern Apennines. For the greater part of the year, CMN observations can be considered representative of the background conditions of the South Europe free troposphere (Bonasoni et al., 2000a), while, during warm months, it can be considered at a transition level between the uplifted boundary layer and the free troposphere (Andrews et al., 2011). Under summer anticyclonic conditions, polluted air masses transport from the regional boundary layer can be detected at CMN, due to thermal transport processes and PBL growth (Marinoni et al., 2008; Cristofanelli et al., 2013). CMN also represents the first mountain ridge impacted by Saharan air masses on their way across the Central Mediterranean basin to Europe.

2.2 Aerosol particle light backscattering coefficient profiles: LiDAR observations

The LiDAR system operating in SPC, described in Cairo et al. (2012), uses a 532 nm Nd-YAG pulsed laser source, with pulse duration of 1 ns, 400 µJ of energy and repetition rate of 1 kHz. The optical receiver of the LiDAR is a Newtonian telescope. Taking into account the distance between the telescope and the laser beam, the overlap of the laser within the FOV begins at few tens of meters (around 50 m) from the system, and is complete at around 300 m. Experimental correction allows the reconstruction of the LiDAR backscattering profile down to around 100 m, with an acceptable uncertainty (close to 10 %) on the backscatter ratio precision (see Rosati et al., 2016, and its supplementary material). Profiles are collected every 10 minutes with a vertical resolution of 7.5 m extending, on average, up to 7 km. In the following discussion, we will make use of the aerosol linear depolarization ratio ($\delta_a$) and the aerosol backscattering ratio $Ra = R - 1$ derived from the total backscattering ratio ($R$), defined as (Cairo et al., 1999; Browell et al., 1990):

$$\delta_a = \frac{\beta_{a_{\text{perp}}}}{\beta_{a_{\text{par}}}}$$

$$R(r) = \frac{\beta_a(r) + \beta_m(r)}{\beta_m(r)}$$

and therefore

$$Ra(r) = R(r) - 1 = \frac{\beta_a(r)}{\beta_m(r)}$$

$\beta_m(r)$ represents the backscatter coefficient from molecular contribution while $\beta_{a_{\text{perp}}}$ and $\beta_{a_{\text{par}}}$ are the backscattered signal components from aerosol particle light scattering, with polarization respectively perpendicular and parallel to the polarization of the emitted light. The inversion of the LiDAR signal is accomplished with the Klett method (Klett, 1985; Fernald, 1984),
finding a suitable region of the profile that is supposed to be free of aerosol to calibrate the signal, and using piecewise constant extinction-to backscattering ratio (LiDAR ratio, \(LR\)) values. Using volume depolarization (\(\delta(r)\), see Cairo et al. (2012)) as proxy of aerosol linear depolarization ratio, we select different values for \(LR\) following what reported in literature: highly depolarizing desert dust (\(\delta(r) < 10\%\)) is associated to a \(LR\) equal to 50 sr (Müller et al., 2007) while for low depolarizing aerosol we assume the values typical for anthropogenic aerosol, \(LR\sim 60–70\) sr (Murayama et al., 1999; Ferrare et al., 2001; Fiebig et al., 2002). In addition we considered different values for water cloud (\(LR = 20\) sr) and ice clouds (\(LR = 30\) sr) (Chen et al., 2002; O’Connor et al., 2004). A more detailed description of the methods used to perform the inversion of LiDAR data, together with a through uncertainty analysis, performance in conditions close to the SPC site and additional experimental set-up details, are given in Cairo et al. (2012) and Rosati et al. (2016) and its supplementary material.

2.3 Aerosol particle number size distribution: APSS and OPSS

Aerosol concentrations at the ground are obtained from an Aerodynamic Particle Sizer Spectrometer (Type TSI, APS model 3321), operating at SPC, that provides real-time aerodynamic measurements of particles from 0.5 to 10 \(\mu m\) at 1 minute time resolution. The aerodynamic diameter is defined as the physical diameter that a unit density sphere will have if settles through the air with a velocity equal to the one of the sampled particle. The aerodynamic diameters of particles is established measuring their transit time between two points when accelerated singly through a well-defined flow field. The aerodynamic diameters are here converted to volume-equivalent particle diameters, following Khlystov et al. (2004) and assuming an effective particle density of 1.8, during dust days, and 1.55 during the remaining days, accordingly to the values retrieved by Putaud et al. (2004). The near surface aerosol number concentrations at the free tropospheric level are instead derived from an Optical Particle Size Spectrometer (Type Grimm, OPSS Particle Size Analyser Mod. 1.108) operating at the CMN station. The OPSS provides particle counts in the diameter (\(Dp\)) range of \(0.3\mu m < Dp < 10\mu m\). The instrument is based on the quantification of the 90° scattering of light by aerosol. According to the specifications, the reproducibility of the OPSS in particle counts is \(\pm 2\%\) (Putaud et al., 2004). Such measurements allow the determination of the fine (\(0.3\mu m < Dp < 1\mu m\)) and coarse (\(1\mu m < Dp < 10\mu m\)) aerosol fractions with a 1 minute time resolution. For the purpose of the paper we make use of the time series of coarse (\(Dp > 1\mu m\)) aerosol concentration observed at CMN, without any further correction to the "optical" diameter, to provide a clear indication on the presence of mineral dust layer in the free troposphere (Marinoni et al., 2008; Duchi et al., 2016).

2.4 Chemical Composition: MARGA

The Monitor for AeRosol and GAses (MARGA, Metrohm Applikon B.V. Schiedam NL) is a wet chemistry instrument that provides continuous measurements of the water soluble inorganic gases and aerosol components that might have a direct effect on air quality (Makkonen et al., 2012; Rumsey et al., 2014). The analytical system allows for the characterization of inorganic aerosol (\(Cl^- , NO_3^- , SO_2^{2-} , NH_4^+ , K^+ , Ca^{2+} , Mg^{2+} , Na^+\)) and gases (\(NH_3, HNO_3, SO_2, HONO, HCl\)) at hourly resolution (Twigg et al., 2015). In the sampling box, the air passes through a Wet Rotating Denuder (WRD) (Keuken et al., 1988) where water soluble gases are stripped from the air stream and collected in water. The sampled air then continues through a Steam-Jet Aerosol Collector (SJAC, Khlystov et al., 1995; Slanina et al., 2001) where the water soluble aerosols are
separated from the air stream and collected. The gas and aerosol samples are then analysed by online ion chromatography with high accuracy (detection limits as low as 0.01 µg m\(^{-3}\) (Twigg et al., 2015)). Size-selective particle cyclones are used in front of the two MARGA sampling boxes so that the size of the particles for analysis can be limited to an aerodynamic diameter of less than 10 (PM10) or 1 (PM1) µm.

### 2.5 Transport modelling

We make use of the FLEXPART Lagrangian particle dispersion model (version 9.02)(Stohl et al., 2005, and references therein) to characterize the transport during the campaign period. FLEXPART is a widely used model system to simulate synoptic and mesoscale transport and diffusion of aerosol and trace gases, as well as loss processes such as dry and wet deposition or radioactive decay (Stohl et al., 2005), and has been validated using large-scale tracer experiments (Stohl et al., 1998; Forster et al., 2007). In our case, the model is driven by pressure level data from NCEP Global Forecast System (GFS) (rda.ucar.edu). Meteorological input is provided every 6 hours (00:00, 06:00, 12:00 and 18:00 UTC) at a resolution of 0.5° × 0.5°. FLEXPART output is the footprint of the retroplume, namely the time of residence of the air parcels over each geographical grid point during the 5 days prior to the moment of the trajectories release. Such quantity, expressed in seconds, gives an indication of which emissive regions are going to contribute (and in which extent) to the mineral dust enrichment of the air parcels, influencing therefore dust burden at the time and position of trajectories release. To give an estimate of the variability in the mass of mineral dust advected over SPC, we compute, for each release, the mass fraction of trajectories that encounter dust emissive regions respect to the total mass of the released cluster. Dust is considered to be injected in the atmosphere when the trajectories are crossing the PBL height (as extracted by FLEXPART itself from the GFS input meteorological field, see Stohl et al. (2010)) over the Sahara desert. The backtrajectory clusters were released every 6 hours, along the whole campaign period, from the SPC station in correspondence of the 1000 – 2000 m and 3000 – 4000 m atmospheric layers.

### 3 LiDAR aerosol type classification

LiDAR observations have been extensively used to identify mineral dust layers and discriminate among different typologies of aerosols, based on a choice of specific ranges of optical parameters considered representatives of distinct aerosol types. Examples are shown in Burton et al. (2012), where the classification among eight different types of aerosol is derived from volume depolarization ratio (\(\delta\)), LiDAR ratio (\(LR\)) and the color ratio (\(CR\)). In Größ et al. (2013) the categorized aerosol types are: sea-salt, mineral dust and mixed dust based on \(LR\) and aerosol linear depolarization ratio \(\delta_a\). The estimate of \(LR\) (that requires an independent information on the extinction that should be derived from the Raman signal), as well as an evaluation of the \(CR\) (based on the adoption of two wavelength channel), is not possible with single wavelength elastic LiDAR as the one deployed at SPC during the PEGASOS campaign. Nevertheless, some typologies of aerosol show distinct values of aerosol linear depolarization ratio. At 532 nm, values of \(\delta_a\) around or higher than 30 % are generally associated with layers of nearly pure mineral dust while smaller values (around 8–10 %) are often detected in correspondence of mixture of mineral dust and
non-depolarizing particles (Murayama et al., 2003; Sugimoto et al., 2006; Tesche et al., 2009a). By contrast, smoke and other anthropogenic aerosols exhibit low values of $\delta_a$ (less than 5 %)(Sun et al., 2012; Größ et al., 2013).

Here we implement a three-types aerosol discrimination scheme, based on the different statistical distribution of optical properties of each class (see Fig. 1), to characterize the vertical and temporal aerosol variability over the region along the campaign period (15 June 2012 – 5 July 2012). The reader should notice that the lower $\delta_a$ values that we observe with respect to what usually found in literature (especially for the dust layers) are likely linked to the calibration process, and in particular to the difficulty in individuating completely aerosol free layers in the vertical span of the adopted LiDAR system (from the ground to 7 Km). In this work, $\delta$ has been calibrated following the "0° calibration" or the "atmospheric calibration" procedure, i.e. making use of a low aerosol height range in the LiDAR signal, where only the molecular contribution could be considered. There, the volume depolarization ratio has been forced to assume the well-known value of the air molecule linear depolarization ratio (Behrendt and Nakamura, 2002). We acknowledge that this calibration is unsatisfactory to produce quantitative results, as the possible residual presence of small amounts of highly depolarizing aerosol, in the assumed clean range, can easily compress the range of variability of the volume depolarization, and underestimate the final depolarization products (Freudenthaler et al., 2009, 2016). However this possible source of inaccuracy does not compromise the purpose of this work. The LiDAR classification, based on the statistical distribution of the overall observed $\delta_a$ and $Ra$ values, is in fact applied here to overcome such limitations. The robustness of the results are then further supported by comparison with lagrangian analysis and in situ measurements.

Figure 1 reports the probability density function, along the whole measurements campaign, of the aerosol occurrence, expressed as a function of $Ra/(Ra + 1)$ (ranging from 0 in aerosol free condition, i.e. $Ra = 0$, to 1 in presence of an opaque aerosol layer, when $Ra$ tends to infinity) and $\delta_a$. The analysis includes 10 minutes resolution observations from the ground up to 5000 m height, for a total of about $1.5 \cdot 10^7$ sampling points. The different aerosol classes can be discerned in three distinct patterns:

1. $0.1 < Ra/(Ra + 1) < 0.8$ and low values of $\delta_a$ (< 3 %); such low values of aerosol linear depolarization ratio are indicative of spherical particles. These particles may be composed of anthropogenic pollution and, for higher values of $Ra$, by droplets, and are defined as non-depolarizing.

2. $0.3 < Ra/(Ra + 1) < 0.7$ and high values of $\delta_a$ (> 10 %); In this class we find the highest values of aerosol linear depolarization ratio (mainly ranging from 10% to 20%) and this can be indicative of the presence of mineral dust particles. This class is defined as depolarizing.

3. $0.2 < Ra/(Ra + 1) < 0.6$ and intermediate $\delta_a$ values (3 % $< \delta_a < 10$ %) which, based solely on $Ra$ and $\delta_a$, cannot be considered as indicative of a dominance of a defined aerosol type unless coupled to a more thorough correlation with additional observations. We will refer to this type as intermediate depolarizing.

The boxes in Fig. 1 delimit the $Ra/(Ra + 1)$ and $\delta_a$ ranges for each of the three classes used to derive the aerosol mask for the whole campaign. The results are reported in Fig. 2 together with the profiles of $Ra/(Ra + 1)$ and aerosol linear depolarization...
ratio $\delta_a$. Overall, non-depolarizing particles (type 1) are dominant throughout the campaign with a total occurrence of 49% of the measurements in the 100-5000 m range. They are observed prevalently below 2000 m height, and are associated with enhanced values of $Ra$ ($Ra/(Ra+1)$ ranging between 0.3 and 0.6). Due to the presence of such particles, the vertical gradient of $Ra$ marks, when not masked by the presence of mineral dust layers or clouds, the PBL evolution. Two events of depolarizing aerosol (19 June – 21 June and 29 June – 02 July), recognized as type 2 and likely related to mineral dust presence, are observed between 2000 m and 5000 m. Such events are also clearly visible as enhancement of $Ra$ in the free troposphere with values of $Ra/(Ra+1)$ ranging from 0.6 and 0.8. The intermediate class (detected with an occurrence of around 19% during the whole campaign) is found in close proximity of the depolarizing aerosol and within the PBL. The observed vertical and time distribution of these intermediate type particles indicates the possibility of mixing of the dust depolarizing layers with local non-depolarizing particulate. Nevertheless, intermediate $\delta_a$ values are also observed systematically after 12:00 UTC (Universal Time Coordinate), between 0 and 1500 m height, for the majority of dust-free days. The nature of such intermediate non-dust depolarizing aerosol is further discussed in Sect. 6.

4 Meteorology and synoptic aerosol regimes

The evolution of the meteorological conditions at SPC is reported in Fig. 3. Vertical profiles from the ground up to 4 km height of wind speed and direction come from radiosondes (Vaisala RS92) launched daily at 00:00, 06:00 and 12:00 UTC. Panel c of Fig. 3 reports also the evolution of ground temperature at 12:00 UTC. PBL evolution description is referred to the PBL height time series presented in Sandrini et al. (2016) and reported also in the supplementary material (see Fig. S1). The observation of wind profiles over SPC highlights a sequence of distinct meteorological regimes:

- Stagnation period: (15 June – 19 June) This first phase is characterised by a situation of stagnant conditions (wind speed less than 4 m s$^{-1}$ below 2000 m height) with a progressive warming of the air masses (from 29 °C up to 34 °C at the ground). The PBL top is limited below 1500 m until 19th June when it reaches 2000 m.

- South-westerly winds: (20 June – 21 June) During this phase higher wind speeds (between 11 m s$^{-1}$ and 16 m s$^{-1}$) are observed above 2000 m. Correspondingly, wind direction profiles indicate a prevalent South and South-West provenience. The arrival of warm Saharan air masses (temperature at the ground around 32°C–33°C) leads to a more intense PBL development (up to 2000 m) respect to the previous stagnation phase. (30 June – 5 July) During the last days of the campaign, strong winds (between 12 and 15 m s$^{-1}$ with a peak of 20 m s$^{-1}$ on 3 July) are observed above 1500 m. While during 30 June winds are coming mainly from South, the following days are characterized by a steering to south-westerly (1 July) and then westerly flow (2 and 3 July). During this phase, temperature at the ground reaches its highest values (35°C–37°C) but the dust layer presence made it difficult to unambiguously retrieve the PBL top. During the immediately following days (3–4 July), the PBL top was detected above 2000 m.

- North-easterly winds: (22 June – 29 June) The radiosounding profiles indicate a prevalence of south-easterly or easterly winds above 2000 m a.g.l. Northerly/north-easterly winds are instead often visible at lower altitudes, in particular
between 23 and 24 June below 1000 m, on 26 June below 500 m (associated also to wind intensities up to 10 m s\(^{-1}\)) and on 27 June between 500 m and 1800 m. During this period, ground temperature first decreases to 28 °C then increases again after 26 June, reaching 35 °C on 29 June. The PBL maximum height varies between 1500 m and 2000 m a.g.l.. Such conditions are favourable for the export of the Po Valley pollution toward the Tyrrhenian Sea and will be extensively discussed in a companion paper.

The evolution of the observed size distribution and optical classification during the distinct meteorological regimes is presented in the following sections.

4.1 Summer stagnant conditions: 15–19 June

Meteorological evolution is compared with the aerosol optical variability from LiDAR (see Fig. 2) and with ground aerosol number concentration and volume size distribution (estimated as the volume of a sphere with diameter corresponding to the volume-equivalent particle diameter) at SPC and CMN (see Fig. 4). Panel a of Fig. 4 shows the time trend of small particle concentration (297 nm < \(D_p\) < 420 nm) from the APSS at SPC. The stagnation period (15–19 June), typical for the Po Valley on hot summer days (Rossa et al., 2012), is characterized by a marked daily cycle in the aerosol concentration and by a progressive day-by-day accumulation of particles in the PBL. This is noticeable in the increase of the particle number size distribution of small particles from APSS (from 5 cm\(^{-3}\) to nearly 20 cm\(^{-3}\)) during the early morning hours (00:00-06:00 UTC), when the lower troposphere is stably stratified. The LiDAR observations (Fig. 2) show a persistent layer of non-depolarizing particles up to 2 km height, attributable to anthropogenic aerosol and modulated vertically by the PBL daily cycle. APSS data in panel b of Fig. 4 show a bimodal aerosol distribution with a clear increase of volume mode due to submicron particles (0.5 µm < \(D_p\) <1 µm) growing from 0.4 µm\(^3\)cm\(^{-3}\) to more than 1 µm\(^3\) cm\(^{-3}\). During this period of the campaign, the OPSS at CMN (panel c Fig. 4) indicates coarse particle number concentrations below 0.4 cm\(^{-3}\).

4.2 Saharan dust events: 19–21 June and 29 June–3 July

During the first event (19 to 21 June) strong south-westerly winds (with speeds greater than 10 m s\(^{-1}\) above 2000 m height) are associated with a stable anticyclonic circulation centred above South Mediterranean and Tunisia, leading to an efficient south/south-westerly circulation. Mineral dust can be clearly observed as an enhancement in LiDAR \(Ra\) profiles (Fig. 2, panel b) above 2000 m until June 21 while, below that height, it is not possible to discern any deviation from the background aerosol signal. The enhancement in \(Ra\) is accompanied with increased aerosol linear depolarization ratio (\(\delta_a\) ~ 10\% – 15\%) during the whole event, with values up to 20 \% above 3000 m during 20 June, resulting in a coherent layer of type 2 particles visible in the aerosol mask (panel a of Fig. 2). In correspondence of the presence of dust aerosol at 2000 m level, the OPSS at CMN (panel c, Fig. 4) detects an increase of coarse particle concentration up to 1.8 cm\(^{-3}\). The peak seen on 21 June at around 09:00 UTC (greater than 5 particles cm\(^{-3}\)) should be attributed to an enhancement in aerosol load, that can be both caused by an intensification of mineral dust burden or by mixing with pollution from the regional PBL (Cristofanelli et al., 2009), as seem to be suggested by the observed corresponding increase in black carbon concentration at CMN (see also Cristofanelli et al.
Intermediate depolarization aerosol type 3 is observed below the depolarizing layer, throughout the mineral dust event, and persisting until 22–23 June. As mentioned in the previous section, this can be a signature of mineral dust mixing with local non-depolarizing particulate. A direct comparison of the aerosol layer structure with the in situ measurement is presented in Fig. 5, focusing on the days of dust transport. From the comparison is possible indeed to observe a simultaneous enhancement of coarse (2 μm < Dp < 5 μm) particles detection by the APSS on 20-22 June (panel a of Fig. 5) in correspondence of the identification of type 3 class near the ground, suggesting mineral dust presence in the layer composition. It’s worth to notice that, while ground measurements do not indicate a clear coarse particle enhancement after 22 June, the LiDAR still observes a lofted layer of intermediate depolarizing aerosol until 23 June. During the second event (29 June to 3 July) high wind speeds above 2000 m (up to 20 m s⁻¹) are associated with a high-pressure area centred above central Italy, leading again to favourable south-westerly circulation. LiDAR data show a second layer of enhanced Ra (Ra/(Ra + 1) around 0.6) lasting from 28 June at 23:00 UTC to July 3 at 00:00 UTC. Depolarization reaches values higher than in the previous event (with mean values of 15 % and maximum exceeding 20 %); this is again visible as a thick and persistent layer of type 2 aerosol that, in this case, extends down to the ground on 1 July (see panel b of Fig. 5). As in the previous case, it is possible to observe the presence of intermediate depolarization particles (type 3) close to the depolarizing layer (type 2). The dust layer appears to be characterized by a more intense contribution of coarse particles respect to the previous event, visible both at lofted level and ground. APSS observations in fact show an increase in coarse particle volume simultaneously to detection of type 2 and 3 particles close to the ground, with values higher than in the previous event (>1 μm³ cm⁻³) and increased contribution from the intermediate particle size (1 μm – 2 μm). Similarly, in correspondence of the presence of the depolarizing layer at 2000 m, the OPC (panel c, Fig. 4) shows coarse particle concentrations nearly doubled compared to the previous event (concentration between 2 and 3 particles cm⁻³).

The upper panels of Fig. 6 shows the footprints of the 5 days retro-plumes released on 20 June at 18:00 UTC (panel a) and on 29 June at 12:00 UTC (panel b). The transport for the first event has a more direct pathway, with an average transport time of 2 days from North Sahara. The second event appears instead to originate from Western Sahara and has a longer pathway revolving around the anticyclonic circulation (around 4 days). Aerosol Optical Depth from multi-model forecasts (SDS-WAS Sand and Dust Storm WMO warning advisory and assessment system, visible at http://sds-was.aemet.es/forecast-products/dust-forecasts/compared-dust-forecasts) indicates a spatial distribution in agreement with the FLEXPART footprints for the two events. The simulated mineral dust mass fraction over the SPC site (panel c, Fig. 6), allows providing an estimate of the evolution (with a time step of 6 hours) of the mineral dust contribution on the SPC site. The simulation confirms the presence of the two desert dust transport periods and the progressive descent of the dust layers advection from 3000–4000 m to 1000–2000 m heights, simultaneously to what shown by LiDAR observations. The maximum mineral aerosol fraction from FLEXPART analysis occurs on June 20, both at the upper layer (9%) and at the bottom layer (9% also). According to FLEXPART, the import of mineral dust at the lower layer persists until the morning of 23 June, when dust presence is not unambiguously inferable from observations but the aerosol mask still indicates the presence of intermediate depolarizing particles below 2000 m. The second desert dust event predicted by FLEXPART again shows the same timing with respect to observation and also confirms the presence of a thick layer of dust that involves at the same time the 1000–2000 m and
3000–4000 m layers. The estimated mass fraction contribution on the contrary, especially in the lower layer (between 2 and 4 \%) appears inferior respect to the previous events. It should be emphasized that a quantitative assessment of the mineral aerosol transport from this method would be difficult, due to uncertainties related to the model estimate of PBL height over the desert, to the variability in the emissions of mineral dust and to the uncertainties on the trajectories dynamics (Stohl et al., 1998). Nonetheless, despite such limitations, the model offers a robust characterization of the dynamics and timing of the events, supporting the interpretation of the data analysis.

5 Mean daily variability

Figure 7 reports the frequency of observations for each of the three classes, integrated for the period 15 June – 5 July. Depolarizing aerosol (panel a, Fig. 7) are associated with the two events of desert dust and hence are mostly observed between 1500 and 5000 m height with a frequency of occurrence ranging between 15 \% and 30 \%. Non-negligible occurrences (\sim 10 \%) are also observed close to the ground and can be attributed to the mineral dust descent during the second event. Non-depolarizing aerosol (panel c, Fig. 7) is dominant throughout the campaign (occurrence up to 80 \% below 2000 m); This class of aerosol appears to be mainly confined below the PBL top (derived by the analysis shown in Fig. S1 of the supplementary material and traced with a black dashed line). During the campaign, diurnal PBL starts to develop on average at 06:00 UTC and reaches its maximum vertical extension, up to 2 km height, between 17:00 and 18:00 UTC. A high occurrence of non-depolarizing particles marks the PBL average daily evolution both during the diurnal formation and at night-time (21:00 to 05:00 UTC), forming the residual layer. A clear minimum of non-depolarizing particles occurrence is observed in the afternoon between 16:00 and 19:00 UTC, when LiDAR indicates instead a maximum (50 \% of observation) of intermediate depolarization type occurrence (panel b, Fig. 7). Such enhancement of type 3 particles detection during late afternoon appears frequently along the campaign (clearly visible on 13 days over 21, see Fig. 2). It should be noticed that on 20–22 June and 30 June–2 July the presence of mineral dust can mask any \( \delta_d \) enhancement in the PBL. On average the vertical extent of each layer of intermediate depolarization is limited within the PBL below 1500 m height (panel b, Fig. 7). Daily average evolution of particle volume size distribution, relatively to dust-free days, is reported in Fig. 8. Fine particles \((Dp < 1 \mu m)\) volume shows a semidiurnal cycle, corresponding to the diurnal cycle of non-depolarizing aerosol near the ground, with concentration increasing during the stable nocturnal layer phase (late night - early morning) and strongly decreasing during the stage of well-developed PBL. The larger particle mode shows two maxima: a first one (volumes < 0.4 \( \mu m^3 cm^{-3} \)), in correspondence to the uplift of the PBL layer around 9:00 UTC, and a second one forming at 15:00 UTC, with a maximum (volumes > 0.5 \( \mu m^3 cm^{-3} \)) at 20:00 UTC, showing a similar timing than the depolarization enhancement described above. Further analysis of the afternoon PBL aerosol composition is reported in the following section.
6 Non desert dust depolarising aerosol

We report in Fig. 9 the $\delta_a$ profiles of a representative case study (3 July) of the late afternoon occurrence of intermediate depolarizing aerosol. The aerosol linear depolarization ratio indicates that the plume starts to develop from 15:00 UTC to 20:00 UTC and reaches the maximum height of 1500 m in the late evening, with a vertical structure suggesting a possible uplift of particles from the ground. An increase in aerosol linear depolarization ratio in a regime of convective PBL was already observed by Gibert et al. (2007a). Their results show a positive correlation of enhanced $\delta_a$ with an increase in vertical wind velocity, possibly indicating a source emission of particles transported upward by convection. The actual nature of the aerosol plume cannot be assessed solely by LiDAR depolarization. An increase in depolarization can be due to the presence of irregularly shaped particles that can belong to a wide range of aerosol types, from soil and desert dust to marine aerosol (Murayama et al., 1999) and ash particles (Nisantzi et al., 2014). The hourly time resolution measurements of PM1 and PM10 aerosol chemical compositions, provided by the MARGA analyser, show no evident correlation between the depolarization increase and the presence of sea salt (not reported here). Similarly, no correlation was found with absorbing aerosol (black carbon), investigated by means of a multi-angle absorption photometer (MAAP, Petzold et al., 2006b) (also not shown). By contrast, MARGA observations highlight a maximum in PM10 calcium concentrations, simultaneously to the afternoon increase in ground depolarization (starting between 15:00 and 20:00 UTC, see panel b of Fig. 7) and in the larger particles detection from APSS (maxima between 18:00 and 20:00 UTC, Fig. 8). The diurnal mean evolution of the calcium ion ($\text{Ca}^{2+}$) fraction, calculated over the total PM10 ions, shows a marked increase after 10:00 UTC with a maximum in the late afternoon (17:00 UTC – 20:00 UTC, up to 0.35, see panel a, Fig 10). This daily behaviour is in agreement with the enhancement in aerosol volume contribution from large particles respect to the fine ones, shown in panel b of Fig. 10. These results reinforce the hypothesis of the crustal origin of the intermediate depolarizing particles observed by the LiDAR. It is possible therefore to explain, at least on a qualitative basis, the recurrent detection of the afternoon aerosol plumes as emissions and resuspension of soil particles from dried land sources. The frequent occurrence of such events during the observational campaign indicates that the Po Valley can effectively act as a source of mineral particles, likely originated from agricultural soils, that under convective atmospheric conditions can be uplifted at the PBL top in late afternoon hours. This is further confirmed by the diurnal evolution of non-desert dust coarse particle concentration at CMN (panel c, Fig. 10), that indicates an enhancement in the coarse particle fraction in late afternoon/evening. Hence, recurrent vertical transport from the Po Valley, triggered by thermal air-mass, can uplift mineral aerosol over the mountains ridge, and potentially impact on particulate transport up to the regional scale (Cristofanelli et al., 2016).

7 Effect of aerosol hygroscopic growth on scattering and depolarization

LiDAR data (Fig. 2) frequently show, during early morning hours, a shallow layer of non-depolarizing aerosol below 300 m height, more easily visible during days characterized by desert dust and mixed dust events (see for instance 00:00-06:00 UTC of 19 June and between 00:00-08:00 UTC of 30 June). The decrease in depolarization is less evident in dust-free atmosphere but is nevertheless observed in several other days of the campaign (18 June, 21 June, 22 June, 29 June, 1 July, 4 July and 5 July),
always below 300 m, before 08:00 UTC and usually associated with high values of $Ra (Ra/(Ra + 1) > 0.6)$. The average profiles of $\delta_a$, relative humidity ($RH$) and potential temperature ($TH$) for dust-free and dust days at 05:00 UTC (selected among the days of dust presence at the ground: 20–22 June and 30 June – 2 July) are reported in Fig. 11. During dust-free days (Fig. 11, panel a), values of $\delta_a$ stay on average below 1% in the upper layer (above 2000 m) and below 2% in the lower layer (below 2000 m). Starting from 800 m, RH increases with decreasing altitude and this is associated with a progressive decrease in the $\delta_a$ values. During dust days (Fig. 11, panel b) a more defined stratification of the atmosphere is observed. This is visible also in the profiles of $TH$ that, compared to the dust-free days, shows a clear passage from a stable layer below 500 m to a more unstable one in correspondence of the depolarizing aerosol. The mineral dust plume is visible between 1500 m and 4000 m height, associated with a layer of higher depolarization ($\delta_a > 10\%$) and drier air respect to the lowermost layer (with $RH$ increasing with height from 40 to 60%). Below 1500 m height, RH varies between 55% and 60% and $\delta_a$ shows lower values compatible with dust presence mixed with local pollution (around 6%). Conversely, the lowermost troposphere (below 500 m), marked by the increased stability shown by the $TH$ profile, is characterized by a sharp decrease in $\delta_a$ (from 7% down to 3%) associated with the increase in $RH$ (from 50% to 75%). The study indicates in both cases a decrease of the depolarization in the lower layers for $RH > 60\%$. During dust-free days the affected aerosol layer (below 400 m) shows a depolarization decrease of about 1% (from an average value of 2% to values around 1% and less). During dust days the process influences the aerosol layers at different levels: for RH values between 60% and 65%, depolarization values decrease is around 2% (from more than 7% to around 5.5%). Closest to the ground, the effect is more evident with increasing RH (up to $\sim 70\%$), corresponding with a decrease of depolarization up to 3.5%. This low depolarization near ground suggests the presence of increasingly spherical particles, which can be originated by two different processes:

- the presence of fine particles of anthropogenic origin that may deliquesce: The stagnant meteorological conditions that characterize the Po Valley during anticyclonic phases are favourable for the formation of secondary inorganic aerosols (especially ammonium nitrate) and of secondary organic aerosol (Sandrini et al., 2016). A recent study (Hodas et al., 2014) showed that, during the same 2012 campaign at SPC, the aerosol liquid water was mainly driven by locally formed nitrate, hence the growth of spherical non-depolarizing aerosol could occur due to deliquescence of fine particles of anthropogenic origin, of which nitrates were the dominant compound.

- Mechanisms explaining the increase of scattering of mineral dust particles, along with a reduction of their depolarization ratio, can also be hypothesized (Ikegami et al., 1993; Murayama et al., 1999; Sassen et al., 2002; Zhou et al., 2002; Nee et al., 2007). It should be emphasized that, during the analysed case study, high relative humidity values (80% or more) are observed in the lowermost non-depolarizing layer (see Fig. 11), suggesting that condensation of water around mineral dust particles coated with (or simply enriched of) hydrophilic components may play a role in the modification of the optical properties of desert dust in this atmospheric layer. Indeed, even if mineral dust is primarily a hydrophobic aerosol, it can become hydrophilic due to chemical reactions occurring on the particle surfaces during long-range transport (Nee et al., 2007; Sullivan et al., 2009a) or locally from the condensation of inorganic and organic soluble materials from ground sources.
During the summer 2012 campaign, under the observed conditions, both processes may have played a relevant role: Fig. 12 shows that, during the stagnation phase (from 14 June until 19 June), aerosol nitrate (NO3-) concentration in both fine and large particles MARGA channels (PM1 and PM10) increases with a marked daily cycle peaking at night. The submicron fraction of nitrate dominates the concentration of PM10 nitrate during this phase. The APSS submicron particle volume concentration follows a daily variability and a buildup similar to that shown by the APSS nitrates concentration, reaching maxima during 19 June (APSS volume concentration up to 25 µm m$^{-3}$ and NO3- PM1 and PM10 concentration up to 15 µg m$^{-3}$ and 18 µg m$^{-3}$, respectively). Such increase, evident during early morning hours in small aerosol from APSS and in PM1 nitrate from MARGA, supports the hypothesis of anthropogenic fine particles deliquescence, explaining therefore the low values of $\delta_a$ observed by the LiDAR in the surface layer during dust-free days. Similarly, in presence of dust, the low depolarization values can be related to external mixing of dust depolarizing particles with such locally formed spherical particles. Nevertheless, after the end of the stagnation phase (19 June) the aerosol nitrate concentration decreased and, during the observed desert dust episodes, the difference between the nitrate PM10 and PM1 fractions became more evident, with PM10 prevailing over PM1. The intensified ventilation established after 19 June may in fact have limited the accumulation of anthropogenic particles, at the same time carrying dryer African air masses and making nitric acid condensation on coarse particles prevails over condensation on accumulation mode aerosol. Consequently coarse-mode nitrate would promote water condensation on the large particles, leading to low aerosol linear depolarization ratio values even in presence of dust.

8 Conclusions

The presented paper provided a characterization of the effects of meteorological evolution and transport patterns on the aerosol variability, based on the observations collected during two major field campaigns (PEGASOS and SuperSito) in the eastern part of the Po Valley. The aim was to contribute to the understanding of the processes that lead to the high concentration and variability of aerosol characterizing the Po Valley during typical summer conditions.

The analysis of meteorological conditions, coupled with observations from LiDAR and in situ aerosol number/size distribution spectrometers, led to the identification of distinct meteorological regimes with a temporal and spatial distribution of different aerosol types.

We identified a first phase (15–18 June), characterized by a stagnation period (weak winds below 2000 m), representative of hot and polluted conditions in the Po Valley whole area, with progressive accumulation of locally emitted aerosol in the lower troposphere and consequent increase of the fine mode aerosol concentration near the ground. Particle concentration at the ground showed therefore a clear diurnal cycle with maxima during the early morning, when PBL uplift and vertical mixing were absent.

Observations and Lagrangian analysis allowed then a detailed description of two events of Saharan dust transport (in line with the average occurrence of 2–3 summer desert dust episodes over the region detected by satellite (Gkikas et al., 2013)). Mineral dust layers were advected over the measurement site from the Sahara desert, travelling along anticyclonic patterns at high level (around 3000–4000 m) and carrying depolarizing aerosol. The study offered evidence of dust transport to the
ground, showing clear dust layers intrusion in the PBL and rapid mixing with local pollution. We showed how this mixed
layer, generally characterized by lower depolarization values, can reach the ground within few hours and we showed, by direct
comparison with ground in situ instruments, the corresponding enhancement of particle volume size distribution in the 2-5
µm range (leading to values higher than 1 µm^3 cm^{-3}). In both events the plumes descended indeed to low height (with a total
occurrence of depolarizing aerosol identification inside the PBL of ~7 % along the whole campaign). As, on a climatological
basis, Saharan dust advection occurs with noticeable frequency over Northern Mediterranean (i.e. Pey et al. (2013) indicated a
frequency of 17 % for the 2001–2011 period), dust intrusion can represent a significant factor in increasing PM concentration
at the ground. Such results give direct evidence to the suggestion of Bonasoni et al. (2004) that hypothesized, based on in situ
measurements in North Italy and back-trajectories analysis, that mineral dust events detected in free troposphere can lead, with
non-negligible frequency, to PM10 exceedance at the ground in the time span of some hours.

The study revealed moreover the presence of events of late afternoon particles resuspension from the soil, not related to
Saharan dust transport, impacting on the PM concentration near the ground. The existence of a contribution to PM10 levels
from resuspension aerosol sources in European regions was already hypothesized by Vautard et al. (2005), based on chemical
transport model study. Here, several events of intermediate depolarizing aerosol (mean diurnal frequency of detection ~22 %),
up to 2000 m height, were observed during late afternoon (17:00–20:00) in dust-free days. The concurrent increase in calcium
particle spectroscopic measurements (with a contribution up to 0.35 of the total PM10 fraction) indicated the crustal nature of
such aerosol, and can therefore be reasonably attributed to processes of vertical uplift of soil particles, likely related to regional
activities (i.e. farming or combustion processes). The vertical extension of such plume, as observed in LiDAR profiles and in
the diurnal variability in the CMN measurements, suggests also that local pollution can be transported above mountains peak
and hence potentially exported outside the orographic boundaries of the region.

The combination of depolarization profiles with meteorological and aerosol measurements also allowed highlighting the
effects of the condition of high RH (typical for this region) on the particle processes. The analysis revealed how, in conditions
of high relative humidity values (RH > 60 %) in a shallow layer near the ground (<500 m), the aerosol linear depolarization
ratio decreases respect to the above atmospheric layer. Such effect is particularly visible when mineral dust particles are present
near the ground. During this period, the temporal evolution and the high values of nitrates ion concentration in the PM1 and
PM10 channels suggest that the origin of such low depolarization particles can be related to processes of secondary organic
aerosol formation and hygroscopic growth on mineral dust particles with a nitrate-enriched surface.

In conclusion, the in-depth analysis of the aerosol light backscattering profiles provided new insights on particles behaviour
from the ground up to free troposphere. Results pointed out particles processes, observed relatively frequently on the period
of the campaign, that impact aerosol variability, air quality and potentially regional climate, and that deserve therefore more
extended analysis from longer-period vertical resolved observations (i.e. EARLINET network). The detailed retrieved infor-
mation (vertical stratification, hygroscopic growth near ground, aerosol evolution inside the PBL) can also support larger scale
studies. As an example, we cite here a recent study, based on MAIAC satellite information (Arvani et al. , 2016), that tries to
assess a method for surface PM retrieval from space observations relying on rough approximations of PBL evolution and RH
effect on aerosols. Accurate studies on aspects as the ones we presented here may, therefore, represent an important contribution to the improvement of more complex and focused atmospheric observation techniques.

*Competing interests.* The authors declare that they have no conflict of interest.

*Acknowledgements.* This work was partly funded by the projects PEGASOS (FP7-ENV-2010-265148), the project SuperSito by the Emilia-Romagna region (DRG no. 428/10), the EU project StratoClim (grant agreement no. 603557), the EU FP7 grants ÉCLAIRE (grant 282910) and the project of National Interest NextData. We would like to acknowledge the Energy Research Centre for the Netherlands (ECN) to provide us with a MARGA instrument to use during these campaigns. This study received fundings also from the FP7 project BACCHUS (grant agreement 603445).
References


5
Khlystov, A., Stanier, C., and Pandis S. N.: An algorithm for combining electrical mobility and aerodynamic size distributions data when
10
aerosol measurements at a Finnish urban site: comparisons with filters, nitrogen in aerosol and gas phases, and aerosol acidity, Atmos.
Marinoni, A., P. Cristofanelli, F. Calzolari, F. Roccato, U. Bonafé, and P. Bonasoni: Continuous measurements of aerosol physical parameters
20
Matassoni L, Pratesi G, Centioli D, Cadoni F, Malesani P, Caricchia AM, Di Menno di Bucchianico, A.: Saharan dust episodes in Italy:
,Chaykovski, A., Chourdakis, G., Comeron, A., Delaval, A., De Tomasi, F., Eixmann, R., Frioud, M., Hagard, A., Iarlori, M., Komguem, L., Kreipl,
25
Monks, P., Granier, C., Fuzzi, S., Stohl, A., Williams, M., Akimoto, H., Amann, M., Baklanov, A., Baltensperger, U., Bey, I., Blake, N., Blake,
R., Carslaw, K., Cooper, O., Dentener, F., Fowler, D., Fragkou, E., Frost, G., Generoso, S., Ginoux, P., Grewe, V., Guen- 25 ther, A., Hans-
son, H., Henne, S., Hjorth, J., Hofzumahaus, A., Huntrieser, H., Isaksen, I., Jenkin, M., Kaiser, J., Kanakidou, M., Klimont, Z., Kulmala,
M., Laj, P., Lawrence, M., Lee, J., Liouesse, C., Maione, M., McFiggans, G., Metzger, A., Mieville, A., Moussiopoulos, N., Orlando, J.,
O’Dowd, C., Palmer, P., Parrish, D., Petzold, A., Platt, U., Pschl, U., Prvt, A., Reeves, C., Reimann, S., Rudich, Y., Sellegr, K., Stein-


Sassen, K.: Polarization LiDAR evidence at FARS for indirect climate forcing from asian dust storms, LiDAR remote sensing in atmospheric and earth sciences 21th conference, 483-484., 2002.


physical characteristics of particulate matter at kerbside, urban, rural and background sites in Europe, Atmos. Environ., 38, 2561-2577, 2004.


Figure 1. Probability Density Function of aerosol particle optical properties over 15 June – 5 July 2012 as a function of $Ra/(Ra+1)$ and $\delta_a$ parameters. The color code indicates the frequency of occurrence.
Figure 2. LiDAR observations along the whole campaign. Panel a: vertical profiles of aerosol particle types resulting from the classification methodology described by Sec. 3: not depolarizing aerosol (yellow), depolarizing aerosol (orange) and intermediate depolarization aerosol (brown). Panel b: vertical profiles of $Ra/(Ra + 1)$. Panel c: vertical profiles of $\delta_a$. 
Figure 3. Panel a: wind provenience direction. Panel b: wind speed. Panel c: Ground temperature at 12:00 UTC
Figure 4. Panel a: APSS particle counts at 297–420 nm of diameter. Panel b: time series of the volume size distribution of aerosol particles as observed by the APSS. The y-axis indicates the volume-equivalent particle diameters in µm while colours report the corresponding volume concentration. Panel c: time series of coarse ($D_p > 1\mu m$) particle number concentration observed at CMN.
Figure 5. Vertical profiles of aerosol particle types (upper panel), as in Fig. 2, and APSS aerosol time series of the volume size distribution of aerosol particles (lower panel), as in Fig. 4, for the first (panel a) and the second (panel b) dust event.
Figure 6. FLEXPART backtrajectories over GFS meteorological input: Panels a and b show the footprint (in ns of residence over each bin) of the trajectories released over SPC at 3000 m. Black triangle indicates the point of release, black squares mark the position of the center of mass every 24 hours. The pattern of trajectories released on 20 June at 18:00 UTC are shown on the left (panel a), while pattern released on 29 June at 12:00 UTC are on the right (panel b). The simulated mass fraction contribution of dust over the SPC site is reported in panel c with a time step of 6 hours. The black line is relative to the particles released at the 1000 – 2000 m atmospheric layer and the blue line to the release at 3000 – 4000 m.
Figure 7. Mean diurnal frequency of the vertical distribution of each aerosol class (computed as the ratio between the number of aerosol class detections and the number of days of measurements): Depolarizing (panel a), Intermediate Depolarizing (panel b) and Non-Depolarizing (panel c). The mean PBL height, derived from LiDAR analysis, is reported in black dashed line over the non-depolarizing particles distribution.
Figure 8. Mean diurnal evolution of aerosol particle volume size distribution in dust-free days from the APSS at SPC
Figure 9. Vertical profiles of LiDAR aerosol linear depolarization ratio on 3 July 2012.
Figure 10. Panel a shows the mean diurnal evolution of the ratio of PM10 concentration of the Calcium ion (Ca\textsuperscript{2+}) over the total PM10 ion concentration (Ca\textsuperscript{2+PM10}/TotalPM10) in SPC. Panel b reports diurnal mean of the ratio of large particles (1 \textmu m < Dp < 5.5 \textmu m) over the fine ones (0.5 \textmu m < Dp < 1 \textmu m) at SPC while panel c shows the diurnal mean of coarse (Dp > 5 \textmu m) particles at CMN. Each mean is computed on dust-free days.
Figure 11. Figure reports the mean vertical profiles of $\delta_a$, RH and TH for dust-free (panel a) and dust days (panel b) at 05:00 UTC. Please notice the different horizontal axis for $\delta_a$. 
Figure 12. APSS fine particles ($D_p < 1 \mu m$) volume contribution (panel a) compared to nitrates ion concentration ($NO_3^-$) both in the PM1 (black) and PM10 (blue) channel (panel b). Zero values corresponds to missing observations.