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# Study of aerosol microphysical properties profiles retrieved from ground-based remote sensing and aircraft in-situ measurements during a Saharan dust event

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Received: 18 August 2015 - Accepted: 20 August 2015 - Published: 9 September 2015

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Published by Copernicus Publications on behalf of the European Geosciences Union.

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ferences in the volume concentration profiles, although somewhat larger, are still within

Mineral dust is estimated to be the most abundant aerosol type in the atmosphere (~ half of the total global aerosol burden) (e.g. Textor et al., 2007; Choobari et al., 2014), with global emission between 1000 and 3000 Mtyr<sup>-1</sup> (Zender et al., 2003, 2004; Shao et al., 2011). Mineral dust directly scatters and absorbs solar and infrared radiation (Miller and Tegen, 1998), and impacts the optical properties of clouds (Ferek et al., 2000; Rosenfeld et al., 2001; Creamen et al., 2013). In addition, mineral dust particles can act as cloud condensation and ice nuclei (Twohy et al., 2009; Ansmann et al., 2009a; DeMott and Prenni, 2010) and affect air quality (Fairlie et al., 2010). The high temporal and spatial variability of dust particles and the complexity in their microphysical and optical properties present a significant challenge to our understanding of how these particles impact the environment.

Numerous field campaigns have been conducted to better characterize mineral dust properties, e.g. the Saharan Mineral Dust Experiments SAMUM-1 and SAMUM-2 (Ansmann et al., 2009b, 2011a and references therein) and the Saharan Aerosol Long-Range TRansport and Aerosol Cloud interaction experiment SALTRACE (http: //www.pa.op.dlr.de/saltrace/), among others. However, the information on mineral dust properties is still guite scarce (Formenti et al., 2011), even though many measurements worldwide have been made using different approaches. Satellites are providing global coverage but the retrievals of particle properties are affected by large uncertainties (Levy et al., 2013). Moreover, the interaction of dust particles with solar and terrestrial radiation is complex due to their irregular shapes and variable refractive indices (Mishchenko et al., 1997). Because of this, in the past years it has been difficult to develop accurate algorithms for the retrieval of dust microphysical properties from optical measurements. Dubovik et al. (2006), one of the first studies that addressed this problem, developed an algorithm that took into account the scattering patterns of non-spherical particles and implemented an inversion method for columnintegrated radiometric measurements in the AERONET (Aerosol Robotic Network) netAMTD

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work (http://aeronet.gsfc.nasa.gov/, Holben et al., 1998). Other approximations using non-spherical particles have also been proposed, e.g. Olmo et al. (2006), Valenzuela et al. (2012a).

Information on the vertical distribution of mineral dust properties is also essential for 5 understanding particle transport processes from regional to intercontinental scales, to improve radiative forcing calculations and to analyze the influence of mineral dust on cloud formation (Ansmann et al., 2008, 2009, 2011a; Seifert et al., 2010). Therefore, it is crucial to develop advanced methods to characterize dust microphysical properties from remote sensors like multi-wavelength Raman lidar or High Spectral Resolution Lidar (HSRL) systems (Müller et al., 2010; Veselovskii et al., 2010). In this framework Veselovskii et al. (2010) implemented the kernel functions of Dubovik et al. (2002) in a regularization technique (Müller et al., 1999; Veselovskii et al., 2002) to obtain vertically-resolved dust properties. This approach was first implemented using data of the SAMUM field campaign (Müller et al., 2013) and from measurements of long-range transport of dust over Europe (e.g. Veselovskii et al., 2010; Papayannis et al., 2012).

Raman or HSRL systems have limitations such as the low signal-to-noise ratio, which limits their use mostly during night-time, and the high cost and maintenance. Hence, their implementation is not very widespread. On the contrary, elastic backscatter lidar systems offer an easier set-up and maintenance and larger signal-to-noise ratio during daytime. Their operation is much more extended worldwide due to implementation in networks such as EARLINET (European Aerosol Research Lidar Network) (Pappalardo et al., 2014) and MPLNET (Micro Pulse Lidar Network). In this context, the LIRIC (Lidar Radiometer Inversion Code) code was developed by Chaikovsky et al. (2012) to obtain vertically-resolved profiles of aerosol microphysical properties by combining elastic lidar measurements and column-integrated sun-photometer microphysical properties retrieved from AERONET.

The first results obtained with LIRIC have already been presented by Wagner et al. (2013) and Granados-Muñoz et al. (2014); however, a complete evaluation, using ancillary information such as in-situ vertically-resolved measurements, is still lack-

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ing. Generally speaking, there have been very few in-situ measurements to directly compare with those made with remote sensors. Murayama et al. (2003) did a direct comparison between lidar derived and in-situ extinction coefficients during the Aerosol Characterization Experiment – Asia (ACE-Asia) and observed that the in-situ measurements produced larger values that the lidar. Comparisons between volume concentrations measured with in-situ instruments and derived from remote sensors are especially rare. Other than the study of Bravo-Aranda et al. (2014), who compared coarse mode mass derived from airborne spectrometer measurements with mass concentrations derived from lidar and observed that the in-situ measurements were larger than the lidar, there are very few direct comparisons in dust layers between the same parameter measured in-situ and remotely. Most comparisons, such as those reported by Weinzierl et al. (2009, 2011) only do a qualitative comparison whereby the relative magnitudes are compared as a function of the altitude rather than directly comparing the absolute quantities.

The objectives of the present study are to analyze the evolution of the microphysical properties of particles in a mineral dust plume using both LIRIC and a regularization technique applied to multiwavelength-Raman lidar measurements (Veselovskii et al., 2010) and to compare LIRIC profiles of particle volume concentration and depolarization ratios with airborne in-situ measurements.

## 2 Experimental site and instrumentation

#### 2.1 Experimental site

The data from ground based-instrumentation were acquired at the Andalusian Institute for Earth System Research (IISTA-CEAMA) located in the city of Granada (37.16° N, 3.61° W, 680 m a.s.l.; Lyamani et al., 2010; Titos et al., 2012; Valenzuela et al., 2012b). Granada is a medium-size city in the South-East of Spain located in a natural basin, delimited on the East by mountains with peaks up to 3000 m a.s.l. Air masses affecting

the area arrive mainly from the Atlantic Ocean, Central Europe, the Mediterranean Basin and North-Africa (Valenzuela et al., 2012a; Perez-Ramirez et al., 2012a). The number of mineral dust events at the Granada station originating in North Africa is quite high, especially during summer, with an occurrence of 45 % of the days in June, July 5 and August (Valenzuela et al., 2012b). These events can transport particles at altitudes of up to 5500 ma.s.l., not always affecting the surface level (Guerrero-Rascado et al., 2008, 2009; Navas-Guzmán et al., 2013). The experimental site is also impacted by anthropogenic particles from local and regional aerosol sources (Lyamani et al., 2008, 2010, 2012; Titos et al., 2014).

#### **Ground-based instrumentation**

The Raman-lidar MULHACEN (based on LR331D400, Raymetrics, Greece) used for aerosol vertical-profiling is described in detail by Guerrero-Rascado et al. (2008, 2009) and Navas-Guzmán et al. (2011). It employees a Nd:YAG laser that emits at three different wavelengths (355, 532 and 1064 nm). The receiving system consists of detectors that split the radiation according to the three elastic channels (355, 532 and 1064 nm), two nitrogen Raman channels (387, 607 nm) and a water vapour Raman channel (408 nm). These Raman measurements have sufficient signal-to-noise ratio only for night-time detection. The system also measures depolarization of the returned signal at 532 nm (532-cross and 532-parallel detection channels) (Bravo-Aranda et al., 2013) for retrieving vertical profiles of the particle linear depolarization ratio  $(\delta_1^P)$ . The estimated uncertainties associated with the lidar signals are between ±15 and 20 % for the aerosol particle backscatter coefficient,  $\beta_{\lambda}^{\text{aer}}$ , and  $\pm$  20% for the aerosol particle extinction coefficient,  $\alpha_{\lambda}^{\text{aer}}$ . These estimates are based on the statistical uncertainties retrieved with Monte Carlo techniques according to Pappalardo et al. (2004) and Guerrero-Rascado et al. (2008). The procedure described by Wandinger and Ansmann et al. (2002) to correct the incomplete overlap of the system is applied to our data. The use of this overlap correction allows to obtain reliable  $\beta_{\lambda}^{aer}$  profiles **AMTD** 

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at 355 and 532 nm down to 320 m above the station (Navas-Guzmán et al., 2011);

however, reliable data are obtained only from  $\sim$  1000 m above the station for  $\alpha_{\lambda}^{\rm aer}$ . The Raman lidar system is part of EARLINET and currently is included in the ACTRIS (Aerosols, Clouds, and Trace gases Research InfraStructure Network) European project (http://www.actris.net/).

Sun photometric measurements obtained at Granada are used to obtain columnintegrated aerosol properties using a CIMEL CE-318. This instrument is included in the AERONET-RIMA network (Iberian Network for Aerosol Measurements, federated to AERONET) (http://www.rima.uva.es/index.php/en/) since 2002 and is calibrated following the AERONET protocols. Details about the CIMEL sun-photometer can be found in Holben et al. (1998), however a brief description is presented here. This instrument makes direct sun measurements at 340, 380, 440, 500, 670, 870, 940 and 1020 nm and sky radiance measurements at 440, 670, 870 and 1020 nm. The direct sun measurements are used to retrieve aerosol optical depth  $(\tau_1)$  at 340, 380, 440. 500, 675, 870 and 1020 nm. The  $\tau_{\lambda}$  uncertainties provided by AERONET are  $\pm 0.02$  for  $\lambda < 400 \,\mathrm{nm}$  and  $\pm 0.01$  for  $\lambda > 400 \,\mathrm{nm}$ . Additionally, the spectral dependency of the  $\tau_1$ has been considered through the Ångström exponent,  $\alpha$  (440–870), calculated in the range 440-870 nm. Also included in the analysis are aerosol optical depths at 500 nm for fine mode ( $\tau_{\text{fine}}$ ) and for coarse mode ( $\tau_{\text{coarse}}$ ) as well as the fine mode fraction  $(\eta)$  (ratio of  $\tau_{\text{fine}}$  to  $\tau$ ), determined using the spectral de-convolution algorithm method developed by O'Neill et al. (2003). In addition, column-integrated aerosol microphysical properties (size distribution, refractive index, volume concentration, etc.) provided by the AERONET code are also used (Dubovik and King, 2000; Dubovik et al., 2002, 2006). For the retrieval of the aerosol microphysical properties both the direct sun and the sky radiance measurements are used. The reported size distribution retrieval uncertainties are  $\pm 10$ –35 %, for the size range 0.1  $\mu$ m < r < 7  $\mu$ m, and outside this range they are as large as ±80-100 %. All the data used here are Level 1.5 data obtained using the AERONET Version 2 algorithm. Only a small number AERONET Level 2.0 were available due to the restrictions imposed by AERONET code ( $\tau_{440\,\mathrm{nm}} > 0.4$  and solar zenith angle > 50°). Therefore, AERONET Level 1.5 (cloud screened data with

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pre- and post-calibrations applied) were used in this study, considering only those data that fulfil the following conditions to assure their quality:  $\tau_{440\,\mathrm{nm}} > 0.2$  and solar zenith angle  $> 50^\circ$ .

The star-photometer EXCALIBUR provides  $\tau_{\lambda}$  measurements from the stars' direct irradiance using interferential filters at 380, 436, 500, 670, 880 and 1020 nm (nominal wavelengths) using a CCD camera as detector during night-time. More details can be found in Perez-Ramirez et al. (2008a, b). A regular calibration of the instrument is performed once a year at a high mountain location. Errors in the  $\tau_{\lambda}$  are  $\pm 0.02$  for  $\lambda < 800$  nm and  $\pm 0.01$  for  $\lambda > 800$  nm (Perez-Ramirez et al., 2011). Data are cloud-screened and quality assured over 30 min intervals (Perez-Ramirez et al., 2012b).

#### 2.3 Aircraft in-situ instrumentation

Meteorological and aerosol particle measurements were made with instruments mounted on a CASA C-212-200 research aircraft operated by the Spanish National Institute of Aerospace Technology (INTA). Details on aircraft measurements can be found in Andrey et al. (2014) and Bravo-Aranda et al. (2015). Here we give a short overview of the instruments used for our study: the Cloud and Aerosol Spectrometer with Polarization detection (CAS-POL) and the the Passive Cavity Aerosol Spectrometer (PCASP-100X).

The CAS-POL measures the light scattered by individual particles passing through a focused 658 nm polarized laser beam (Baumgardner et al., 2001). The instrument makes use of 30 channels logarithmically distributed to cover the 0.51–50  $\mu m$  size range and depolarization ratios are derived from the light back-scattering patterns from individual particles. In this study, CAS-POL measurements are used to obtain both the aerosol volume concentration and the depolarization ratio. A detailed description of the polarization features of the CAS-POL and associated uncertainties are included in the Appendix.

Additionally, the Passive Cavity Aerosol Spectrometer (PCASP-100X) that provides aerosol size distributions in the  $0.1-3\,\mu m$  diameter range in 15 different bins has been

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used in order to enhance the size range provided by the CAS-POL measurements. The measuring principle and uncertainties (20% for the size distributions) have been described in detail by Cai et al. (2013), Rosenberg et al. (2012) and references therein.

#### 3 Methodology: retrievals of aerosol particles microphysical properties

#### 3.1 LIRIC code

The LIRIC algorithm is applied to lidar and sun-photometer data to retrieve aerosol particle microphysical properties profiles during daytime (Chaikovsky et al., 2008, 2012; Wagner et al., 2013; Granados-Muñoz et al., 2014). This algorithm uses as input data the column-integrated optical and microphysical properties retrieved from AERONET code (Dubovik et al., 2002, 2006) and measured lidar elastic backscattered signals at three different wavelengths (355, 532, and 1064 nm). The depolarization information from lidar data can optionally be used. To perform the retrieval, an aerosol model, based on the AERONET code, which assumes a mixture of randomly oriented spheroid and spherical particles defined by the column-integrated volume concentrations of each mode, is used (Dubovik and King, 2000; Dubovik et al., 2006). Subsequently, an iterative procedure based on the Levenberg-Marquardt method is applied. The combined lidar and sun-photometer information provides volume concentration profiles for the fine and coarse modes, distinguishing also between coarse spherical and coarse spheroid particles if depolarization information is considered. The separation between the fine and the coarse modes is made by searching the radius located at the minimum of the AERONET bimodal size distribution in the radii range between 0.194 and 0.576 µm. Uncertainties of the volume concentration profiles related to the different user-defined parameters used within LIRIC algorithm are usually below 15% (Granados-Muñoz et al., 2014), proving LIRIC to be guite stable. A detailed analysis on the uncertainty due to the input lidar and sun-photometer data uncertainties is still lacking, but it can be estimated for our case. The uncertainty in the retrieved volume

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concentration is related to the size distribution that is retrieved from AERONET. As discussed above, this uncertainty can range from  $10-35\,\%$  if the particle radii are less than 7 µm but can be as much as  $100\,\%$  if there is significant volume in particles larger than this. As will be illustrated below, measurements of the volume size distribution with the CAS-POL show that there are particles larger than 7 µm radius and in the dust layer they can be larger than  $30\,\mu\text{m}$ . Hence, a reasonable estimate of the retrieved volume uncertainty is approximately  $50\,\%$ , similar to what is estimated for the volume measured by the CAS-POL (see Appendix).

# 3.2 Inversion of Raman lidar measurements to retrieve microphysical properties

The Raman lidar system provides  $\alpha_{\lambda}^{\text{aer}}$  profiles at 355 and 532 nm,  $\beta_{\lambda}^{\text{aer}}$  at 355, 532, 1064 nm and the  $\delta_{\lambda}^{\text{P}}$  at 532 nm. This  $3\beta + 2\alpha + 1\delta$  data set was inverted to retrieve the aerosol particle microphysical properties using the regularization approach, described in detail by Müller et al. (1999) and Veselovskii et al. (2002, 2004).

To account for mineral dust particles non-sphericity the model of randomly oriented spheroids was used, as described in Veselovskii et al. (2010). Following this approach, the particle volume concentration and effective radius were estimated with an uncertainty of about 50 and 25 %, respectively. The real part of the refractive index,  $m_{\rm r}$ , was also estimated, with an uncertainty of  $\pm 0.05$ .

# 3.3 Inversion of $\tau_{\lambda}$ spectral measurements from sun- and star-photometry measurements

In the present study, we used column-integrated aerosol properties provided by AERONET. These variables were retrieved by inversion of  $\tau_{\lambda}$  and sky radiances using the AERONET operational algorithm. In addition to these AERONET aerosol retrievals, we inverted  $\tau_{\lambda}$  spectral values obtained from both the sun- and the star-photometer with the Linear Estimation (LE) algorithm to get aerosol particle properties (e.g. integrated

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volume and effective radius) with high temporal resolution (Kazadzis et al., 2014). The LE algorithm is described in Veselovskii et al. (2012) and more details are provided by Perez-Ramirez et al. (2015), where correction functions to get accurate aerosol data close to that of the operational AERONET code were introduced.

#### Results and discussion

#### Overview of the measurement campaign

Different models such as NAAPS (Navy Aerosol Analysis and Prediction System) (Christensen, 1997) and BSC-DREAM8b (Pérez et al., 2006a, b; Basart et al., 2012), forecast mineral dust over the Granada station on the 27 June 2011. Based on these forecasts, a measurement campaign was launched using all the available instrumentation at the IISTA-CEAMA experimental station in coincidence with a simultaneous flight of the CASA C-212-200 research aircraft. The flight took place around 10:30 UTC and the aircraft performed a pseudo spiral track flying from 1200 to 5200 ma.s.l. at a distance of approximately 8 km from the IISTA-CEAMA station. A detailed description of the field campaign and an analysis of the aerosol optical properties retrieved from the ground-based instrumentation are presented by Bravo-Aranda et al. (2015), with a main focus on the analysis of the entrainment and mixing processes of the mineral dust layer within the planetary boundary layer. In our study, however, the main focus is on the evolution of the aerosol optical and microphysical properties evaluated with different instrumentation and different approaches, including a comparison of LIRIC volume concentration and lidar depolarization ratio profiles with in-situ aircraft measurements.

Figure 1 shows the range corrected signal (RCS) time series measured by the lidar on the 27 June 2011. Measurements were performed between 00:00 and 01:00 UTC and 06:45 and 12:10 UTC. According to the lidar measurements, a strong backscattered signal associated with long-range transport of mineral dust (as we will show later on) was observed up to 5 km a.s.l. during both sampling periods.

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Figure 2 shows columnar aerosol optical and microphysical properties from the sunand star-photometry measurements. Figure 2a illustrates the temporal trends in the  $\tau_{500\,\mathrm{nm}}$ , and the corresponding contributions of fine  $(\tau_{\mathrm{fine}})$  and coarse  $(\tau_{\mathrm{coarse}})$  modes using the Spectral Deconvolution Algorithm (O'Neill et al., 2001a, b; 2003). The  $\tau_1$ - $_{5}$  related Ångström exponent (AE( $\lambda_{1}-\lambda_{2}$ )) computed using the wavelengths between 440 and 870 nm (436 and 880 nm for star-photometry) and the fine mode fraction ( $\eta$ ) are shown in Fig. 2b. As can be observed, there was a smooth temporal-evolution of the aerosol properties with small variations.  $\tau_{500\,\mathrm{nm}}$  time series indicated that the aerosol load slightly varied during the analysed period, with values ranging between 0.27 and 0.37.  $\tau_{\text{coarse}}$  was significantly larger than  $\tau_{\text{fine}}$  during the end of the night and day-time measurements whereas  $\tau_{\text{fine}}$  was almost constant around 0.1 during the study period. The simultaneous increase of  $\tau_{500\,\mathrm{nm}}$  and  $\tau_{\mathrm{coarse}}$  from midnight suggests an increase of the incoming mineral dust in the atmospheric column. However, during the first hours of the night, the fine mode had a relevant contribution to the total  $\tau_{500\,\mathrm{nm}}$ . The AE ranged between 0.80 at night-time and 0.4 during day-time, suggesting a clear predominance of coarse particles that can be associated with the transport of dust particles. However, the larger values obtained at the beginning of the night period are higher than those acquired from AERONET measurements in the presence of mineral dust (e.g. Dubovik et al., 2002), and thus the contribution of fine particles to the aerosol mixture is also considerable.

The evolution of the effective radius,  $r_{\text{eff}}$ , and the column volume concentration, V, obtained both by LE and AERONET retrievals during day-time and by LE during night-time for the 27 June 2011 is shown in Fig. 2c and d. Good agreement between AERONET and LE retrievals was obtained during daytime corroborating the good performance of the LE method. In addition, continuity between the star- and sunphotometer measurements was observed, which shows the utility of combining both instruments to perform a continuous monitoring of aerosol properties. As observed in Fig. 2c and d, both  $r_{\rm eff}$  and V were slightly lower during the night compared to daytime. The  $r_{\rm eff}$  values were around 0.3–0.4 µm during night-time and almost constant around

0.5 µm during daytime. These values suggest a predominance of coarse particles in the atmospheric column. The V values were increasing from  $0.15\,\mu\text{m}^3\,\mu\text{m}^{-2}$  during the night up to  $0.25\,\mu\text{m}^3\,\mu\text{m}^{-2}$  in the early morning and then they decreased again down to 0.1 µm<sup>3</sup> µm<sup>-2</sup> at noon. The changes observed in aerosol properties suggest a mixture 5 of different aerosol types changing from night to day.

During the daytime, a comprehensive analysis of the vertically resolved optical properties retrieved by means of Klett-Fernald (Fernald et al., 1972; Fernald, 1984; Klett, 1981) method was presented by Bravo-Aranda et al. (2015). However, a brief discussion is included here to properly discuss the microphysical properties profiles retrieved with LIRIC during daytime.

Figure 3 shows the analysis of backscattered elastic lidar signals for measurements at daytime on the 27 June retrieved by means of the Klett-Fernald inversion method (Fernald et al., 1972; Fernald, 1984; Klett, 1981) using lidar ratio values between 40 and  $48 \, \mathrm{sr}^{-1}$ . Namely, aerosol particle backscatter profiles at 532 nm,  $\beta_{532 \, \mathrm{nm}}^{\mathrm{aer}}$ , backscatter related Ångström exponent between 355 and 532 nm,  $\beta$ -AE(355–532 nm), and linear particle depolarization profiles,  $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$ , obtained during day-time from lidar elastic measurements are shown. According to the analysis of these optical properties profiles, two different layers can be distinguished. The first layer is in the height range between 3000 and 5000 ma.s.l. In this layer, the  $\beta_{532\,\mathrm{nm}}^{\mathrm{aer}}$  values are decreasing during the morning. For this height range,  $\beta$ -AE(355–532 nm) profiles suggest the predominance of coarse particles with values close to zero during the analysed period, in coincidence with  $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$  values ( $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$  in the range 0.23–0.28) that suggest an important contribution of non-spherical particles (Gross et al., 2011). However, below 2000 ma.s.l.,  $\beta$ -AE(355–532 nm) was decreasing from 2 in the early morning down to  $\sim 0.5$  around midday. At the same time,  $\delta_{\rm 532\,nm}^{\rm P}$  was increasing from 0.08 up to 0.20. During the morning, these results suggest the influence of anthropogenic particles from local origin that is mixed with the mineral dust due to convective processes within the planetary mixing layer (Bravo-Aranda et al., 2015).

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LIRIC retrievals obtained for the morning of the 27 June 2011 are shown in Fig. 4. A clear predominance of the coarse spheroid mode is observed from the surface up to 5000 ma.s.l., as expected for mineral dust events. In addition, a decrease of the total volume concentration values was observed throughout the morning, in agreement with the decrease in  $\beta_{532}^{\text{aer}}$  (Fig. 3) and the decrease in the integrated volume concentration V (Fig. 2d). A maximum peak in the volume concentration of the coarse spheroid mode was observed between 4000 and 4500 m a.s.l., in coincidence with the maximum in  $\beta_{532\,\mathrm{nm}}^{\mathrm{aer}}$  profiles, indicating the presence of an aerosol layer at this height. It is worth noting that fine particles were also observed during the different analysis periods as indicated by the volume concentration profiles, but in low concentrations ( $\sim 6 \, \mu m^3 \, cm^{-3}$ ). The fact that the profiles of the fine mode volume concentration show a maximum peak at the same height as the coarse spheroid mode suggests that these fine particles could have been advected with the mineral dust. Therefore, they might correspond to the fine mode of mineral dust. There is also the possibility that they correspond to anthropogenic aerosol originating from the industrial areas in the North of Africa together with the mineral dust (Rodríguez et al., 2011; Valenzuela et al., 2012c), as indicated by the backward trajectories analysis performed with HYSPLIT model using GDAS meteorological database (http://ready.arl.noaa.gov/HYSPLIT\_traj.php, figure not shown). Another possible explanation is that it could be an artifact introduced by LIRIC due to problems with the incomplete overlap or in cases of non-homogeneous layering (Granados-Muñoz et al., 2014), distributing the contribution of local pollution at high altitudes.

The coarse spherical mode volume concentration slightly increased during the morning reaching its maximum values (~ 7  $\mu m^3$  cm<sup>-3</sup>) between 11:15 and 11:45 UTC. This slight increase in the volume concentration of the coarse spherical mode (from 0 up to  $5 \, \mu m^3 \, cm^{-3}$ ) around 4500 m a.s.l. was in agreement with a slight decrease in  $\delta_{532 \, nm}^{P}$ values (from 0.28 to 0.23) in the same height range, indicating a larger contribution of spherical particles and corroborating once again the coherence between the aerosol optical and microphysical properties profiles obtained.

The layering observed in the optical properties profiles (Fig. 3) indicated the presence of mineral dust in the upper layer between 3000 and 5000 m a.s.l. and the presence of anthropogenic particles in the lower part of the troposphere that was not detected by LIRIC. This can be explained by the fact that LIRIC assumes several AERONET retrieved properties (i. e., refractive index, size distribution, sphericity, etc.) as height-independent and therefore LIRIC results are much more vertically homogeneous regarding the distribution of the modes than those retrieved from the lidar data with Klett-Fernald algorithm. A combined inversion of lidar and sun-photometer data without such assumptions, as the one proposed in GARRLIC (Generalized Aerosol Retrieval from Radiometer and Lidar Combined data) (Lopatin et al., 2013), might provide more accurate results regarding the vertical distribution of the aerosol properties, although GARRLIC is out of the scope of this work. Meanwhile, the interpretation and analysis of the LIRIC-retrieved volume concentration profiles needs to be carefully made when the atmospheric aerosol layers comprise different aerosol types, as shown in Granados-Muñoz et al. (2014).

## 4.2 Night-time retrieval of mineral dust microphysical properties profiles

The Raman  $3\beta + 2\alpha + \delta$  measurements on the night previous to the flight were inverted to retrieve vertical profiles of volume concentration,  $r_{\rm eff}$  and the real part of the refractive index using the regularization approach (Veselovskii et al., 2010). The comparison of these profiles with LIRIC results using the first data of the day, acquired at 06:15–06:45 UTC, allow us to see the evolution of the vertical structure of the aerosol microphysical properties. A comparison of both techniques over the same time period was not possible due to the lack of simultaneous data.

Figure 5a and b show the optical properties retrieved with the Raman method during night-time (averaged between 00:00 and 01:00 UTC). The optical properties shown in Fig. 5a and b indicate values of  $\beta_{532\,\mathrm{nm}}^{\mathrm{aer}}$  similar to those observed in Fig. 3, corresponding to the period between 06:15 and 06:45 UTC. However, the vertical structure of the aerosol layer was different, corroborating the evolution of the aerosol vertical structure

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ture during the night. As can be observed, two peaks were detected during night-time around 3200 and 4200 ma.s.l. whereas a much more homogeneous structure was observed during daytime, with only one maximum at 4100 ma.s.l. The  $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$  profiles and its inverse correlation with  $\beta$ -AE(355-532 nm) in Fig. 5b corroborates also the pres-5 ence of two differentiated regions with different aerosol types during night-time, namely there were mineral dust particles above 2250 ma.s.l. ( $\delta_{532\,\mathrm{nm}}^{\mathrm{P}} \sim 0.25$ ) and a mixing of local anthropogenic aerosol with mineral dust below this height. Values of the different aerosol properties in both regions are shown in Table 1.

The overlap effects prevented using extinction data below 2000 m and therefore inversion of microphysical properties was not possible below this height. For the inversion, the optical data were averaged in 250 m layers for the heights between 2000 and 3600 ma.s.l. Above that height averaging was done in 1000 m layers. Figure 5c-e shows the corresponding microphysical properties obtained by the regularization technique. As we can observe in Fig. 5c, reff values varied in the different layers. The highest values, ~ 1.76 μm, related to larger particles (mineral dust), were found in the layer between 2500 and 3500 m a.s.l., while the smallest values, ~ 0.53 µm, were found in the lower layer. Values around 4200 m a.s.l. ( $\sim 1.1 \, \mu m$ ) are still larger than those in the lower layer, but lower than in the maximum around 3200 ma.s.l. This decrease in the radius with height can be due to an artifact of the algorithm because of the larger averaging height interval used in the upper part of the profiles and not necessarily related to a decrease in the size of the particles. Volume concentration values were also larger for the dust layer than for the one with mixed particles. This fact can be explained by both the volume concentration dependence on particle radius and also the larger aerosol load in this layer (as observed in the layering structure of  $\beta_{532\,\mathrm{nm}}^{\mathrm{aer}}$  in Fig. 5a). Two regions were also clearly distinguished in the  $m_r$  profile obtained with the regularization technique. The  $m_r$  values were larger in the upper part of the profile, corresponding to the mineral dust layer and lower values were obtained in the lower part of the profile, due to the presence of the mixture of mineral dust and anthropogenic particles. The mean value of  $m_r$  in the profile was 1.55  $\pm$  0.05. Figure 5e shows the volume size dis-

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tributions for different altitudes. The volume size distribution at the surface level was obtained from the Aerodynamic Particle Sizer (APS-3321, TSI) (Volcken and Peters, 2003; Lyamani et al., 2010). The APS is an aerosol optical counter that measures aerosol particle size distribution and concentration in the aerodynamic radius range between 0.25 and 10 µm in 52 nominal size bins. Therefore most of the fine mode is not detected. The volume size distributions at higher altitudes were obtained by the regularization technique. In addition, the closer in time column-integrated AERONET size distribution (26 June 2011 at 18:15 UTC) was included. From both APS and lidar measurements, it was observed a clear increase in the coarse mode radius with height, as the location of the maximum is displaced towards larger radii, in agreement with  $r_{\rm eff}$ profiles. There was also an increase of the aerosol load since the maximum in the volume concentration strongly increased with height. Fine mode was almost insignificant in the different layers. However, the AERONET column-integrated distribution showed a small contribution of the fine mode aerosol particles and also the coarse mode radius shifted to smaller values. This can be explained by the presence of mineral dust particles at higher altitudes while at lower levels dust was mixed with local anthropogenic pollution. In the first 400 m above ground it seemed that fine aerosol particles predominated, but because of the lidar incomplete overlap and the limitations of the APS this could not be confirmed. However, in-situ measurements presented by Bravo-Aranda et al. (2015) pointed in this direction.

### Comparison of airborne in-situ measurements and ground-based retrieved profiles

Figure 6a shows the vertical profiles of the volume concentration derived from the AERONET and lidar measurements using the LIRIC algorithm and from the aircraft in-situ measurements. Since AERONET size distribution includes only particles up to 10 µm radius, in order to make a coherent comparison only particles below 20 µm diameter measured by the CAS-POL were considered in the calculations of the volume concentration profile (red profile in Fig. 6a). A second profile including the complete size

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distribution from the CAS-POL (black profile in Fig. 6a) is also included for completeness. For the retrieval of the volume concentration profiles from the in-situ aircraft measurements, a refractive index of 1.54 (real part) (McConnell et al., 2010) was assumed and Mie theory was applied assuming spherical aerosol particles as explained by Andrey et al. (2014). This refractive index value is very similar to the one obtained with the regularization technique for the night-time measurements, which was  $1.55 \pm 0.05$ . The agreement in the vertical structure between the aircraft and LIRIC volume concentration profiles was quite good with respect to the location of the peaks. Similar layering was detected with both LIRIC and the airborne data, distinguishing two maximum peaks around 3500 and 4200 km a.s.l. The height of the first maximum obtained by aircraft in-situ measurements was located at a slightly higher altitude (3450 m a.s.l.) than by the LIRIC algorithm (3250 ma.s.l.). The geometrical thickness of the different layers observed were also very similar for both LIRIC and the aircraft data.

Regarding the volume concentration values, the differences are within the expected uncertainties, marked by the horizontal bars in Fig. 6. In general, average in-situ values exceed those from LIRIC by less than 20 µm<sup>3</sup> cm<sup>-3</sup>, except for the concentration maximum between 3200 and 3500 m if we consider only particles below 10 µm. Differences are much larger if we compare LIRIC retrieval to the volume concentration profiles retrieved using the complete size distribution provided by the CAS-POL. The difference between the two profiles retrieved from the aircraft in-situ measurements reveals a likely underestimation of the volume concentration profiles retrieved by LIRIC by some fraction that depends on the amount of particle volume in sizes larger than 10 µm (in radius) not considered in AERONET. A similar behaviour was found by Bravo-Aranda et al. (2014): when comparing the dust mass concentration retrieved by applying the POLIPHON (Polarizing Lidar Photometer Networking) method to lidar data (Ansmann et al., 2011b) with the coarse-mode mass concentration derived from the aircraft measurements (assuming a density of 2.6 g cm<sup>-3</sup> for the mineral dust) larger differences where observed in the maximum peaks and a similar displacement of the aerosol layers was found. The difference between the in-situ and remote sensing concentrations

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raises the question as to which technique is providing the more accurate value. Both techniques rely on certain assumptions that could potentially bias the results. In the discussion of the CAS-POL uncertainties in the Appendix, the two most important assumptions are related to the refractive index and shape of the particles. If the refractive index of the particles differ from what is assumed, this can bias the sizing either smaller or larger, but no more than 50%. The assumption that the particles are spherical will bias the sizes, and hence the volumes, perhaps even as much as a factor of two if the dust particles are very aspherical. This could explain the discrepancies between the two techniques over most of the profiles where the differences are within a factor of two. Countering that argument, however, is the comparison of the size distributions measured with the combined PCASP-100X and the CAS-POL and those derived from AERONET. Figure 7 shows average size distributions in the five altitude ranges labelled (1) through (5) in Fig. 6a. When compared with the size distributions derived from AERONET (Fig. 5d), we observe that both measurements show a very similar distribution (note that AERONET reports in radius whereas the CAS-POL measurements are in EOD), e.g. the peak volume falls at 10 µm EOD (5 µm radius). This suggests that the larger volume reported from the CAS-POL is not a result of oversizing.

In situ measurements of depolarization ratios from aircraft have never been published before so the results presented here are the first opportunity to compare profiles from aircraft with those derived from remote sensors. The CAS-POL mean depolarization ratio at a wavelength of 680 nm is compared with the  $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$  profile retrieved at 10:30 UTC from the lidar data (Fig. 6b). The uncertainty of the depolarization ratio from the CAS-POL measurements is estimated to be around 30 % (see Appendix), similar to what is estimated for the lidar. The horizontal bars on each curve illustrate these uncertainties. In general, we observe reasonable agreement in the trends derived from lidar  $\delta_{532 \, \text{nm}}^{\text{P}}$  profiles and CAS-POL, especially in the layer between 4000 and 4500 ma.s.l., where discrepancies are close to 0. In the rest of the profile, the CAS-POL values are somewhat larger than the lidar-retrieved  $\delta_{\rm 532\,nm}^{\rm P}$ . Nonetheless, the discrepancies are below 15% for most part of the profile, which is within the uncertainty limits, corroborating the good performance of both techniques. It is worth noting that not only does the CAS-POL measure the scattered light at a longer wavelength than the lidar, but in addition, whereas the lidar measures at 180° backscatter, the CAS-POL is measuring over backscattering angle from 168 to 176°. The importance of this difference can be seen in Fig. 8 that shows theoretical calculations of polarization ratio as a function of scattering angle and different shapes (courtesy of P. Yang, Texas A&M University). The shapes are those of different types of ice crystals but serve equally well to demonstrate the degree of depolarization as a function of the measurement angle. The polarization ratio defined here are from Stokes matrix components  $P_{11}$  and  $P_{22}$ . For linearly polarized incident light, the ratio of  $P_{22}/P_{11}$  is the inverse of what is reported from the lidar and CAS-POL. What is important to note is that for all of the shapes except rough aggregates, the ratio is smaller at 180° than for the 168 to 176° range. Rough aggregates are quasi-spherical whereas dust is likely more aspherical leading to the observation that the higher polarization ratios measured by the CAS-POL may be mostly due to the difference in collection angles.

#### 5 Summary and conclusions

An exhaustive measurement campaign was performed on the 27 June 2011 at Granada station during a Saharan dust outbreak. Active and passive remote sensing techniques were used to continuously monitor the atmosphere together with in-situ measurements on board an aircraft. This study focused particularly on the analysis of the microphysical properties, column integrated and vertically resolved, implementing different processing algorithms, i.e. LIRIC, the regularization method and the Linear Estimation technique. To our knowledge this is the first time that these techniques are processed and compared together along with a direct comparison between in-situ aircraft and remote sensing measurements.

Column-integrated properties were retrieved by means of star and sun-photometer measurements. Night-to-day evolution of the aerosol optical depth at 500 nm and the

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Ångström exponent followed a smooth behaviour explained by the natural variability of the aerosol particle population. The aerosol optical depth at 500 nm ranged between 0.27 and 0.37 during the analysis period, while the Angström exponent varied from 0.80 at night-time to 0.4 during the day-time. Column-integrated microphysical properties retrieved with the operational AERONET and Linear Estimation codes were in good agreement. Moreover, the Linear Estimation allowed analysis of the temporalevolution from night- to day-time of effective radius and volume concentration. A smooth temporal-evolution has been observed, with the effective radius ranging between 0.3 and 0.5 µm. The derived Ångström exponent and effective radius suggest a predominance of coarse particles but with some contribution by the fine mode particles, which was significant during some periods.

The analysis of the aerosol optical properties profiles highlighted the presence of two aerosol layers during daytime: a lower layer corresponding to a mixture of anthropogenic aerosol with mineral dust and a higher layer corresponding to pure mineral dust. As expected in presence of mineral dust, the LIRIC analysis indicates a clear predominance of the coarse spheroid mode, with average volume concentrations around 30 µm<sup>3</sup> cm<sup>-3</sup>, reaching 60 µm<sup>3</sup> cm<sup>-3</sup> at the maximum peaks. Fine particles were observed in lower concentrations (~ 6 µm<sup>3</sup> cm<sup>-3</sup>) at high altitudes, suggesting that they could be anthropogenic particles originated in the industrial areas in the North of Africa or they might correspond to the mineral dust fine mode. However, they could also be related to a LIRIC artifact that distributes the local anthropogenic aerosol at higher altitudes. Results obtained with LIRIC are in agreement with the results obtained from the analysis of the optical properties retrieved from the lidar data in the upper layer corresponding to mineral dust. However, there is some disagreement in the lowermost part of the profiles, below 2000 ma.s.l., which corresponds to the mixture of mineral dust with anthropogenic aerosol. These discrepancies are mainly related to the assumption of height-independent aerosol properties in LIRIC and indicate that LIRIC profiles need to be carefully interpreted in cases of non-homogeneous aerosol layers.

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The regularization technique applied to the Raman lidar data during the night previous to the flight was used to evaluate the temporal evolution of the aerosol microphysical properties profiles. According to the lidar data, the aerosol backscatter coefficient profiles,  $\beta_{532nm}^{aer}$ , showed larger values during the night than in the early morning in certain layers indicating changes in the aerosol vertical structure due to the temporal evolution of the dust event during the night. The retrievals of total volume concentration profiles at night (with the regularization technique) and in the early morning (with LIRIC) also indicated a decrease in the volume concentration from maximum values of 75-50 μm<sup>3</sup> cm<sup>-3</sup> and changes in the aerosol vertical structure. However, column integrated values showed small changes in the aerosol properties, remaining almost constant during the night. These results underscore the need for vertically resolved measurements to adequately monitor the evolution of the aerosol properties. The discrepancies between regularization and LIRIC results are mainly explained by the natural variability of the aerosol during the night. Because of this temporal variability, results obtained from the two different approaches are not comparable in absolute terms. However, from our analysis we can conclude that the combined use of LIRIC and the regularization technique improves our capability for evaluating the evolution of microphysical properties profiles during night- and day-time.

For a direct intercomparison of LIRIC with the regularization technique more simultaneous datasets are required. Advances in vibrational Raman will allow measurement of the extinction coefficients during the daytime and measurements from HSRL systems could also help in this aspect. On the other hand, advances in techniques to retrieve microphysical properties from star-photometry will allow the use of LIRIC during night. Improvements in the application of the Linear Estimation method also could be addressed in this way. Such approaches are essential both to study the temporal evolution of aerosol microphysical profiles and also to define strategies for evaluating satellite products.

Profiles of linear particle depolarization retrieved with the lidar data and volume concentration retrieved with LIRIC were compared with CASA C-212-200 aircraft measure-

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ments. This study represents the first time that in-situ measurements of particle depolarization ratio have been compared with lidar measurements. The CAS-POL measurements were compared with linear particle depolarization retrieved with the lidar data. The two separate techniques produces results that were within the estimated un-5 certainties over the whole vertical extent of the measurements. Similarly, the vertical profiles of the volume concentration, retrieved from the combined PCASP-100X and CAS-POL size distributions and LIRIC retrievals were very similar in general shape, i.e. maxima at 3200-3500 and 4100-4400 m were identified by both techniques. The average values were within the estimated uncertainties with discrepancies generally below 20 µm<sup>3</sup> cm<sup>-3</sup> when the same size range was considered for both techniques. The volume concentrations from the CAS-POL are likely overestimates due to the asphericity of the dust particles but the LIRIC derived values are underestimates because of the presence of particles with equivalent optical diameters larger than 20 µm.

The synergy of instruments and different techniques presented in this study reveals the potential to obtain a complete characterization of the aerosol properties using a combination of the measurements. However, future efforts should aim to reduce the uncertainty of the measured and retrieved aerosol properties in order to obtain more reliable and accurate aerosol properties databases.

### Appendix: CAS-POL measurement principles and uncertainties

The CAS-POL measures the light that is scattered by individual particles that pass through a focused, 658 nm, polarized laser beam (Baumgardner et al., 2001). Optical components are positioned to collect some of the light that is scattered into a forward cone from 4-12° and a backward cone from 168-176°. The backward scattered light is divided into two components: one that is measured with a detector that is behind an optical filter that passes only the scattered light with polarization that is perpendicular to the polarization of the incident light and another detector with no filter (Glen and Brooks, 2013). The equivalent optical diameter (EOD) of each particle in the nominal

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size range from 0.6-50 µm is derived from the peak intensity of the collected forward scattered light using Mie theory (Mie, 1908) and an assumption of particle sphericity at a known refractive index. The terminology, EOD, is used here to underscore that ambient aerosol particles are only spherical if they are liquid or have a liquid coating 5 and the refractive index can vary over a wide range. Hence, the EOD refers to the size of a spherical particle with known refractive index that would have scattered the equivalent intensity of light.

The three signals, forward scattering (FS), backward scattering (BS) and polarized (POL) provide three pieces of information that, from a relative perspective, can differentiate regions of air masses that have different particle characteristics. Although there are currently studies underway that will provide for a more quantitative interpretation of this, for the analysis used in the current study the metrics derived from these signals will be used to compare particle properties in the boundary layer and the free troposphere with and without a dust layer.

The interaction of the linearly polarized laser radiation with a particle leads to scattered light with some of the polarization in the same plane as the incident light and some fraction at a perpendicular plane (Nicolet et al., 2007, 2010, 2012; Schnaiter et al., 2012; Baumgardner et al., 2014). In laboratory studies, Glen and Brooks (2013) showed that the relationships between the FS, BS and POL signals were related to the type of dust, i.e. hematite, guartz and zeolite could be clearly differentiated by comparing the three signals. In the analysis of CAS-POL measurements in the current study, we will only report the depolarization ratio (defined below).

The measurement uncertainties are associated with the accuracy of determining the sample volume and the derivation of the EOD. The estimated uncertainty in the sample volume is ±20% primarily due to the optical technique used to qualify particles within the beam (Baumgardner et al., 2001). The sizing uncertainty ranges from 20-50% and depends on the variation in the refractive index and the asphericity of the particles. Figure 9a shows the theoretical scattering cross section (Mie, 1908) of spherical particles as a function of diameter for three refractive indices and the collection angles

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of the CAS-POL. Only a small size range is shown in order to illustrate the potential magnitude of uncertainty. The dashed lines show that particles with four different sizes have the same scattering cross section. The scattering intensity is directly proportional to the particle optical cross section; hence, when we measure scattering of this magni-5 tude with the instrument, we do not know if the particle was a water droplet with 1.25, 2.0 or 2.5 µm EOD, a salt particle (NaCl) with a 1.75 or 2.5 µm EOD or a sulfate particle (refractive index of 1.44) with an EOD of 2.5 µm.

The average uncertainty due to deriving an EOD of unknown refractive index was estimated by analyzing the variation in size when assuming a particle had a refractive index of 1.48 then finding what particle size would have the same scattering cross section if it actually had a different refractive index. The calculation was carried out over a range of refractive indices from 1.33 (water) to 1.60 (some types of organics). Figure 9b presents the results of this evaluation where the ordinate is the average derived value for the given size at a refractive index of 1.48 and the vertical bars are one standard deviation around the mean. The red lines are the one-to-one (solid) and ±20 % (dashed) around the one to one. From this figure we see that the uncertainty falls within ±20% of the assumed values except for the EOD between 2 and 4 μm. Hence, the uncertainty due to the refractive index variations is on average ±20%.

The uncertainties that are related to the asphericity of a particle are more difficult to estimate since they are dependent on the complexity of the morphology, the degree of asphericity and the orientation of the particle when it passes through the beam. Borrmann et al. (2000) applied T matrix theory to estimate the amount of undersizing of prolate and oblate spheroids dependent on their aspect ratios. The change in polarization caused by a particle is also dependent on aspect ratio and orientation. Baumgardner et al. (2014) ran simulations using T matrix theory to calculate the depolarization ratio as a function of aspect ratio, size and orientation. The depolarization ratio is defined here as the signal from the polarization detector divided by intensity of scattered light measured by the total backscatter detector (Glen and Brooks, 2013). The variation in the shape and orientation of the particle in the beam leads to uncer-

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tainties of the depolarization ratio around 30%, depending on the aspect ratio of the dust. Figure 10 illustrates the response of the CAS-POL to a 2 µm particle with aspect ratio of 0.9 as a function of its orientation in the beam (alpha and beta designate the angles with respect to the incident laser beam). From this figure we observe that the depolarization ratio can vary from almost zero to more than 0.15, depending on how it is oriented. With no other information available to determine a particle's asphericity, the estimated sizing uncertainty due to the assumption of sphericity is of the order of 50%.

The errors in the sample volume and size are propagated using the root sum squared (RSS) approach to estimate the uncertainty in the derived bulk parameters of number and volume concentrations and the median volume diameter. These estimated uncertainties are  $\pm 20$ ,  $\pm 90$  and  $\pm 50$  %, respectively.

Acknowledgements. This work was supported by the Andalusia Regional Government through projects P12-RNM-2409 and P10-RNM-6299, by the Spanish Ministry of Science and Technology through projects CGL2010-18782, and CGL2013-45410-R; by the EU through AC-TRIS project (EU INFRA-2010-1.1.16-262254); and by the University of Granada through the contract "Plan Propio. Programa 9. Convocatoria 2013". CIMEL Calibration was performed at the AERONET-EUROPE calibration center, supported by ACTRIS (European Union Seventh Framework Program, (FP7/2007–2013) under grant agreement no. 262254). Granados-Muñoz was funded under grant AP2009-0552. The authors thankfully acknowledge the computer resources, technical expertise, and assistance provided by the Barcelona Supercomputing Center for the BSC-DREAM8b model dust data. The authors express gratitude to the NOAA Air Resources Laboratory for the HYSPLIT transport and dispersion model and those at the NRL-Monterey that helped in the development of the NAAPS model. We also express our gratitude to the developers of LIRIC algorithm and software. We also thank N. T. O'Neill for providing the Spectral Deconvolution Algorithm used in the star-photometer. Thanks are also due to INTA Aerial Platforms, a branch of the Spanish ICTS program, and the Spanish Air Force for their efforts in maintaining and operating the aircraft. We would like to thank Ping Yang of Texas A&M University for providing the results of his simulations of light scattering from different types of ice crystals.

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**Table 1.** Aerosol properties on region 1, corresponding to the mixture of anthropogenic aerosol and mineral dust below 2250 ma.s.l. and region 2, corresponding to the mineral dust particles located above 2250 ma.s.l.

(1)	oaer 532 nm Vm <sup>-1</sup> sr <sup>-1</sup> )		β-AE (355–532 nm)	r <sub>eff</sub> (μm)	ν (μm <sup>3</sup> cm <sup>-3</sup> )
Region 1 3		0.15	0.18	0.61	44
Region 2 8		0.25	0.05	1.23	54

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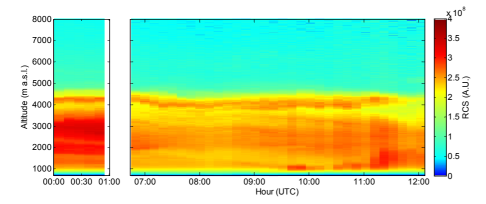
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**Figure 1.** Time series of the RCS for the period 00:00–01:00 UTC and 06:45–12:10 UTC on the 27 June 2011.

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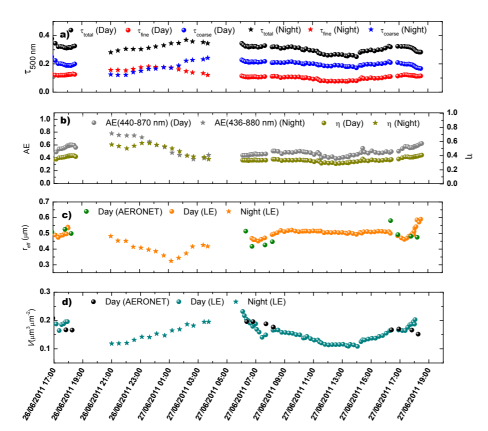
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**Figure 2.** Night-to-day temporal evolution of **(a)** aerosol optical depth  $(\tau_{\lambda})$  including also its separation between fine  $(\tau_{\rm fine})$  and coarse  $(\tau_{\rm coarse})$  mode at 500 nm, **(b)** Ångström exponent (AE) between 440 and 870 nm (436–880 nm for star photometry) and fine mode fraction  $\eta$  **(c)** effective radius  $(r_{\rm eff})$  and **(d)** column integrated volume concentration (V) on the 27 June 2011.

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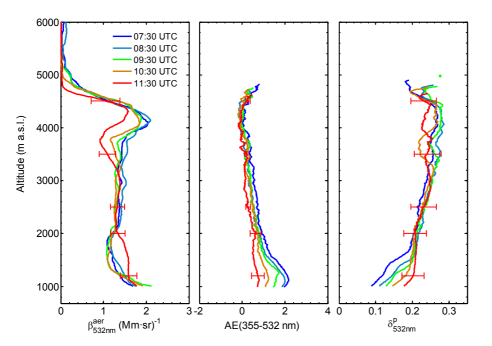


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**Figure 3.** Profiles of **(a)** aerosol particle backscatter at 532 nm ( $\beta_{532\,\mathrm{nm}}^{\mathrm{aer}}$ ) **(b)** backscatter-related Ångström exponent between 355 and 532 nm ( $\beta$ -AE(355–532 nm)) and **(c)** linear particle depolarization ratio at 532 nm ( $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$ ) retrieved from lidar elastic measurements at different hours between 07:30 and 11:30 UTC on the morning of the 27 June 2011.

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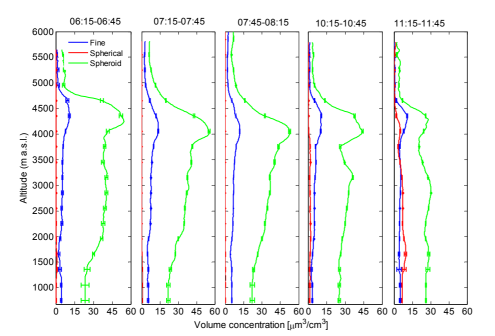


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**Figure 4.** Volume concentration profiles of the fine, coarse spherical and coarse spheroid modes obtained with LIRIC from 30 min averaged lidar data for different periods on the 27 June 2011. The error bars are obtained as indicated in Granados-Muñoz et al. (2014).

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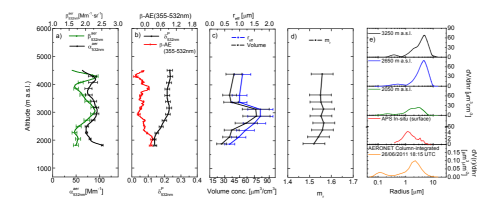
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**Figure 5.** (a)  $\beta_{532\,\mathrm{nm}}^{\mathrm{aer}}$  and  $\alpha_{532\,\mathrm{nm}}^{\mathrm{aer}}$  retrieved with Raman technique and (b) derived  $\beta$ -AE(355– 532 nm) and  $\delta_{532\,\mathrm{nm}}^{\mathrm{P}}$  at 00:00–01:00 UTC on the 27 June 2011. (c) Total volume concentration,  $r_{\rm eff}$  and (d)  $m_{\rm r}$  retrieved for the same period with the regularization technique applied to the  $3\beta + 2\alpha + 1\delta$  lidar data and (e) Volume size distributions retrieved at different height levels using the APS (surface level) and the regularization technique applied to the  $3\beta + 2\alpha + 1\delta$  lidar data at 00:00-01:00 UTC on the 27 June 2011. Column-integrated AERONET size distribution corresponding to the 26 June 2011 at 18:15 UTC is included (bottom).



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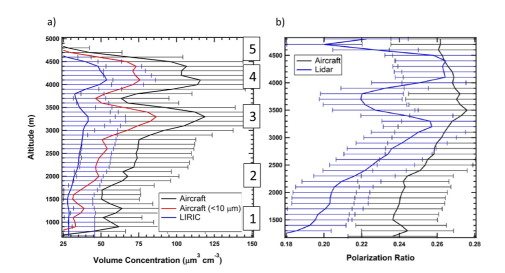


Figure 6. (a) 30 min averaged volume concentrations retrieved with LIRIC (blue line) centered at 10:30 UTC and the combined PCASP-100X and CAS-POL concentrations for particles smaller than 10 µm radius (red line) and for the complete size distritubion (black line) retrieved during the flight ascent on the 27 June 2011. (b) Depolarization ratio retrieved from the CAS-POL measurements (black line) and  $\delta P$  532 nm retrieved from the lidar data (blue line) between 10:30 and 11:00 UTC. Horizontal bars show the estimated uncertainties.



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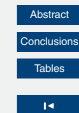
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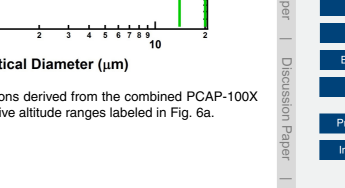
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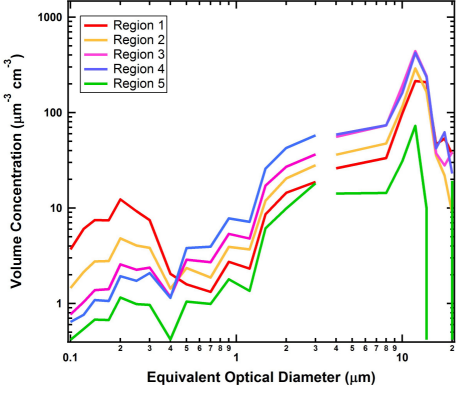
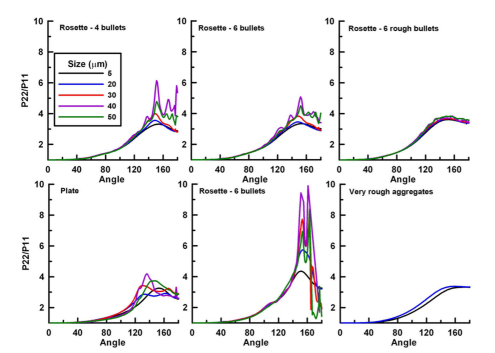


Figure 7. Size distributions of volume concentrations derived from the combined PCAP-100X and CAS-POL measurements averaged over the five altitude ranges labeled in Fig. 6a.



**Figure 8.** Theoretical calculations of the light scattering at 680 nm from individual ice crystals of different shapes and sizes (expressed in diameter) showing the ratio of the  $P_{22}$  and  $P_{11}$  components of the Stokes scattering matrix (calculations courtesy of Ping Yang, Texas A&M University).

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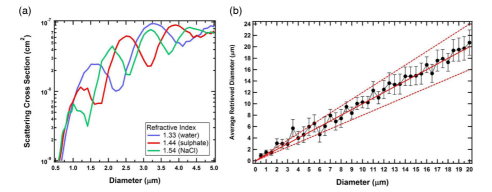
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**Figure 9. (a)** Theoretical scattering cross section (Mie, 1908) of spherical particles as a function of diameter for three refractive indices and the collection angles of the CAS-POL. Only a small size range is shown in order to illustrate the potential magnitude of uncertainty. The dashed lines show that particles with four different sizes have the same scattering cross section. **(b)** Average retrieved diameter for a given size at a refractive index of 1.48. Vertical bars represent one standard deviation around the mean. The red lines are the one-to-one (solid) and  $\pm 20\%$  (dashed) around the one to one.

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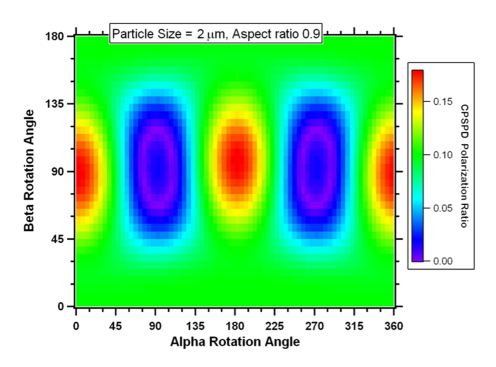


Figure 10. Response of the CAS-POL to a 2 µm particle with aspect ratio of 0.9 as a function of its orientation in the beam (alpha and beta represent the angles with respect to the incident laser beam).

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