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Validation of LIRIC aerosol concentration retrievals using airborne measurements during a biomass burning episode over Athens



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ABSTRACT

In this paper we validate the Lidar-Radiometer Inversion Code (LIRIC) retrievals of the aerosol concentration in the fine mode, using the airborne aerosol chemical composition dataset obtained over the Greater Athens Area (GAA) in Greece, during the ACEMED campaign. The study focuses on the 2nd of September 2011, when a long-range transported smoke layer was observed in the free troposphere over Greece, in the height range from 2 to 3 km. CIMEL sun-photometric measurements revealed high AOD (~0.4 at 532 nm) and Ångström exponent values (~ 1.7 at 440/870 nm), in agreement with coincident ground-based lidar observations. Airborne chemical composition measurements performed over the GAA, revealed increased CO volume concentration (~ 110 ppby), with 57% sulphate dominance in the PM₁ fraction. For this case, we compare LIRIC retrievals of the aerosol concentration in the fine mode with the airborne Aerosol Mass Spectrometer (AMS) and Passive Cavity Aerosol Spectrometer Probe (PCASP) measurements. Our analysis shows that the remote sensing retrievals are in a good agreement with the measured airborne in-situ data from 2 to 4 km. The discrepancies observed between LIRIC and airborne measurements at the lower troposphere (below 2 km), could be explained by the spatial and temporal variability of the aerosol load within the area where the airborne data were averaged along with the different time windows of the retrievals.

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1. Introduction

According to the report of the Intergovernmental Panel on Climate Change (IPCC; Myhre et al., 2013), anthropogenic aerosols result into

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a net cooling globally through their interaction with radiation and clouds, by an amount that remains difficult to quantify accurately and which could be comparable in magnitude to the net warming effect of greenhouse gases. Moreover, according to the World Health Organization (WHO), there is a significant linkage between suspended particles and human's mortality (WHO, 2006). Aerosol effects are determined, among others, by particle's size, their chemical composition, and number concentration. To understand how atmospheric particles are affecting the Earth's climate, the scientific community has established and operates global networks equipped with active and passive remote

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sensing instrumentation. More precisely, the AErosol RObotic NETwork (AERONET) provides almost real-time columnar aerosol optical and microphysical properties, based on the operation of >400 sun-sky radiometers distributed worldwide (Holben et al., 1998). Several aerosol lidar networks are used for aerosol/cloud research including well established infrastructures and networks like: the European Aerosol Research Lidar Network (EARLINET; Pappalardo et al., 2014), the Micro Pulse Lidar Network (MPLNET; Welton and Campbell, 2002) and the Asian Dust and aerosol lidar observation network (AD-Net; Sugimoto et al., 2014).

Depending on the capabilities of each system and the employed techniques, lidar measurements are used to retrieve the vertical distribution of (a) the aerosol backscatter coefficient (β_{qer}), (e.g. Fernald et al., 1972; Klett, 1981); (b) the aerosol extinction coefficient (a_{aer}), (Ansmann et al., 1990; Ansmann et al., 1992) and (c) the particle linear depolarization ratio δ_{aer} (e.g. Cairo et al., 1999; Sassen, 2005; Freudenthaler et al., 2009). The aerosol backscatter and extinction coefficients can be retrieved from the backscatter, Raman (Ansmann et al., 1992; Whiteman et al., 1992; Whiteman, 2003), or High Spectral Resolution Lidar (HSRL) technique (e.g. Eloranta, 2005). The selected technique defines also the accuracy of the retrieved aerosol products. Many studies have demonstrated that the spectral information of the aforementioned aerosol optical properties, makes feasible the provision of accurate retrievals in the fine mode, regarding aerosol microphysical parameters namely: aerosol size distribution, aerosol effective radius, number and volume concentration (Müller et al., 1999; Veselovskii et al., 2002; Veselovskii et al., 2010). In case that the aerosol extinction coefficient is provided by Raman technique, the microphysical retrievals are typically limited to night-time measurements since their accuracy depends on the error of the optical parameters provided as initial inputs.

During the last years, there are increased efforts to retrieve aerosol concentration profiles also during day-time. For example, the polarization lidar photometer networking (POLIPHON) technique is capable of retrieving the concentration profiles of dust and non-dust particles using single wavelength backscatter and depolarization coefficient lidar profiles (Ansmann et al., 2011; Ansmann et al., 2012). In this technique, AERONET microphysical retrievals are used to provide columnar volume-to-AOD values needed to convert the optical properties to concentration. On a continuation effort, Mamouri and Ansmann (2014), have expanded the technique to separating the contribution of fine and coarse dust modes, based on laboratory measurements of fine and coarse dust depolarization ratios.

In the framework of Aerosols, Clouds, and Trace gases Research Infrastructure (ACTRIS), two algorithms have been developed for retrieving concentration profiles from the synergy of lidar and sunphotometric measurements. The Generalized Aerosol Retrieval from Radiometer and Lidar Combined (GARRLiC; Lopatin et al., 2013) inversion algorithm retrieves vertical profiles of both fine and coarse aerosol concentrations as well as the size distribution and complex refractive index for each mode. Based on similar approach, the LIdar-Radiometer Inversion Code (LIRIC; Chaikovsky et al., 2016) considers that the fine and coarse particle intensive properties are constant with height, taken equal to the column-integrated values provided by AERONET, with only their concentration varying along the atmospheric column. LIRIC, GARRLiC and POLIPHON techniques have been used by many EARLINET-AERONET stations, during large and medium scale dust events over the European continent, for evaluating dust model performance in terms of dust layer geometrical properties (height range and centre of mass) as well as dust load (particle concentrations) (e.g. Binietoglou et al., 2015; Granados-Muñoz et al., 2016a, 2016b). For a case study of Saharan dust outbreak over Athens, Greece, Tsekeri et al. (2013), found a satisfactory agreement between LIRIC output and dust concentration profiles, simulated by the regional dust model BSC-DREAM8b (Pérez et al., 2006a; Pérez et al., 2006b; Basart et al., 2012). Furthermore, comparisons of LIRIC output, with dispersion models of other aerosol types than dust (e.g. volcanic dust), like Lagrangian dispersion model FLEXPART (Stohl et al., 1998; Stohl et al., 2005), showed a Pearson's coefficient (R) varying from 0.69 to 0.84 (Kokkalis et al., 2013). Moreover, Wagner et al. (2013), inter-compared LIRIC and POLIPHON concentration profiles, by applying both techniques on two case studies of irregularly shaped particles in the atmosphere (i.e. one Sahara dust outbreak and one volcanic dust event). The comparison between the two techniques revealed acceptable agreement, however the potential of LIRIC to retrieve optical properties, namely particle backscatter coefficient, lidar ratio and Ångström exponent, was found to demonstrate systematic deviations compared to corresponding measurements obtained with a Raman lidar. In addition, Papayannis et al. (2014) showed that the relative difference between LIRIC and POLIPHON mass concentration retrievals, is in the range of \pm 20% for the case of coarse non-spherical particles.

Furthermore, LIRIC retrievals of volume concentration have been compared with aircraft in-situ measurements, by Granados-Muñoz et al. (2016a) during a Saharan dust episode over Granada, Spain. The case study of coarse mode (dust), non-spherical particles, is introducing limitations regarding, the potential of in-situ instrumentation to measure size distribution in the size range above 3 µm (diameter). Thus, during their study, they combined: in-situ depolarization measurements from Cloud and Aerosol Spectrometer with Polarization detection (CAS-POL; Baumgardner et al., 2001), operating at the size range 0.6-50 µm (diameter), and a Passive Cavity Aerosol Spectrometer (PCASP 100×; Rosenberg et al., 2012; Cai et al., 2013) measuring aerosol size distribution in 0.1–3 µm diameter range. Correcting the retrieved size distributions, for refractive index assumptions, they demonstrated volume concentration discrepancies $<20 \, \mu m^3 \, cm^{-3} \, (0.02 \, ppbv)$ and they attributed them to CAS-POL overestimation due to the asphericity of dust particles and to the possible underestimation of LIRIC, despite the fact that the derived size distributions from CAS-POL and lidar, were found to be in a good agreement.

In this study, we validate the volume concentration retrieved by LIRIC, with independent in-situ measurements of chemical composition, for a case of predominant fine mode particles in the atmosphere, over the Greater Athens Area (GAA; Saronic Gulf, Evoikos Gulf and Aegean), Greece. To our knowledge, this is the first time that fine mode LIRIC retrievals are validated against airborne. The case of fine mode particles is favourable for comparing remote sensing and in-situ observations since in this case there are fewer limitations on the instrumental side for the in-situ measurements while for remote sensing part the Mie scattering simulations are applicable since the particles are mainly spherical. In Section 2 we present the instrumentation and methodology used. Section 3 presents a brief description of ACEMED campaign along with the case study used for the evaluation of the LIRIC fine mode aerosol concertation retrieval. Our analysis contains, a thorough characterization of the aerosol load monitored over the GAA, in terms of their optical properties and chemical composition. In the second part of Section 3 we compare the retrieved concentrations with the independent in-situ airborne measurements. Finally, our conclusions are given in Section 4.

2. Instrumentation and method

2.1. Backscatter-depolarization lidar

At the National Technical University of Athens (NTUA, 37.97° N, 23.79° E, elevation: 212 m) a six-wavelength Raman-backscatter lidar system (EOLE) operates since February 2000, as a member of the EARLINET network (Bösenberg et al., 2003; Pappalardo et al., 2014). The emission unit is based on an Nd:YAG laser, emitting high energy laser pulses at 355, 532 and 1064 nm with a repetition rate of 10 Hz. The respective emitted energies per pulse are of the order of 240, 300 and 260 mJ. A Galilean type beam expander (\times 3) is mounted, just before the emission of the laser beam in the atmosphere, for reducing the laser beam divergence and increasing the beam diameter, almost

with the same efficiency for all the emitted wavelengths. The optical receiver is based on a Cassegrainian telescope with 600 mm focal length and clear aperture diameter 300 mm, directly coupled with an optical fiber, to the wavelength separation unit, detecting finally signals at 355, 387, 407, 532, 607 and 1064 nm.

During day time operation, the system is capable to provide aerosol backscatter profiles (β_{aer}) at 355, 532 and 1064 nm, based on the standard backscatter lidar technique and employing the Klett inversion method (Klett, 1981). The technique assumes, the existence of an aerosol-free region (e.g. upper troposphere), and a linear relationship between aerosol backscatter and extinction coefficient, the so called lidar ratio (S_{qer}) , constant in the laser-telescope path. A variety of studies performed in the framework of EARLINET, revealed a wide range for the lidar ratios, covering values from 20 to 100 sr (Ackermann, 1998; Mattis et al., 2004; Amiridis et al., 2005; Müller et al., 2007; Papayannis et al., 2008; Amiridis et al., 2009; Groß et al., 2011; Groß et al., 2013; Groß et al., 2015; Giannakaki et al., 2015). In elastic backscatter lidar technique, the assumption of a constant lidar ratio value throughout the laser sounding range, is the most critical for solving the lidar equation, while the overall uncertainty, including both statistical and systematic errors, on the retrieved β_{aer} values, is of the order of 20-30% (Bösenberg et al., 1997; Comerón et al., 2004; Rocadenbosch

Next to EOLE system, a depolarization lidar system (AIAS), was also operating continuously during September 2011. AIAS is capable to detect both the parallel- and perpendicular- components of the backscattered light at 532 nm, with respect to the linear polarization plane of the initially emitted laser beam. The emission unit of AIAS is based on a Nd:YAG laser, emitting short laser pulses at 532 nm with energy of the order of 95 mJ per pulse. The backscattered light is collected by a Dall-Kirkham/Cassegrainian telescope with 1000 mm focal length and 200 mm clear aperture diameter, and guided to a Polarization Beam Splitter Cube (PBSC) where the two polarization components are separated and directed to the corresponding detectors (Sassen, 2005). The calibrated ratio of these two components is known as volume depolarization ratio, and the key for deriving accurate particle linear depolarization measurements (δ_{aer}) lies in obtaining a reliable calibration of the lidar system. Various calibration methods exist in the literature (e.g. Biele et al., 2000; Reichardt et al., 2003; Alvarez et al., 2006), however the $\pm 45^{\circ}$ calibration technique (Freudenthaler et al., 2009), has been employed in our case.

Both lidar systems are affected by the overlap height and below this region are not capable to provide trustworthy aerosol products. The geometrical specification of EOLE system makes feasible the full overlap of the laser beam with the receiver field of view to be reached at heights 700–900 m above ground (Kokkalis et al., 2012). Regarding the depolarization lidar AIAS, the measured volume depolarization ratio is reliable to about 50 m above ground since overlap effects widely cancel out due to the signal ratios needed for the calculation of the depolarization ratio. However, in this study the data are analysed for heights above 850 m. Below that height we set the measured aerosol related physical quantities to height-independent values.

2.2. CIMEL sun-sky radiometer

In this study, the reported columnar aerosol optical properties, have been retrieved by a CIMEL sun-sky radiometer (Holben et al., 1998), located on the roof of the Research Center for Atmospheric Physics and Climatology of the Academy of Athens (37.99° N, 23.78° E, elevation: 130 m). The radiometric station is in the city centre, approximately 10 km away from the coastal line and 1.6 km North from the lidar station. The instrument is part of NASA's global sun photometric network AERONET, and is capable to perform automatic measurements of the direct solar irradiance at the common wavelengths of 340, 380, 440, 500, 675, 870, 940 and 1020 nm and diffuse sky radiance at 440, 675, 870 and 1020 nm, respectively. Those measurements are further used to

provide, both optical and microphysical aerosol optical properties in the atmospheric column (Dubovik and King, 2000; Dubovik et al., 2006). The CIMEL data used in this study, are the cloud screened and quality assured level 2.0 data products, providing information regarding the columnar aerosol optical depth (AOD), fine and coarse mode fractions of AOD at 500 nm, the particle volume size distribution in the size range of 0.05 to 15 μm in terms of particle radius, and the Ångström exponent. The separation of fine and coarse, size distribution is done by finding the minimum concentration values in the particle's radius range 0.194–0.576 μm . The AOD uncertainty is <±0.02 for UV wavelengths and <±0.01 for wavelengths longer than 440 nm (Eck et al., 1999). The uncertainty of the aerosol size distribution retrieved by the sky radiance measurements is based on the calibration uncertainty of each wavelength, assumed to be <±5%.

2.3. Lidar-sun-photometric inversion algorithm (LIRIC)

The LIRIC algorithm has been developed by the Belarusian Institute of Physics in Minsk, in collaboration with the French Laboratoire d'Optique Atmosphérique in Lille. LIRIC combines the elastically backscattered lidar signals at 355, 532 and 1064 nm, along with the radiometric measurements from CIMEL, and is capable of retrieving the fine $(C_f(z))$ and coarse mode particle volume concentration profiles $(C_c(z))$ (Chaikovsky et al., 2016), in parts per billion volume (ppbv). In case that depolarization measurements are available, LIRIC algorithm is capable to provide also the concentration profiles of coarse spherical and coarse spheroid modes. However, in the present study this capability has not been examined, since an older version of the LIRIC code has been used. The threshold value for defining fine and coarse mode particles in terms of their size, is obtained from the columnar volume size distribution retrieved by AERONET inversion algorithms. Thus, in the case of LIRIC, fine mode particles, may assume to be particles with radius approximately <0.5 µm. Finally, the retrieval is based on a maximumlikelihood estimation of the concentration profiles so that the lidar signals are reproduced within their measurement uncertainty and the integral of the retrieved aerosol concentrations matches the total volume concentration of the fine and coarse modes, derived from sunphotometric measurements.

LIRIC's retrieval uncertainty, depends both on the regularization parameters defined by the operator, as well as on the uncertainty of the input data. More precisely, for various atmospheric conditions, lidar geometrical characteristics, and LIRIC's regularization parameters, the uncertainties in the retrieved aerosol parameters were found to be maximum 30%, for cases of complex aerosol structures, however taking values below 10%, during cases of homogeneous aerosol mixing and simple aerosol structures (Granados-Muñoz et al., 2014). Uncertainties due to the regularization parameters were found to be below 2%. Overall, according to Chaikovsky et al. (2016), the retrieved volume concentration profiles may be provided with a standard deviation in the range 5–20% of the maximum aerosol layer concentration, with the concentration uncertainty to be significant only for cases of low concentration values. In the studied case, we considered that the retrieved concentration profiles are provided with an overall uncertainty of 20%.

2.4. In-situ airborne aerosol instrumentation

The size-resolved chemical composition and mass concentration, have been measured with an Aerodyne time-of-flight Aerosol Mass Spectrometer (TOF-AMS; Allan et al., 2003; Canagaratna et al., 2007; Morgan et al., 2010). The AMS is capable to measure the fine mode particle's mass concentration (M), of the following chemical compounds: sulphates (SO_4) , nitrates (NO_3) , ammonium (NH_4) , chloride (Cl) and organic matter (OM). Particle's aerodynamic diameter, are related to their physical diameters, ranging usually between 0.1 and 1 μ m. The initially reported mass concentrations, are measured in

micrograms per standard cubic meter (μ g Sm⁻³), at standard temperature-pressure conditions (T = 273.15 o K and P = 1013.25 hPa).

Volume concentration of gaseous pollutants, have been also measured during the flights, using the following gas analysers: Aero-Laser (AL) 5002 VUV Fast Fluorescence analyser, for carbon monoxide (CO) measurements, and a Thermo-Electron (TE) 49C UV photometric analyser, for ozone (O_3) measurements.

The size-resolved particle's number density has been measured with a Passive Cavity Aerosol Spectrometer Probe (PCASP). The instrument is capable to measure particle's number size distribution, over the diameter range of 0.1 to 3 µm, in 30 size bins. The principle of operation of PCASP is based on the scattering of a laser light from suspended particles. The scattered radiation over an angular range of 35°-120° (primary angle) and 60°-145° (secondary angle), is collected by a parabolic mirror and is focused onto a photodetector, producing an electronic pulse. The pulse height for each particle is linearly related to a particle's cross section and therefore nonlinearly related to the particle's diameter. The number of pulses counted per second is proportional to the concentration. However, since scattering cross section depends among other parameters also on particle's shape and refractive index, a correction regarding refractive index have to be applied. Thus, for deriving calibrated particle's number density distributions we followed the procedure demonstrated by Rosenberg et al. (2012), and applied also by Tsekeri et al. (2016), for ACEMED campaign dataset.

Finally, temperature sensors (Rosemount/Goodrich) and hygrometers (Lyman-alpha) mounted on the aircraft, are capable to provide measurements of temperature and relative humidity. The overall calibration uncertainties of the temperature measurements are of the order $\pm\,0.3$ °K.

2.5. Modelling

Source-receptor relationships between the measurement areas and the potential emission sources, are investigated with the use of the particle dispersion model FLEXPART-WRF (Stohl et al., 2005; Brioude et al., 2013). The model is driven by WRF_ARW (Skamarock et al., 2008) meteorological fields at a resolution of 12×12 km over the area of interest. A two-way nested 3×3 km grid over the greater Athens area is also enabled for the description of the local sea breeze flow during the experimental period. Initial and boundary conditions for WRF are from the National Center for Environmental Prediction (NCEP) final analysis (FNL) dataset at $1^{\circ} \times 1^{\circ}$ resolution. The sea surface temperature (SST) is from the NCEP $1^{\circ} \times 1^{\circ}$ analysis and a total of 10.000 tracer particles are assumed for each release in FLEXPART simulations. Backward trajectories and emission sensitivity studies during the sampling period indicate the possible sources for the aerosol layers, detected both by in-situ and remote sensing instruments.

3. Results from ACEMED campaign and discussion

During the period from 1st to 9th of September 2011, the European Fleet for Airborne Research (EUFAR) supported airborne measurements over a wide domain over Greece by deploying the FAAM-BAe146 research aircraft. Two flights, one on the 2nd and one on the 9th of September were performed in the framework of the ACEMED experimental campaign, in order to retrieve detailed information about the physical and chemical aerosol properties, along the flight track of the CALIPSO satellite. ACEMED campaign, supported the collection of quality assured and coordinated ground-based, airborne in-situ and spaceborne measurements, to generate representative case studies that will be further used to study the aerosol type classification scheme, applied on CALIPSO dataset.

This study is focused on the 2nd of September flight (FAAM flight ID number B638), where the research aircraft took-off from Chania (Crete Island), covering areas from Southern Greece to Athens, at various heights. The flight duration was approximately 4 ½ hours (~ 08:00–

12:30 UTC), and the aircraft flight track is depicted in Fig. 1-a with the black dashed line. The green circle, of 220 km radius in Fig. 1-a, is depicting the GAA used for the spatial and temporal averaging of the on-board in-situ measurements. The GAA, is further zoomed in Fig. 1-b, presenting also the locations of the ground based instrumentation (i.e. stations of sun-sky radiometer and lidar), along with the B638 flight track. Unfortunately, the flight of 9th of September, was performed during night time, and thus cannot be used for the validation of LIRIC. Additional information regarding ACEMED and concurrent campaigns (Aegean-Game and CarbonExp) may be found in Bezantakos et al. (2013), Tombrou et al. (2015) and Tsekeri et al. (2016).

During the entire ACEMED campaign period, high values of fine mode AOD were observed by the CIMEL sun-sky radiometer. The fine mode fraction of columnar AOD at 500 nm was >89%, reaching the highest values of 91% and 93% (Fig. 2), on 2nd and 3rd of September 2011, respectively. Moreover, large Ångström exponent values were observed during this 8-day period, varying from 1.30 to 1.60 (Fig. 2). The similarity in optical properties observed with AERONET are indicative of the presence of a consistent aerosol type over Greece, during the entire campaign period, especially between, 2nd-3rd and 7th-9th of September.

3.1. Case study description

For the 2nd of September 2011 (case study examined here), we present in Fig. 3, the temporal variability of the columnar aerosol optical properties from AERONET. The Ångström exponent varied from 1.77 to 1.90 with a mean value of 1.84 \pm 0.02, while the AOD at 340 and 500 nm increased during morning and noon and remained constant during afternoon hours, with mean values of 0.82 \pm 0.01 and 0.44 \pm 0.01 respectively (Fig. 3). The same pattern of the AOD is followed also by the columnar water vapor (in cm) as depicted by AERONET. An increase of the columnar water vapor is observed from around 07:00 UTC to 14:00 UTC varying from ~2.00 to 2.80 cm, demonstrating the high temporal variability of water content over the atmosphere (Fig. 3). Moreover, the volume size distribution retrievals from the photometric data indicate the presence of rather fine mode particles in the atmospheric column (Fig. 4). The dominance of fine mode particles is highlighted by the bi-modal size distribution with separation radius ranging from 0.44 um at 13:30 UTC to 0.57 um at 14:20 and 15:52 UTC.

Both backscatter and depolarization lidar systems (EOLE and AIAS) were operating during that day from 09:58 to 17:32 UTC. In Fig. 5-a we present the spatio-temporal evolution of the backscattered signal obtained by EOLE at 1064 nm, visualizing the evolution of a distinct aerosol layer extending from 2 up to ~3.5 km over the station. Cloud formation is also found at 3 km during 11:00-13:00 UTC. In Fig. 5-b the vertical profiles of relative humidity and potential temperature are demonstrated as obtained by radiosonde data on 2nd of September (11:22 UTC). The launching area is located at a coastal station around 15 km south-west of the lidar station. The transport of moister marine air due to the initiation of a sea breeze circulation results in RH increase reaching up to 60% at the layer 2–3 km. As seen also by the WRF model results in Fig. 5-c, the sea breeze flow penetrates the Attica basin and results in SSW surface winds exceeding 7 m s⁻¹ at 11:00 UTC. Mechanical elevation of these marine air masses along the Attica Mountains leads in the formation of the shallow orographic clouds that are found by the lidar measurements. This is also shown in Fig. 5-d for the area of NTUA lidar station where the model predicts RH values up to 100% at the layer 2-3 km during the period 12:00-16:00 UTC. Condensation is also evident in the model and the formation of a warm phase orographic cloud with a maximum mixing ratio of 0.15 g kg^{-1} is shown in Fig. 5-d around 11:00-13:00 UTC in accordance with the lidar observations.

In order to avoid the contamination of our retrievals with the presence of low altitude scattered clouds, developed at the top of that layer from 11:30 to \sim 13:00 UTC, we focused our analysis in the time window 13:00–15:00 UTC, indicated with a grey rectangle overlaid in

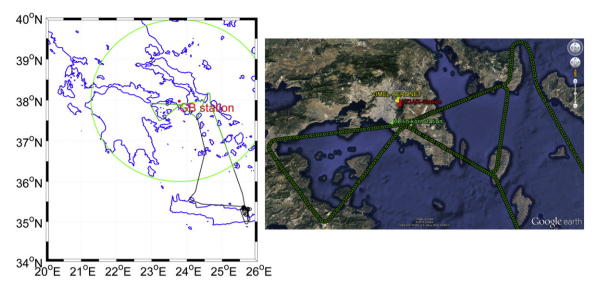


Fig. 1. (a) The aircraft ground track (dashed black line) during B638 flight and ground based station (red cross). The GAA is denoted with a green circle of 220 km radius. The GAA is zoomed in (b) denoting the flight track, and the locations of the ground-based instrumentations (i.e. sun-sky radiometric and lidar stations). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 5-a. For the aforementioned time period, the vertical profiles of aerosol backscatter coefficients ($\beta_{aer}(z)$) at 355, 532 and 1064 nm have been retrieved using the Klett inversion method (Fig. 6-a), using the lidar ratio value of 70 sr at 532 nm. Similar lidar ratio values pointed out by AERONET on the same day. More precisely, by using the values of single scattering albedo ($\omega(\lambda)$) and phase function at 180° ($P(180^\circ)$), retrieved with AERONET inversion algorithm, the mean daily columnar lidar ratio value at 532 nm, was found to be equal to 65 \pm 4 sr (Eq. 1).This lidar ratio value is typical of smoke (e.g. Amiridis et al., 2009; Müller et al., 2007).

$$S_{aer} = \frac{4\pi}{\omega(\lambda) \times P(180^{\circ})} \tag{1}$$

The backscatter-related Ångström exponent at VIS/IR is > 1.5, in agreement with the corresponding values retrieved from AERONET

(Fig. 6-b), denoting also the dominance of fine mode particles at higher altitudes. For the same time period, the particle linear depolarization ratio ($\delta_{aer}(z)$) profile at 532 nm, was calculated from AIAS lidar signals (Fig. 6-c). $\delta_{qer}(z)$ depicts an almost constant value of 4% (3.97 \pm 0.19%), for heights above 2 km, while at the height range between 1 and 2 km a mean value of 5.57 \pm 0.66% is observed. The aforementioned values are indicative for the presence of rather spherical particles. In summary, the high values of backscatter-related Ångström exponent along with the low particle linear depolarization ratio values in the height range ~ 2-3.5 km, are considered as typical for smoke as also reported by Burton et al. (2012) and Groß et al. (2013). In Fig. 6-d, the LIRIC retrievals from the synergy of lidar signals and sun-sky radiometric measurements are shown $(C_{f/c/t}(z))$. The aerosol fine mode is dominant in the entire vertical atmospheric profile and the contribution of the coarse mode is limited to ~25% of the total. More precisely, the fine mode aerosol volume is almost 85% larger than the coarse mode,

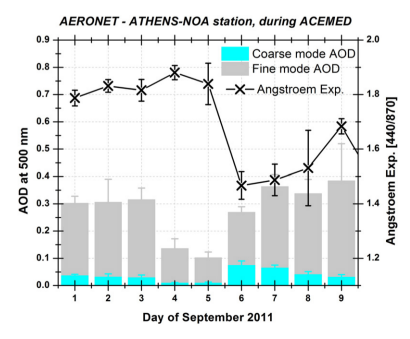


Fig. 2. AERONET fine/coarse mode AOD retrievals over Athens at 500 nm, for the time period between 1st-9th of September 2011, along with the Ångström exponent of 440 nm/870 nm for the same time period.

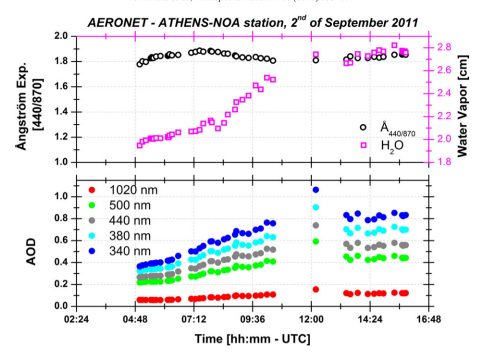


Fig. 3. Temporal variability of aerosol optical depth at 1020, 50, 440, 380 and 340 nm, and Ångström exponent (440/870) along with the columnar water vapor (in cm), retrieved by CIMEL-AERONET station in Athens, during 2nd of September 2011.

inside the free-tropospheric layer extended from 2 to approximately 3 km, reaching a maximum value of 0.03 ppbv.

The presence of smoke particles over GAA for our case study has also been identified from model simulations performed for our case. Specifically, we performed backwards emission sensitivity analysis employing the FLEXPART-WRF starting from three different areas inside the GAA (Saronic Gulf, Evoikos Gulf and Aegean Sea). The results indicate a homogeneous transport of air masses from the area of West Balkans. The active fires that are detected during the same time period at this area by MODIS (red triangles in Fig. 7) are likely the source for the biomass smoke layer, detected in our measurements.

On 2nd of September, and for the time period 09:15–10:38 UTC, the research aircraft FAAM-BAe146 flew at various heights from 1.6 up to 4.5 km, over the GAA, performing measurements of fine mode particle concentration (PM₁; particles with diameter $<1~\mu m$), with the on-

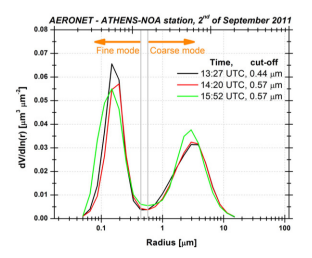


Fig. 4. Columnar aerosol volume size distributions retrieved by AERONET, on 2nd of September 2011 during noon hours. With grey vertical lines the cut-off points of the bimodal distributions, are also denoted.

board AMS instrumentation. The AMS measurements of Fig. 8-a show that the most dominant chemical components of the total measured PM_1 are sulphates and organic matter. The error bars shown for each chemical compound, correspond to the standard deviation due to the temporal averaging of the measured mass concentrations. Their low values indicate that the aerosol variability over the GAA was negligible. In the height range from 1.6 to 4.1 km, the fraction of sulphates in the total measured PM_1 was found to be 57.6% while the corresponding organic fraction is 31.6%. The high percentage of sulphate and organic species in the submicronic non-refractory total concentration could indicate the presence of smoke particles at this altitude, in accordance to the findings of Pratt et al. (2011) and Sun et al. (2015) and literature given therein. The smoke particles usually present increased fractions of organics and sulphates in both fresh and aged states according to Pratt et al. (2011).

More precisely, above the flight height of 2.5 km, the mass concentration of sulphates was $M_{SO_4}=6.60\pm4.59~\mu \mathrm{g}~m^{-3}$, while the corresponding mean values regarding organics, nitrates, chlorides and ammonium are found to be respectively, $M_{OM}=5.18\pm2.85~\mu \mathrm{g}~m^{-3}$, $M_{NO_3}=0.17\pm0.08~\mu \mathrm{g}~m^{-3}$, $M_{Cl}=0.012\pm0.005~\mu \mathrm{g}~m^{-3}$, $M_{NH_c}=1.48\pm0.75~\mu \mathrm{g}~m^{-3}$.

Simultaneous measurements of the gaseous pollutants revealed higher values of volume concentrations in the height range above 2 km, reaching even the 110 ppbv for CO and 52 ppbv for O₃ respectively (Fig. 8-b), indicating the presence of smoke in that height range. These results are consistent with similar results published by Verma et al. (2009), who reports values of O₃ concentration in the free troposphere of approximately 55 ppbv corresponding to the smoke plume, while CO concentrations ranges between 120 and 250 ppbv. Moreover, due to the low relative humidity (RH) values in our case study (<50%), no humidity corrections are needed for the airborne in-situ retrievals (Fig. 8-c). Similar values to our observations has been reported also by Tombrou et al. (2015) and Athanasopoulou et al. (2015).

3.2. Validation of volume concentration retrievals

LIRIC provides the concentration profiles of fine and coarse mode particles ($C_{f/c}(z)$) in ppbv. In contrast, the mass concentration profiles

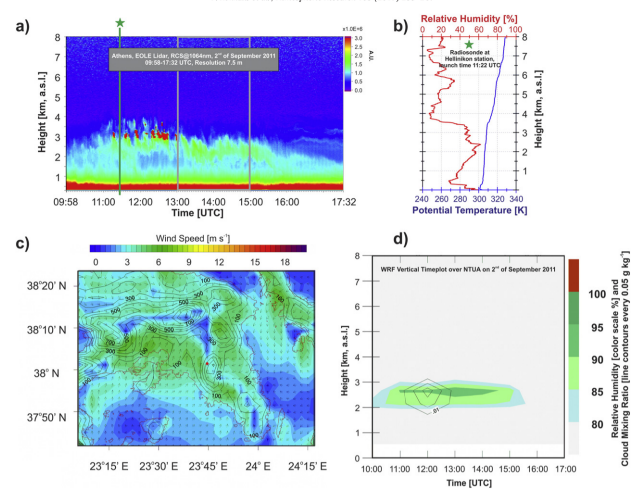


Fig. 5. (a) Spatio-temporal evolution of range corrected lidar signal (RCS) in arbitrary units (A.U.) at 1064 nm, between 09:58–17:32 UTC, during 2nd of September 2011. The two vertical grey lines refer to the time window of our analysis. (b) Potential temperature (K) and relative humidity – RH (%) vertical profiles obtained by radiosonde data on 2nd of September 2011 (11:22 UTC). (c) WRF simulated 10 m wind speed (m s⁻¹) and topographic contours every 100 m at 11:00 UTC. The red dot indicates the location of NTUA lidar station. (d) Vertical time-height plot of WRF simulated RH (%) and cloud mixing ratio (g kg⁻¹) above NTUA lidar station on 2nd of September 2011. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

 $(M_i(z))$ retrieved by the AMS instrument, are expressed in μ g cm⁻³, and since the chemical composition is known, we are able to convert the mass to volume concentration, and directly compare them with the LIRIC retrievals. For the conversion we use the following equations:

$$C_{i}(z) = \frac{M_{i}(z)}{\rho_{i}} \tag{2}$$

$$C_f(z) = C_{PM_1}(z) = \sum_{i=1}^{i=5} C_i(z)$$
 (3)

where the subscript i denotes each constituent of the PM₁ fraction measured by the AMS. The denominator in Eq. 2 is the particle density (ρ_i) for each chemical compound: $\rho_{SO_4}=1.75~g~cm^{-3}$,(Lide, 2007); $\rho_{OM}=1.2~g~cm^{-3}$,(Turpin and Lim, 2001); $\rho_{NO_3}=1.75~g~cm^{-3}$, $\rho_{Cl}=1.75~g~cm^{-3}$, $\rho_{NH_4}=1.75~g~cm^{-3}$ (Lide, 2007). The total fine mode volume concentration ($C_{PM_1}(z)$) is retrieved by summing the volume concentration of each constituent (Eq. 3).

In order to cross-validate our retrievals independently with a second instrument on board, we further estimate the PM₁ volume concentration, from size distribution measurements obtained with PCASP instrumentation at six discrete height levels. The PCASP volume size distribution $(\frac{dV(r)}{dh(r)})$ is calculated from the number size distribution

PCASP data assuming spherical particles, as follows (Seinfeld and Pandis, 2006):

$$\frac{dV(r)}{dln(r)} = v(r) \times \frac{dN(r)}{dln(r)} = \frac{4}{3} \times \pi \times r^3 \times \frac{dN(r)}{dln(r)} \tag{4}$$

where v(r) the volume of particles with radius r and $\frac{dN(r)}{d\ln(r)}$ the number of particles in the infinitesimal size range $r \pm d\ln(r)$.

To evaluate the PCASP derived volume size distributions, we compare with the AERONET size distribution. In Fig. 9, the volume concentrations retrieved with PCASP at various aircraft flight heights over the GAA, along with the effective-column volume size distribution retrieved by AERONET at 13:27 UTC are shown. The error bars on the size distributions of PCASP, denote the refractive index correction uncertainty (according to Rosenberg et al., 2012). For a direct comparison of the effective-column AERONET data (in units of $\mu m^3 \, \mu m^{-2}$) with the heightresolved PCASP data (in units of µm³ cm⁻³), the AERONET volume size distribution has been divided by a layer depth of 2 ± 1 km (black solid and dashed lines in Fig. 9). The layer depth of 2 ± 1 km was chosen for demonstrating the variability of this approach. From the lidar sounding seems that aerosol particles are present up to 3 km height. However the under study free tropospheric layer is having a layer depth of 1 km, extending from 2 to 3 km a.s.l. On the other hand, the volume size distribution retrieved from AERONET is related to the entire atmospheric

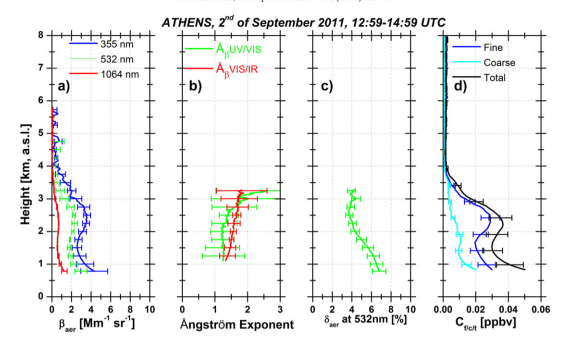


Fig. 6. From left to right vertical profiles of: (a) the aerosol backscatter coefficient at 355, 532 and 1064 nm (Mm⁻¹ sr⁻¹), (b) the backscatter related Ångström exponent, (c) the particle depolarization ratio at 532 nm (%), retrieved from backscatter and depolarization lidar measurements and (d) volume concentration (ppbv) of fine/coarse aerosol mode, retrieved from LIRIC, on 2nd of September 2011 (12:59–14:59 UTC).

column, in contrast to the size distributions retrieved from PCASP measurements at various height bins. It is observed that in the height range of 2–3 km, the volume concentration of fine mode particles is predominant over the corresponding coarse mode, as also observed in LIRIC retrievals. Moreover, a good correlation of the two distributions is found in the fine mode range of the retrievals. The cut-off radius of fine mode has been estimated by AERONET at 0.439 μ m, with almost 7% difference from the in-situ observations performed with PCASP (0.472 μ m). Additionally, the fine mode width (FWHM) of the size distribution measured by PCASP was found to vary from 0.143 μ m (at

2.92~km) to $0.168~\mu m$ (at 3.25~km). The corresponding value retrieved from AERONET columnar measurements was found to be 8.5~to 27.5% lower ($0.132~\mu m$). Regarding the modal radius of the fine mode distributions, PCASP measurements revealed the value of $0.133~\mu m$ while AERONET columnar observations showed a modal radius at $0.148~\mu m$.

Following the aforementioned methodology, we present in Fig. 10 the volume concentrations acquired with the airborne in-situ instruments ($C_{PM_1}(z)$ in ppbv; red circles AMS; black circles PCASP) along with the volume concentration, of fine aerosol mode ($C_f(z)$ in ppbv), as derived from the LIRIC algorithm (blue line). Integrating the volume

Date: 2nd of September 2011, 14:00UTC

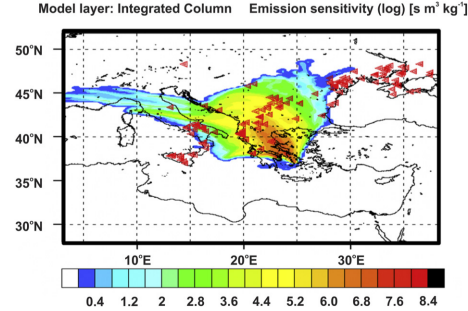


Fig. 7. Emission sensitivity (log) in s m^3 kg $^{-1}$ for 3 days FLEXPART-WRF backwards calculation starting at 2nd of September 2011–14:00 UTC. The arrival (receptor) layers are at 2–3 km above Saronic Gulf, Evoikos Gulf and Aegean Sea (GAA). The red triangles indicate the location of active fires as detected by MODIS during the simulation period. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

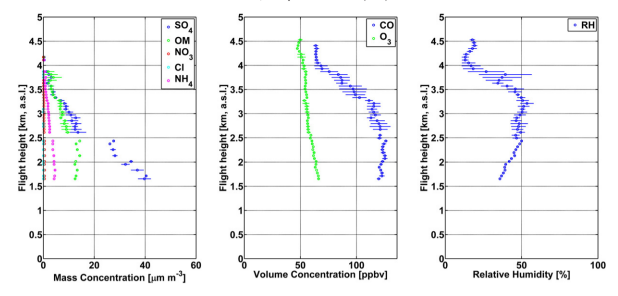


Fig. 8. Vertical profiles of (a) PM_1 mass concentration (in $\mu g/m^3$) (b) gaseous pollutant (CO, O₃) volume concentration (in ppbv) and (c) relative humidity (%), retrieved from various instruments on board, during B638 flight and for the time period 09:15–10:38 UTC.

concentration retrievals of PCASP in the size range bin 0.06–0.472 μm (in terms of radius), we estimated the volume concentration of PM₁, for each height bin. Concentrations retrieved by PCASP measurements are slightly lower than the corresponding measurements of AMS (mean relative difference ~ 30%), however within the error bars.

The airborne AMS dataset was obtained during morning hours (09:15–10:38 UTC) under low to moderate RH conditions (~ 50%), while the LIRIC retrievals obtained under higher RH conditions, influenced probably from a water uptake effect. Consequently, we need to take account this effect in order to make a straight forward comparison of LIRIC and AMS concentration retrievals. Assuming that for the entire time period (09:15–15:00 UTC), the aerosol chemical composition was constant above our measurement site but the meteorological conditions varied, we estimated the growth of the total mass concentration by employing the ISORROPIA II model.

ISORROPIA-II (Fountoukis and Nenes, 2007) calculates the particle water associated with inorganics for the PM1 aerosol fraction based on a thermodynamic equilibrium between an inorganic aerosol (NH₄-SO₄-NO₃-Cl-Na-Ca-K-Mg-water) and its gas phase precursors (Guo et al., 2015; Bougiatioti et al., 2016). Here, ISORROPIA-II was operated in the "forward mode" assuming a metastable aerosol state. The inputs to ISORROPIA-II were the inorganic ions measured by the AMS, while values for Na⁺, Ca²⁺, K⁺ and Mg²⁺ were considered to be equal to zero as their contribution to the submicron fraction is negligible, and

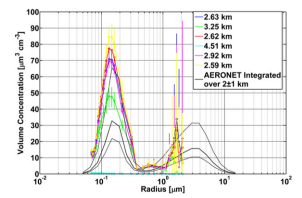


Fig. 9. Size distributions of volume concentrations obtained by PCASP at discrete height levels, and size distribution retrieved by AERONET at 13:27 UTC (solid and dashed black lines). AERONET size distributions have been divided by layer depths of $2\pm 1~{\rm km}$ (black solid and dashed lines) for direct comparison with PACASP measurements.

RH and T measured by the aircraft. Particle water concentrations were subsequently revisited by changing the respective RH and T values to match the ambient data and the model was reinitiated, comparing finally the derived mass concentration values with the initial ones. The results from simulations with ISORROPIA II, are demonstrated as inset figure in Fig. 10. The growth of the total mass remains constant (1) up to almost 60% RH, taking the values of 1.33 and 1.78 at 70% and 80% RH respectively. Since in our dataset we do not have a solid information regarding the vertical variability of RH, we consider indicative RH values of the order of 70–80%. We applied the growth factors of total mass concentration derived from ISORROPIA II to the AMS data set (Fig. 10 open turquoise and green circles). As demonstrated by the inset figure, as the RH approaches the 100% LIRIC assumption of a vertically uniform fine mode properties, could lead to large errors.

For the fine mode concentration profiles as retrieved with LIRIC algorithm and AMS measurements, shown in Fig. 10, the mean relative difference of LIRIC retrievals from AMS measurements found to be -24.2% with a mean bias of -0.003 ppbv. This mean bias of LIRIC concentration becomes -0.001 ppbv for data points obtained above 2 km height. The

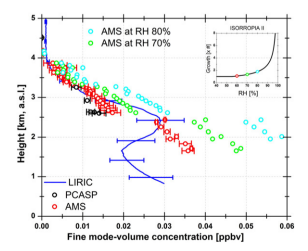


Fig. 10. Fine mode volume concentration profiles retrieved by, LIRIC (blue line), AMS (red open circles), and PCASP (black open circles). The inset figure demonstrates simulations of ISORROPIA II, regarding the total mass concentration growth for various RH values. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

RMSE value found to be 0.006 with mean fractional bias and mean fractional error of the order of -0.136 and 0.315 respectively.

A comparison between LIRIC retrievals and airborne measurements at specific height bins, where in-situ AMS data are available, is shown in Fig. 11. For this comparison we used the AMS dataset instead of the PCASP, due to the largest availability of AMS measurements during the ACEMED campaign. The total number of data points used are 29. The correlation coefficient between LIRIC $C_f(z)$ and AMS $C_{PM_1}(z)$ is 0.85, showing the very good agreement between LIRIC retrievals and in-situ measurements. However this correlation becomes stronger (0.97) when excluding the data points sampled at heights below 2 km. These height bins, were found to be the most distant from the AERONETlidar station, and belong to the lowest part of troposphere which is linked mostly to anthropogenic activity, strongly contributing to the aerosol load inside the planetary boundary layer. This load can vary significantly inside the GAA in a horizontal distance of 220 km. Overall, the $C_f(z)$ values present a very slight underestimation up to 0.001 \pm 0.001 ppbv compared to $C_{PM_1}(z)$. However this bias should be considered negligible since as can be seen from Fig. 11, is driven by the three data points sampled below 1.9 km. Beside the large spatial variability of locally produced particles up to 1.9 km in a horizontal distance of 220 km, the observed discrepancies below that height may also be attributed to LIRIC limitations. According to the LIRIC retrievals and the linear particle depolarization ratio (~ 6%) below 2 km, a mixture of fine and coarse particles is observed in the lower part of the atmosphere, while above 2 km mostly fine particles are observed (Fig. 6). The usage of constant with height AERONET values in LIRIC retrievals may lead to high uncertainties in the retrieval of the volume concentration profiles, especially in cases with non-homogeneous aerosol mixing (Granados-Muñoz et al., 2014), as observed in this case. Moreover, LIRIC volume concentration retrievals are also affected by the incomplete overlap of the lidar systems.

The case study of fine particles described here, was an excellent case for validating LIRIC with in-situ airborne instrumentation since there are no limitations regarding (i) the presence of non-spherical particles and (ii) the inlet size and pipeline loses of the in-situ instruments for particles up to 1.5 µm in radius. Due to (i) it is feasible to use Mie scattering calculations for estimating the number size distribution values and applying the refractive index correction (Rosenberg et al., 2012). Under those conditions LIRIC revealed a good performance regarding the fine mode aerosol concentration retrievals, especially for the height range above 2 km where the fine mode contribution is highly predominant, and the LIRIC code is less influenced by the overlap height. The

observed discrepancies below 2 km could be partly explained to the presence of mixed coarse and fine particles, which introduce difficulties in LIRIC to accurately distinguish between the fine and coarse particles, and to the lidar signals which are affected by the overlap height. However, our results below 1.9 km are not conclusive since there are only three available data points below that height.

4. Summary

In this study we demonstrated a good performance for the LIRIC algorithm regarding the retrieval of the volume concentration profile in the fine mode. The evaluation was done against high-quality and wellestablished in-situ airborne measurements. This validation is performed for the first time, for a case of fine mode particles, specifically for a smoke case study over Greece on 2nd of September 2011. We analysed the case study using all measurements available and characterized the aerosol load in terms of optical and microphysical/chemical properties specifically as follows: (i) we analysed in detail the increased values of aerosol optical properties observed in the atmospheric column, with AERONET (AOD 0.82 at 340 nm), and we estimated the range dependent aerosol optical properties, with backscatter-depolarization lidar measurements. (ii) The synergistic use of passive and active remote sensing measurement made also feasible the aerosol characterization in terms of volume concentration, demonstrating values of 0.03 ppbv regarding fine mode. (iii) The available airborne in-situ measurements of chemical composition, revealed the presence of sulphates (57.6%) and organic carbon (31.6%) fractions of fine mode in the atmosphere, and justified the biomass origin of the detected layer. (iv) In addition the airborne in-situ measurements have been used for validating the fine mode volume concentration retrievals of LIRIC during the specific case study. The correlation coefficient between AMS and LIRIC values was found to be 0.85 and reach even the 0.97, if we exclude the data points at heights below ~2 km, showing a satisfactory agreement between LIRIC retrievals and in-situ measurements. The $C_f(z)$ values are slightly underestimated compared to $C_{PM_1}(z)$ up to 0.003 ppbv, with a mean relative difference of -24.2% and a RMSE of 0.006. However, for the height range below 2 km, higher discrepancies are observed which may be attributed: (i) to the time difference, of around two to three hours, between LIRIC retrievals and the AMS measurements, and (ii) the aerosol spatial variability over the GAA, especially for heights below 2 km which are strongly affected by the anthropogenic activity (iii) to the lidar incomplete overlap region (iv) to LIRIC limitations due to non-homogeneous aerosol mixing.

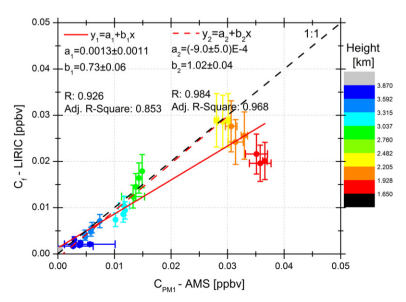


Fig. 11. AMS versus LIRIC fine mode volume concentrations retrieved at various heights (indicated with colour scale).

Acknowledgment

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