Atmos. Chem. Phys. Discuss., 15, 32723–32757, 2015 www.atmos-chem-phys-discuss.net/15/32723/2015/ doi:10.5194/acpd-15-32723-2015 © Author(s) 2015. CC Attribution 3.0 License.



This discussion paper is/has been under review for the journal Atmospheric Chemistry and Physics (ACP). Please refer to the corresponding final paper in ACP if available.

Temporal consistency of lidar observables during aerosol transport events in the framework of the ChArMEx/ADRIMED campaign at Menorca Island in June 2013

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Received: 9 October 2015 - Accepted: 9 November 2015 - Published: 19 November 2015

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



Abstract

We performed synergetic daytime and night-time active and passive remote sensing observations at Menorca (Balearic Island, Spain), over more than 3 weeks during the Chemistry-Aerosol Mediterranean Experiment/Aerosol Direct Radiative Effect in the Mediterranean (ChArMEx/ADRIMED) special observation period (SOP 1a, June-July 2013). We characterized the aerosol optical properties and type in the low and middle troposphere using an automated procedure combining Rayleigh-Mie-Raman lidar (355, 387 and 407 nm) with depolarization (355 nm, AERONET Cimel[®] sunphotometer data. Results show a high variability due varying dynamical forcing. The mean column-averaged lidar backscatter-to-extinction ratio (BER) was close to 10 0.024 sr^{-1} (lidar ratio of ~ 41.7 sr), with a large dispersion of ±33 % over the whole observation period due to changing atmospheric transport regimes and aerosol sources. The ground-based remote sensing measurements, coupled with satellite observations, allowed to document (i) dust particles up to 5 km a.s.l. in altitude originating from Morocco and Algeria from 15 to 18 June with a peak in aerosol optical thickness 15

- (AOT) of 0.25 ± 0.05 at 355 nm, (ii) a long-range transport of biomass burning aerosol (AOT = 0.18 ± 0.16) related to North American forest fires detected from 26 to 28 June 2013 by the lidar between 2 and 7 km and (iii) mixture of local sources including marine aerosol particles and pollution from Spain. During the biomass burning event, the high
- value of the particle depolarization ratio (8–14%) may imply the presence of dust-like particles mixed with the biomass burning aerosols in the mid troposphere. We show also linearity with SEVIRI retrievals of the aerosol optical thickness within 35% relative bias, which is discussed as a function of aerosol type.



1 Introduction

The Mediterranean has been identified as one of the "hot-spots" in projections of future climate change (Giorgi and Lionello, 2008) and it has been recently shown that aerosol direct and semi-direct effects, which were not properly taken into account in global climate change simulations (IPCC, 2014), have a significant impact on surface temperature, evaporation, and precipitation at the regional scale (Nabat et al., 2015a), i.e. a likely positive feedback on the trend for future dryer and thus more turbid Mediterranean summers. Due to the variability of aerosol properties over the Mediterranean basin, this calls for a more representative description of aerosol optical properties and spatiotemporal distribution by both observations and models.

Regional experiments including measurements of the vertical distribution of aerosols were performed some time ago to characterize aerosols over the Mediterranean Sea (i) in the framework of the MEditerranean DUSt Experiment (MEDUSE) in 1997 (Hamonou et al., 1999), (ii) the Scientific Training and Access to Aircraft for Atmo-

- spheric Research Throughout Europe (STAAARTE) airborne flights in 1997 (Dulac and Chazette, 2003) and 1998 (Formenti et al., 2002), (iii) with a lidar deployed in Crete (Gobbi et al., 2000) or an instrumented ultralight aircraft in Lampedusa (Di Iorio et al., 2003) during the Photochemical Activity and Ultraviolet Radiation (PAUR II) campaign in 1999, (iv) over the eastern Mediterranean basin during the Mediterranean Inten-
- sive Oxidant Study (MINOS; Lelieveld et al., 2002;) and Mediterranean Israeli Dust Experiment (MEIDEX; Levin et al., 2005) in 2001; and (v) over the urban and industrial region of Marseille–Fos–Berre on the French Mediterranean coast also in 2001 (Cros et al., 2004; Cachier et al., 2005), (vi) in the framework of the EARLINET network (Papayannis et al., 2008). Such past experiments have produced very useful informa-
- tion about the vertical distribution of Mediterranean aerosol optical properties, based on in-situ observations and lidar measurements. During those preceding campaigns in the Mediterranean region, the use of aerosol lidars was focused on rather short time periods, but they appear as a very powerful tool to identify the wide spectrum of



aerosol types encountered in the tropospheric column (e.g. Chazette, 2003; Chazette et al., 2005a; Berthier et al., 2006; Groß et al., 2011; Tesche et al., 2011; Nisantzi et al., 2014). The multidisciplinary programme Mediterranean Integrated Studies at the Regional and Local Scales (MISTRALS; http://www.mistrals-home.org), initiated by

⁵ CNRS/INSU in 2010 to study the future habitability of the Mediterranean region, offered the opportunity, within the Chemistry–Aerosol Mediterranean Experiment (ChArMEx, http://charmex.lsce.ipsl.fr), to conduct ground-based and airborne lidar observations at the scale of the Western Mediterranean basin.

The ChArMEx/Aerosol Direct Radiative Effect in the Mediterranean (ADRIMED) special observation period (SOP-1a) was set-up from 11 June to 3 July to study aerosol optical properties and radiative effects in the western Mediterranean during the dry season, which shows a maximum in aerosol optical depth (Nabat et al., 2013). The campaign involved several surface stations throughout the western Mediterranean, research aircrafts, and instrumented balloons (Mallet et al., 2015). In this work, we fo-

- ¹⁵ cus on both active and passive remote sensing observations performed at Menorca (Balearic Island, Spain) during this campaign. The main goal of the paper is to demonstrate the benefit of continuous daytime and night-time lidar measurements during at least 3 weeks to derive aerosol optical properties. It improves both the assessment of the diurnal variation of the aerosol distribution related to the planetary boundary layer
- 20 (PBL) growth and the probability to detect long range transports of aerosol plumes. In Sect. 2, we first present the experimental set-up. Retrieved aerosol optical properties for both night-time and daytime conditions are analysed in Sect. 3 to give an overall identification of aerosol types. In Sect. 4, we discuss the different origins of aerosol particles before discussing comparison with the Spinning Enhanced Visible and InfraRed Imagen (OEV(ID)) attributes and completing in Sect. 5.
- ²⁵ Imager (SEVIRI) retrievals and concluding in Sect. 5.



2 Ground-based remote sensing measurements

During the campaign, our custom-made Raman lidar WALI (Chazette et al., 2014) was operated together with an AERONET sunphotometer at Cap d'en Font (http://aeronet.gsfc.nasa.gov/new_web/photo_db/Cap_d_En_Font.html) on the south-eastern coast

- of the Balearic island of Menorca, Spain. The instruments were located within ~ 6 m from each other, at 39°49′32.9″ N, 04°12′29.3″ E, at ~ 10 m above the mean sea level (a.m.s.l.) and less than 70 m from a small cliff on the sea shore. The choice to use only remote sensing instruments is driven by the lack of representativeness of the ground-based in situ measurements, which are mainly affected by local dynamical forcings.
- ¹⁰ This is especially true in coastal regions (Chazette, 2003). The selected location is mainly affected by Saharan and Spanish air masses. Figure 1 shows the location of the station approximately in the centre of the western Mediterranean basin. The campaign average aerosol optical thickness (AOT₅₅₀, at 550 nm) distribution derived from SEVIRI on-board the geostationary Meteosat Second Generation (MSG) platform is reported
- ¹⁵ in this figure. It shows a classical North–South decreasing gradient in the western Mediterranean Basin due to African dust with maximum values between 0.20 and 0.25 in the Alboran \bigcirc , and minimum values of ~ 0.12 in the Gulf of Lyon. Intermediate values of ~ 0.17 are found around Menorca.

2.1 Raman lidar

- ²⁰ The WALI instrument uses an emitted wavelength of 354.7 nm and is designed to fulfil eye-safety conditions. The instrument, its calibration and the associated errors are documented in Chazette et al. (2014) and will not be detailed here. During all the experiment, the acquisition was performed continuously with a vertical resolution of 15 m for mean profiles of 1000 laser shots leading to a temporal sampling close to
- 1 min. The UV pulse energy is ~ 60 mJ and the pulse repetition frequency is 20 Hz. It is equipped with four detection channels: an aerosol optical board including co-polarized and cross polarized channels with respect to the laser emission, and a Raman opti-



cal board including a channel dedicated to the detection of the water vapour Raman signal at 407.5 nm and a fourth channel dedicated to the recording of the atmospheric di-nitrogen Raman signal at 386.6 nm. We briefly recall the systematic approach used to invert the lidar profiles in the following.

- ⁵ The presence of clouds was visually detected in the lidar time series of rangecorrected lidar backscattered profile and the corresponding periods were removed. Depending on the parasitic sky radiance contribution to the measured signal, two measurement synergy types have been used to retrieve the aerosol optical properties from the lidar. They have both been validated during previous studies (e.g. Dieudonné et al. 2015). The N Baman measurements from our system are impressive above.
- et al., 2015). The N₂-Raman measurements from our system are imprecise above 2 kma.m.s.l. during daytime and cannot be used to characterize the aerosol optical properties in the upper troposphere. Hence, during daytime the sunphotometer AOT₃₅₅ is considered as a constraint for the lidar inversion as in Chazette (2003). Note that using the total AOT only allows us to retrieve a column-averaged or equivalent
- ¹⁵ backscatter-to-extinction ratio (BER, product of the backscatter phase function and the single scattering albedo, inverse of the lidar ratio LR), integrating all the aerosol layers. During night-time (low sky radiance), the two elastic and the N₂-Raman channels of the lidar are used to determine simultaneously the aerosol BER, the vertical profile of the aerosol extinction coefficient (α_e), and the particulate depolarization ratio (PDR).
- ²⁰ All methodological details are well presented in Royer et al. (2011a). The approaches to retrieved the PDR and the water vapour mixing ratio (WVMR) are described in detail in Chazette et al. (2012b, 2014), respectively.

Two representative examples of AOT and BER retrieval are given in Fig. 2 corresponding to the main aerosol sources, biomass burning and desert dust observed ²⁵ during this campaign. They demonstrate the good agreement between the cumulative AOT derived from the N₂-Raman and the elastic channels. The calculations have been performed using the average profile of night-time measurements during the nights of 16–17 and 27–28 June, for biomass and dust cases, respectively. To improve the inversion, the mean profiles have been inverted using an altitude-variable BER and a regu-



larization approach (Royer et al., 2011). For the first example, the BER (LR) is close to $0.04 \, \text{sr}^{-1}$ (25 sr) in the marine boundary layer (MBL) and decreases with the altitude to reach values between 0.02 and $0.025 \, \text{sr}^{-1}$ (50 and 40 sr) between 2 and 3 km a.m.s.l. The values of BER are similar for the second example in the MBL, but after decreasing below $0.02 \, \text{sr}^{-1}$ in the aerosol layer above the MBL, they significantly increase above $4 \, \text{km a.m.s.l.}$ to reach ~ $0.025 \, \text{sr}^{-1}$. These two profiles correspond to the main contributions of aerosol sources encountered during this period: maritime aerosol in the MBL (BER ~ $0.04 \, \text{sr}^{-1}$), dust (BER ~ $0.025 \, \text{sr}^{-1}$) and biomass burning or local pollution (BER < $0.02 \, \text{sr}^{-1}$).

- ¹⁰ Uncertainties on the retrieval of aerosol optical properties from similar detection channels have already been extensively discussed in Chazette et al. (2010) and Royer et al. (2011). In general, a vertical profile of BER can be derived during night-time, after averaging the lidar measurements for at least 1 h to reach a sufficient signal-to-noise ratio (SNR > 10), in order to extract the information for the highest aerosol layers (above 3 km amsl). The relative uncertainty on the BER is ~ 5% (~ 10%) during night-time
- ¹⁵ 3 km amsl). The relative uncertainty on the BER is ~ 5% (~ 10%) during night-time (daytime). The relative uncertainties on the PDR are close to 10% for the encountered AOT at 355 nm (AOT₃₅₅ > 0.2). The relative uncertainty on the AOT is less than 2%. The relative uncertainty on the water vapour mixing ratio (WVMR) is between 7 and 11% within the first kilometres of the atmosphere (Chazette et al., 2014).

20 2.2 Sunphotometer

The Cimel[®] sunphotometer is part of the Aerosol Robotic Network (AERONET; http: //aeronet.gsfc.nasa.gov/cgi-bin/type_piece_of_map_opera_v2_new; Holben et al., 1998). It performs measurements of solar light extinction at 8 wavelengths in the solar spectrum between 340 and 1020 nm to retrieve the AOT at 7 wavelengths. The instru²⁵ ment field of view is about 1° and the channel bandwidths are less than 20 nm. The instrument was calibrated prior to and after the campaign by the observation service Photométrie pour le Traitement Opérationnel de Normalisation Satellitaire (PHOTONS; http://loaphotons.univ-lille1.fr/), the French component of AERONET. We have used



Level-2 quality assured data. The AOT is retrieved with a maximal absolute uncertainty of 0.02, independent of the aerosol load. The aerosol optical thickness at the lidar wavelength of 355 nm (AOT₃₅₅) has been assessed using the Ångström exponent (Ångström, 1964) and the sunphotometer AOT at 380 and 440 nm. Sunphotometer

⁵ AOT values at 500 and 675 nm are also used in this work for a better comparison to satellite products described below. Additionally, these saturements were checked against and completed by a SOLAR Light[®] Microtops II manual sunphotometer, calibrated by PHOTONS shortly before the campaign (AERONET instrument #695).

3 Temporal continuity of the aerosol optical properties

10 3.1 Vertically integrated aerosol optical properties derived from the sunphotometer

Both times series of AOT at 500 nm (AOT₅₀₀) and Ångström exponent between 440 and 675 nm, as directly measured by the sunphotometer, are plotted in Fig. 3.

- AOT and Ångström exponent both exhibit a strong variability due to the succession
 of aerosol events of different types, as revealed by the large range of variation of the Ångström exponent between ~ 0.4 and 2.15. The fine mode fraction of AOT also reported in the figure clearly co-varies with the Ångström exponent. The coarse mode contribution is dominant from 16 to 20 June (coarse mode fraction of AOT between 50 and 80%) and also important on 24–25 June (35 to 70%). The AOT appears to be
 higher with values larger than 0.2 during such periods. The AOT, which is below 0.38 (on 18 June) during the first 2 weeks of campaign, significantly increases on 26–28 June, showing several maxima (up to more than 0.6 on 27 June). Variations with particularly large amplitude appear on the 26 and 27 June. Usually such peaks are due to North-African dust aerosol transport over the Western Mediterranean basin (e.g. Moulin et al., 1998; Hamonou et al., 1999). In our case, the Ångström exponent rang-
- ing between 1 and 1.6 appears too high to support the hypothesis a dominant presence



of dust particles. Values of the Angström exponent over 1.5 are typical of pollution-like or biomass burning aerosols (Chazette et al., 2005a) and an average value of 1.80 has been observed for non-dust conditions over the Mediterranean by Paronis et al. (1998). Computations by Hamonou et al. (1999) suggest that a dust contribution to this AOT cannot be excluded but should be under a 0.45 fraction for the observed range of

- Angström exponent. The lowest AOT values observed on 10 June together with a low range of Ångström exponent (0.01–0.21) are typical of a clear marine atmosphere with an aerosol population dominated by sea-salt particles. We notice that the uncertainty on the Angström exponent grows as the AOT decreases. For AOT < 0.15, the meaning
- of the Ångström exponent is subject to caution.

3.2 Aerosol optical properties derived from the ground-based lidar WALI

The temporal evolution of the BER derived from both the lidar measurements and the sunphotometer are compared in Fig. 4. The sunphotometer-derived column-integrated BER of the aerosols can be computed from the single scattering albedo and the phase function at a scattering angle of 180° and 440 nm wavelength derived from the opera-15 tional algorithm of AERONET (Dubovik et al., 2000). The root mean square error (rmse in gray area) on the lidar-derived BER, determined as the variability over 20 min, is close to 0.004 sr⁻¹ on average, which is comparable with the one retrieved by Chazette et al. (2012a) with a similar lidar system set-up in Menorca in the 2012 autumn season. We note a good coherence with the BER at 440 nm derived by the AERONET 20 sunphotometer. Yet, the sunphotometer-derived BER seems to be underestimated by ~ 0.004 to 0.01 sr⁻¹ comparatively with the lidar between 19 and 26 June; the larger variability bars at this period are due to a lighter aerosol load (see Fig. 3), which may explain part of this discrepancy. The higher values of BER seen by the lidar would also be consistent with the hygroscopic properties of aerosols within the PBL where the rel-

25 ative humidity significantly increases (Fig. 4b) to reach more than 90%. Indeed, BER may increase with the growth of aerosols. Moreover, the surface wind speed rose on



26 June, with gusts reaching 12 m s^{-1} , which may seed the atmosphere with marine aerosols (Blanchard et al., 1984).

For the sake of checking the consistency between the inversion procedures used during night-time and daytime, the histograms of the equivalent BER are compared in ⁵ Fig. 5 for daytime, night-time, and the whole day. These histograms account for all lidar data inverted in cloud-free conditions. The values greater than 0.045 sr^{-1} are not significant and may represent situations where the inversion process does not converge. Hence, ~ 10% of lidar profiles have not been considered in the synthesis. The nighttime BER distribution, with a BER value of $0.024 \pm 0.008 \text{ sr}^{-1}$, is only slightly smaller than the daytime distribution ($0.026 \pm 0.007 \text{ sr}^{-1}$). Hence, the BER values are deemed consistent between daytime and night-time, and the synthesis on the entire experiment

period shows an average of $0.024 \pm 0.008 \text{ sr}^{-1}$. The temporal evolutions of the vertical profile of aerosol extinction coefficient and PDR are shown in Fig. 6. PDR is an effective parameter to separate the contribution

- of the more spherical particles from the ones due to dust-like aerosols (e.g. Chazette et al., 2012a). Between 16 and 19 June the PDR value is between 10 and 27 %, which is representative of non-spherical dust-like aerosols (Müller et al., 2007; Tesche et al., 2011) as identified in Fig. 4a for BER between ~ 0.021 and 0.028 sr⁻¹. Except between 26 and 28 June, the aerosol content is dominated by spherical particles. Between
- 20 26 and 28 June a depolarizing layer is observed between ~ 5 and 7 km a.m.s.l. The PDR ranges between 8 and 14 % suggesting that dust aerosols were mixed with other aerosol sources or were processed during their transport to Menorca. We will further discuss this case in Sect. 4. When considering the temporal evolution of AOT also given in Fig. 6a, we note that lidar- and sunphotometer-derived AOT significantly differ
- on several occasions, especially in the cloudy periods (11, 20, 24, 25 and 27 June). It is due to residual cloud layers in the lidar profiles, which are not seen on the line-of-sight of the sunphotometer, with a positive bias explained by the higher BER of these thin layers. In addition, from 26 to 28 June, the presence of high altitude aerosol layers also probably causes a strong heterogeneity of the aerosol BER in the tropospheric



column (see Fig. 2b), which may explain part of the previous discrepancies because the aerosol types may be very different against the altitude. This shows the limited relevance of the notion of column-equivalent BER in heterogeneous cases.

3.3 Evidence of contributions by aerosol type as discriminated by lidar

- ⁵ The temporal evolution of the observed aerosol species can be derived from the analysis of the equivalent BER and PDR. Indeed, these two parameters, only calculated from the lidar profiles, are sufficiently discriminating to the profiles of the main aerosol types in most cases (Burton et al., 2012). Three aerosol families are considered: (i) dust-like aerosols with values of BER and PDR red on 0.022 sr⁻¹ and 20%, respectively,
- (ii) pollution aerosols with BER and PDF centred on 0.015 sr⁻¹ and 2%, respectively, and (iii) marine aerosols with mean values of BER and PDR centred on 0.04 sr⁻¹ and 0%, respectively. Figure 7 gives the temporal evolution of the aerosol type after defining a specific colour map as a function of BER and PDR. The lidar profiles were here averaged during 1 h with a vertical resolution of ~ 30 m. The aerosol backscatter coef-
- ficient (ABC) is coded by colour density: the more saturated, the larger the ABC (white corresponds to ABC = 0). A specific colour scale is affected to the couple of variables BER and PDR during night-time. As the inversion using the N₂-Raman channel is not possible when the sun is up, the colour map has been only associated with the PDR during daytime.
- ²⁰ Such a graphic representation allows the refinement of the identification of the aerosol types that are presented in Fig. 4a. The higher BERs retrieved in Fig. 4a between 19 and 26 June are due to a larger contribution of reflective aerosols in the lower layers, likely sea salt particles, which may also be very hydrophilic. Pollution aerosols are present all along the measurement period except during the dust event between 16
- and 19 June. It is more difficult to attribute the layer above 5 km a.m.s.l. during the night of 26–27 June to a single aerosol source because depolarization is observed simultaneously with low BER. This layer arrives above a layer of biomass burning or polluted aerosols which spread between ~ 3 and 5 km a.m.s.l. A succession of pollution plumes



originated from different locations along the Spanish coast contributes to the aerosol pollution load in the lower free troposphere over Menorca according to the air mass trajectories (not shown). However for the 26–27 June period the long range transport revealed by the backtrajectories discussed by the backtr

American biomass burning aerosol sources. The intermittent plumes, lifted as the PBL develops over Spain each afternoon, explain the periodic behavior observed in Fig. 4a for the temporal evolution of the column-equivalent BER. This temporal evolution of aerosol types based on the unique analysis of the lidar data is quite consistent with the column-integrated observations of the AERONET sunphotometer, as discussed in Sect. 3.1 and reported in Fig. 3.

3.4 Regional representativeness as seen by spaceborne measurements

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The observations conducted from the Menorca Island station are relevant to the local atmospheric column. In the following we put them in a more regional context using the measurements performed by SEVIRI (e.g. Bennouna et al., 2009) and the spaceborne instruments Moderate Resolution Imaging Spectroradiometers (MODIS; Salmonson et al., 1989; King et al., 1992; http://modis.gsfc.nasa.gov). The MODIS data above land (few data are available above sea due to sun-glitter) and the SEVIRI data above sea are combined in a single map to check the reliability in terms of continuity between sea and continent.

- The spatial resolution of the MYD04_L2 product of MODIS is $10 \text{ km} \times 10 \text{ km}$ at nadir. The predicted uncertainty on the AOT at 550 nm over land is $\pm 0.15 \cdot \text{AOT} \pm 0.05$. The spatial and temporal resolutions of SEVIRI measurements are $10 \text{ km} \times 10 \text{ km}$ and 15 min, respectively. The uncertainty on the SEVIRI-derived AOT is very dependent on the aerosol type (Bennouna et al., 2009). Compared to AERONET products from
- ²⁵ coastal stations, Thieuleux et al. (2005) do not highlight any significant bias on the AOT at 550 nm derived from SEVIRI for values between ~ 0.07 and 1. Their comparison based on observations in 2003 indicates that the SEVIRI AOT product is of somewhat lower quality at the sunphotometer sites directly affected by a desert dust plume from



northern Africa. This is attributed to the fact that the aerosol models used to compute the look-up table does not include a specific desert dust model. Bréon et al. (2011) report a bias of 0.07 from their more exhaustive evaluation with AERONET sunphotometers over the period from June 2005 to December 2010. From a similar linear fitting between MODIS and AERONET, they found a smaller bias of ~ 0.02 and a correlation slope close to 1.

Figure 8 shows the inter-comparison between qua -hourly products from SEVIRI and from the coincident AERONET sunphotometer of Menorca, including the AOT at 550 nm (Fig. 8a) and the Ångström exponent (Fig. 8b, computed between 630 and

- 810 nm, and 675 and 870 nm for SEVIRI and the sunphotometer, respectively). For the AOT, a linear least square fit highlights a significant deviation from the 1 : 1 relationship with a factor of 0.65. The additive bias is low, positive and close to 0.03. The mean rmse is ~ 0.066. The main discrepancies are mostly observed for the highest AOTs, occurring between 17–19 and 26–28 June when marine and dust aerosol are mixed
- and when biomass burning aerosols arrived above the site. The latter case is likely also associated with aerosol mixing. The discrepancies can be due to the resulting difficulty of the inversion process to identify a proper aerosol model, even for dust particles which never completely prevail in terms of AOT. We note the larger dispersions for the mixing of marine particles with dust or pollution aerosols. In the following, we have corrected
- ²⁰ by -35 % the SEVIRI AOT₅₅₀ product. Whereas the sunphotometer-derived Ångström exponent seems coherent with our previous classification, our results suggest that the SEVIRI Ångström exponent product (Fig. 8b) relatively large discrepancies at all AOT (although less at larger values), mostly related to aerosol type, and this microphysical properties. The dispersion is lower for the dust (red in the figure) and biomass burning
- (brown in the figure) events, but with overestimation and underestimation, respectively. Consequently, the SEVIRI-derived AOT₅₅₀ product over ocean, that relies on the evaluated Ångström exponent, should be carefully checked before use.

The situations with the strongest AOT contrasts above the western Mediterranean basin are shown in Fig. 9. We can notice the very good continuity, after the correction



of the SEVERI-derived AOT, between sea and continent (MODIS-derived AOT). The main aerosol events are linked with either the highest PDR observed between 16 and 19 June, or the highest altitude transport (above 5 km a.m.s.l.) between 26 and 28 June. The first event is due to desert dust aerosols off the Moroccan and Algerian coasts (see also Fig. 4). The second event reveals a plume crossing the Mediterranean from North to South and will be discussed hereafter. It is associated with a decrease of the BER after 26 June as shown in Fig. 4a after 26 June.

4 Discussion

The pollution transport events observed at Menorca in the first part of the campaign,
12–18 June (Fig. 4), are associated with the lowest values of the BER. To investigate their origins, we ran the HYbrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Draxler and Rolph, 2014) with 3 hourly archived meteorological data provided by the US National Center for Environmental Prediction (NCEP) Global Data Assimilation System (GDAS) at the horizontal resolution of 0.5°. Two-day back-trajectories (not shown) clearly trace those polluted air masses back to Spain. Still in the same period, we note an increase of the BER during night-time. It may be due to a higher relative contribution of hygroscopic aerosols below 1 kma.m.s.l. as explained in Sect. 3. Between 18 and 26 June (Fig. 4), the BER reaches ~ 0.04 sr⁻¹ (LR = 25 sr) as observed by Flamant et al. (2000) for marine aerosols over the open ocean. Nevertheless, we also note weak-medium surface wind speeds between 2 and 8 m s⁻¹ not

²⁰ ertheless, we also note weak-medium surface wind speeds between 2 and 8 m s⁻¹ no favourable to a strong contribution of sea salt particles in the lower troposphere.

Satellite data show the arrival of an African dust plume from the Alboran Sea over the Balearic Islands starting slowly on 15 June and leaving Menorca on 19 June. The AOT slightly increases from 16 to 18 June, when the densest part of the dust plume passes over the Menorca Island, to reach $AOT_{355} = 0.25 \pm 0.05$. As highlighted by 3 day back-trajectories (not shown), the dust plume came from Morocco and Algeria, as also illustrated by the satellite image in Fig. 9a. In the dust layer above 1 kma.m.s.l., the



PDR is 20±5% and the mean BER (LR) is 0.024±0.002 sr⁻¹ (~41.7±4 sr). Note that, as shown Fig. 2a and previously discussed, the BER significantly evolves within the low and meddium troposphere from 0.04 sr⁻¹ (LR = 25 sr) in the MBL to ~0.020–0.025 sr⁻¹ (LR ~ 50–40 sr) in the dust layer situated above ~ 2 km a.m.s.l. A relative
⁵ peak is observed in the BER profile between 1 and 2 km a.m.s.l. associated with smaller values of the aerosol extinction coefficient (Fig. 6). This intermediate layer is associated with PDR < 2% and may be mainly affected by both marine and pollution aerosols.

The high-altitude aerosol event observed between 4 and 7 km a.m.s.l. above the western Mediterranean basin from 26 to 28 June is not usual in its nature since it results

- from a very turbid plume (AOT₅₅₀ > 0.6) arriving from the NE Atlantic as visible on the Bay of Biscay on 26 June in Fig. 10. Formenti et al. (2002) have already documented with airborne measurements such an occurrence of 10 day aged haze layers from Canadian fires over the eastern Mediterranean in August 1998. Seven days-back trajectories have been computed with the HYSPLIT model (Fig. 10). The back trajectories
- ¹⁵ are superimposed on a MODIS AOT image combining data from 24 June 2013. Dense aerosol plumes appear all along a transport pathway over the Northern Atlantic, with a dark red colour associated with AOT₅₅₀ values larger than 1. Note that the AOT even reaches 5 for many pixels, maybe because cloud contribution is also included as can be seen in the true colour image available on https://earthdata.nasa.gov/labs/worldview/.
- As shown in Fig. 10, the biomass burning plume observed at 6 km pver Menorca on 27 June crossed the Atlantic Ocean at altitudes between 4 and 8 km.s.l. The plume is associated with forest burning occurring in Canada and Colorado.

The possible source regions are discussed in detail in the companion paper of Ancellet et al. (2015). It includes contributions from two different plumes: biomass burn-

²⁵ ing aerosol from North America and dust transported westward over the Atlantic by the trade winds. Trajectories from the other plume detected close to 4 km a.m.s.l. over Menorca by the lidar on 28 June (Fig. 6) also come from North America with little contribution from the Atlantic Saharan dust. The North American aerosol event of 26– 27 June is characterized by $AOT_{355} = 0.18 \pm 0.16$ above Menorca. It represents ~ 50 %



of the total columnar AOT (AOT₃₅₅ = 0.41 ± 0.12) encountered during this day. It is associated with a moister air mass, with a WVMR close to $1-2 \text{ gkg}^{-1}$, comparatively to the clean free mid-troposphere (0.5 gkg^{-1}), as derived from the H₂O-Raman lidar channel. The uplifting of air masses from the lower troposphere occurs either above

- ⁵ the continental US or above the Atlantic ocean as discussed in Ancellet et al. (2015). The equivalent BER has been assessed to be $0.023 \pm 0.002 \,\mathrm{sr}^{-1}$ (LR ~ 43.5±4 sr) and corresponds about to the mean value of the value retrieved in the MBL correspond to marine aerosols. The BER in the mixture of biomass burning aerosol and dust is determined as $0.025 \pm 0.002 \,\mathrm{sr}^{-1}$
- (LR ~ 40 ± 3 sr) with a PDR between 8 and 14 %. PDR values between 8 and 18 % were measured over Cyprus by the lidar of Limassol (Nisantzi et al., 2014) for almost fresh biomass burning aerosols mixed with dust-like particles uplifted by thermal convection and transported above the lidar site. In our case, the aerosols are more aged (at least 7 days) and may be more spherical due to water vapour condensation dur-
- ¹⁵ ing transport over the Atlantic Ocean. We note cloud formation along some filaments created from the initial plume (not shown). Such a phenomenon decreases the PDR. Nevertheless, the value of PDR is higher and may indicate the presence of dust-like particles within the biomass burning plume. The high vertical resolution of the CALIOP lidar (30–60 m) can be processed to derive optical properties of the aerosol layers (e.g.
- ²⁰ Vaughan et al., 2004; Thomason et al., 2007; Kim et al., 2008; Berthier et al., 2006) as the PDR. The 8–14 % PDR measured at 355 nm by the WALI lidar appears comparable to the 10 % PDR at 532 nm observed by CALIOP (the ground track is given in Fig. 9d) off the Mediterranean Spanish coast at 1° E in a layer between 38 and 39° N on 28 June, 02:00 UTC (see Ancellet et al., 2015).

25 5 Conclusions

Aerosol optical properties in the tropospheric column were derived from the measurements performed continuously, during three weeks in June and early July 2013, at



Menorca Island during the Chemistry–Aerosol Mediterranean Experiment/Aerosol Direct Radiative Effect in the Mediterranean (ChArMEx/ADRIMED) special observation period (SOP-1a). The measurements sampled air masses with very different aerosol content and a large range of optical thicknesses (AOT₃₅₅ = 0.29 ± 0.17), which has been shown as representative of the years 2011 to 2013 (AOT₃₅₅ = 0.24 ± 0.15). There are only a few cases where the aerosol layers are not composed of a mixture of different aerosol types. They are originating from the surrounding sea, the Spanish coastal cities, the North Africa deserts and even distant forest fires in North America. We have

noted that the complex mixing of aerosols likely impact the retrieval of the AOT from
 SEVIRI leading to a relative bias close to 35 %.

The instrumental synergy, coupling either the sunphotometer or the N_2 -Raman channel with the elastic channel, allows a well-constrained processing of the lidar measurements, from which we were able to follow the evolution of the aerosol optical properties between night and day. In particular, the continuity of column-equivalent BER measure-

- ¹⁵ ments is ensured. Lidar observations allowed locating scattering layers in the troposphere, and in particular identifying a complex aerosol transport from North America in the middle troposphere (between 2 and 7 km a.m.s.l.). Air masses took between 5 and 7 days to arrive over the Mediterranean Sea. There has been a great variability in the nature of aerosols in the troposphere during this period from 26 to 28 June 2013. This
- ²⁰ variability is evidenced by the BER profile estimated from the Raman lidar WALI, with a strong variance (BER (LR) = $0.024 \pm 0.008 \text{ sr}^{-1}$ (~ $41.7 \pm 14 \text{ sr}$), above 4 km a.m.s.l. and < 0.02 sr^{-1} (> 50 sr) between 2 and 4 km a.m.s.l.). However, such variability has weak impact on the AOT measurements, be it during night-time or daytime. We have also observed the presence of depolarizing particles (PDR ~ 8–14% @355 nm) in
- ²⁵ a biomass burning plume originating from North America corresponding to Saharan dust re-circulated over the Atlantic Ocean, as discussed in the companion paper by Ancellet et al. (2015).

These results show that an assessment of the radiative budget of aerosols over the western Mediterranean basin can be easily performed by considering the average op-



tical properties of the particles. Nevertheless, for the evaluation of atmospheric heating rates and possible associated effects on cloud formation, account must be taken of the single scattering albedo, which is linked to the vertical evolution of the aerosol types given by our classification. Moreover, the single scattering albedo may be constrained
⁵ by the lidar-derived BER as in Randriamiarisoa et al. (2004) or Raut and Chazette (2008). The latter has indeed been shown in this campaign to be very variable, both in time and altitude, due to the mixing of very different aerosol contributions over the Mediterranean Sea.

Acknowledgements. This work was supported by the French space agency (CNES) and the
 Commissariat à l'Energie Atomique (CEA). The campaign at Menorca was also supported by CNRS/INSU and Météo-France through the MISTRALS/ChArMEx programme, and by the Spanish Ministry of Science and Innovation and FEDER funds under the projects TEC2012-34575, UNPC10-4E-442 and CGL2011-13580-E/CLI. Matthieu Jeannot from LPC2E, François Dulac and Sahar Hassanzadeh from LSCE are acknowledged for their technical help during
 the campaign. Juan Ramon Moreta Gonzalez and Miguel Angel Heredia Jodar from AEMET

are acknowledged for AERONET data from Palma.

References

20

Ancellet, G., Pelon, J., Totems, J., Chazette, P., Bazureau, A., Sicard, M., Di Iorio, T., Dulac, F., and Mallet, M.: Long range transport and mixing of aerosol sources during the 2013 North American biomass burning episode: analysis of multiple lidar observations in the Western

Mediterranean basin, Atmos. Chem. Phys. Discuss., 15, 32323–32365, doi:10.5194/acpd-15-32323-2015, 2015.

Ångström, A.: The parameters of atmospheric turbidity, Tellus, 16, 64–75, 1964.

- Bennouna, Y. S., de Leeuw, G., Piazzola, J., and Kurmierczyk-Michulec, J.: Aerosol remote sensing over the ocean using MSG-SEVIRI visible images, J. Geophys. Res., 114, D23203, doi:10.1029/2008JD011615. 2009.
 - Berthier, S., Chazette, P., Couvert, P., Pelon, J., Dulac, F., Thieuleux, F., Moulin, C., and Pain, T.: Desert dust aerosol columnar properties over ocean and continental Africa from



Lidar in-Space Technology Experiment (LITE) and Meteosat synergy, J. Geophys. Res., 111, D21202, doi:10.1029/2005JD006999, 2006.

Berthier, S., Chazette, P., Pelon, J., and Baum, B.: Comparison of cloud statistics from spaceborne lidar systems, Atmos. Chem. Phys., 8, 6965–6977, doi:10.5194/acp-8-6965-2008, 2008.

5

10

- Blanchard, C. D., Woodcock, A. H., and Cipriano, R. J.: The vertical distribution of concentration of sea sealt in the marine atmosphere near Hawaii, Tellus B, 36, 118–125, 1984.
- Bréon, F.-M. M., Vermeulen, A., and Descloitres, J.: An evaluation of satellite aerosol products against sunphotometer measurements, Remote Sens. Environ., 115, 3102–3111, doi:10.1016/j.rse.2011.06.017, 2011.
- Burton, S. P., Ferrare, R. A., Hostetler, C. A., Hair, J. W., Rogers, R. R., Obland, M. D., Butler, C. F., Cook, A. L., Harper, D. B., and Froyd, K. D.: Aerosol classification using airborne High Spectral Resolution Lidar measurements – methodology and examples, Atmos. Meas. Tech., 5, 73–98, doi:10.5194/amt-5-73-2012, 2012.
- ¹⁵ Cachier, H., Aulagnier, F., Sarda, R., Gautier, F., Masclet, P., Besombes, J.-L., Marchand, N., Despiau, S., Croci, D., Mallet, M., Laj, P., Marinoni, A., Deveau, P.-A., Roger, J.-C., Putaud, J.-P., Van Dingenen, R., Dell'Acqua, A., Viidanoja, J., Martins-Dos Santos, S., Liousse, C., Cousin, F., Rosset, R., Gardrat, E., and Galy-Lacaux, C.: Aerosol studies during the ES-COMPTE experiment: an overview, Atmos. Res., 74, 547–563, 2005.
- ²⁰ Charlson, R. J., Schwartz, S. E., Hales, J. M., Cess, R. D., Coakley Jr., J. A., Hansen, J. E., and Hoffman, D. J.: Climate forcing by anthropogenic aerosols, Science, 255, 423–430, doi:10.1126/science.255.5043.423, 1992.
 - Chazette, P.: The monsoon aerosol extinction properties at Goa during INDOEX as measured with lidar, J. Geophys. Res., 108, 4187, doi:10.1029/2002JD002074, 2003.
- ²⁵ Chazette, P., Couvert, P., Randriamiarisoa, H., Sanak, J., Bonsang, B., Moral, P., Berthier, S., Salanave, S., and Toussaint, F.: Three-dimensional surveyof pollution during winter in French Alps valleys, Atmos. Environ., 39, 1035–1047, 2005a.
 - Chazette, P., Randriamiarisoa, H., Sanak, J., Couvert, P., and Flamant, C.: Optical properties of urban aerosol from airborne and ground-based in situ measurements performed during the
- Etude et Simulation de la Qualité de l'air en Ile de France (ESQUIF) program, J. Geophys. Res., 110, D02206, doi:10.1029/2004JD004810, 2005b.
 - Chazette, P., Raut, J.-C., Dulac, F., Berthier, S., Kim, S.-W., Royer, P., Sanak, J., Loaëc, S., and Grigaut-Desbrosses, H.: Simultaneous observations of lower tropospheric continen-



tal aerosols with a ground-based, an airborne, and the spaceborne CALIOP lidar system, J. Geophys. Res., 115, D00H31, doi:10.1029/2009JD012341, 2010.

- Chazette, P., Bocquet, M., Royer, P., Winiarek, V., Raut, J. C., Labazuy, P., Gouhier, M., Lardier, M., and Cariou, J. P.: Eyjafjallajökull ash concentrations derived from both lidar and
- ⁵ modeling, J. Geophys. Res.-Atmos., 117, D00U14, doi:10.1029/2011JD015755, 2012a. Chazette, P., Dabas, A., Sanak, J., Lardier, M., and Royer, P.: French airborne lidar measurements for Eyjafjallajökull ash plume survey, Atmos. Chem. Phys., 12, 7059–7072, doi:10.5194/acp-12-7059-2012, 2012b.

Chazette, P., Marnas, F., and Totems, J.: The mobile Water vapor Aerosol Raman Lldar and

- its implication in the framework of the HyMeX and ChArMEx programs: application to a dust transport process, Atmos. Meas. Tech., 7, 1629–1647, doi:10.5194/amt-7-1629-2014, 2014.
 Cros, B., Durand, P., Cachier, H., Drobinski, P., Fréjafon, E., Kottmeier, C., Perros, P., Peuch, V., Ponche, J.-L., Robin, D., Said, F., Toupance, G., and Wotham, H.: The ESCOMPTE Program: an overview, Atmos. Res., 69, 241–279, 2004.
- Dieudonné, E., Chazette, P., Marnas, F., Totems, J., and Shang, X.: Lidar profiling of aerosol optical properties from Paris to Lake Baikal (Siberia), Atmos. Chem. Phys., 15, 5007–5026, doi:10.5194/acp-15-5007-2015, 2015.

Draxler, R. R. R. and Rolph, G. D. D.: HYSPLIT (HYbrid Single-Particle Lagrangian Integrated Trajectory) Model access via NOAA ARL READY Website, available at: http://www.arl.noaa.

- 20 gov/HYSPLIT.php (last access: November 2015), NOAA Air Resources Laboratory, College Park, MD, 2014.
 - Di Iorio, T., di Sarra, A., Junkermann, W., Cacciani, M., Fiocco, G., and Fuà, D.: Tropospheric aerosols in the Mediterranean: 1. Microphysical and optical properties, J. Geophys. Res., 108, D02201, doi:10.1029/2002JD002815, 2003.
- ²⁵ Dulac, F. and Chazette, P.: Airborne study of a multi-layer aerosol structure in the eastern Mediterranean observed with the airborne polarized lidar ALEX during a STAAARTE campaign (7 June 1997), Atmos. Chem. Phys., 3, 1817–1831, doi:10.5194/acp-3-1817-2003, 2003.

Flamant, C., Pelon, J., Chazette, P., Trouillet, V., Quinn, P. K., Frouin, R., Bruneau, D., Francois

Leon, J., Bates, T. S., Johnson, J., and Livingston, J.: Airborne lidar measurements of aerosol spatial distribution and optical properties over the Atlantic Ocean during a European pollution outbreak of ACE-2, Tellus B, 52, 662–677, doi:10.1034/j.1600-0889.2000.00083.x, 2000.



- Formenti, P., Boucher, O., Reiner, T., Sprung, D., Andreae, M. A., Wendisch, M., Wex, H., Kindred, D., Tzortziou, M., Vasaras, A., and Zerefos, C.: STAAARTE-MED 1998 summer airborne measurements over the Aegean Sea. 2. Aerosol scattering and absorption, and radiative calculations, J. Geophys. Res., 107, 4451, doi:10.1029/2001JD001536, 2002.
- ⁵ Giorgi, F. and Lionello, P.: Climate change projections for the Mediterranean region, Global Planet. Change, 63, 90–104, doi:10.1016/j.gloplacha.2007.09.005, 2008.
 - Gobbi, G. P., Barnaba, F., Blumthaler, M., Labow, G., and Herman, J. L.: Observed effects of particles nonsphericity on the retrieval of marine and desert dust aerosol optical depth by lidar, Atmos. Res., 61, 1–14, 2000.
- Groß, S., Tesche, M., Freudenthaler, V., Toledano, C., Wiegner, M., Ansmann, A., Althausen, D., and Seefeldner, M.: Characterization of Saharan dust, marine aerosols and mixtures of biomass-burning aerosols and dust by means of multi-wavelength depolarization and Raman lidar measurements during SAMUM 2, Tellus B, 63, 706–724, 2011.
- Hamonou, E., Chazette, P., Balis, D., Dulac, F., Schneider, X., Galani, E., Ancellet, G., and Pa payannis, A.: Characterization of the vertical structure of Saharan dust export to the Mediter ranean Basin, J. Geophys. Res., 104, 22257, doi:10.1029/1999JD900257, 1999.
- Holben, B. N., Eck, T. F., Slutsker, I., Tanré, D., Buis, J. P., Setzer, A., Vermote, E. F., Reagan, J. A., Kaufman, Y. J., Nakajima, T., Lavenu, F., Jankowiak, I., and Smirnov, A.: AERONET a federated instrument network and data archive for aerosol characterization, Remote Sens. Environ., 66, 1–16, 1998.
- IPCC: Climate Change 2014: Impacts, Adaptation, and Vulnerability. Part A: Global and Sectoral Aspects. Contribution of Working Group II to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, edited by: Field, C. B., Barros, V. R., Dokken, D. J., Mach, K. J., Mastrandresa, M. D., Bilir, T. E., Chatterjee, M., Ebi, K. L., Estrada, Y. O., Gen-
- ova, R. C., Girma, B., Kissel, E. S., Levy, A. N., MacCracken, S., Mastrandrea, P. R., and White, L. L., Cambridge University Press, Cambridge, UK and new York, NY, USA, 1132 pp., 2014.
 - Kim, S.-W., Berthier, S., Raut, J.-C., Chazette, P., Dulac, F., and Yoon, S.-C.: Validation of aerosol and cloud layer structures from the space-borne lidar CALIOP using a ground-based
- ³⁰ lidar in Seoul, Korea, Atmos. Chem. Phys., 8, 3705–3720, doi:10.5194/acp-8-3705-2008, 2008.



King, M. D., Kaufman, Y. J., Menzel, P., and Tanré, D.: Remote-sensing of cloud, aerosol, and water-vapor properties from the Moderate Resolution Imaging Spectroradiometer (MODIS), IEEE T. Geosci. Remote, 30, 2–27, 1992.

Klett, J. D.: Lidar inversion with variable backscatter/extinction ratios, Appl. Optics, 24, 1638, doi:10.1364/AO.24.001638, 1985.

5

20

- Lelieveld, J., Berresheim, H., Borrmann, S., Crutzen, P. J., Dentener, F. J., Fischer, H., Feichter, J., Flatau, P. J., Heland, J., Holzinger, R., Korrmann, R., Lawrence, M. G., Levin, Z., Markowicz, K. M., Mihalopoulos, N., Minikin, A., Ramanathan, V., de Reus, M., Roelofs, G. J., Scheren, H. A., Sciare, J., Schlager, H., Schultz, M., Siegmund, P., Steil, B.,
- ¹⁰ Stephanou, E. G., Stier, P., Traub, M., Warneke, C., Williams, J., and Ziereis, H.: Global air pollution crossroads over the Mediterranean, Science, 298, 794–799, 2002.

Le Treut, H., Somerville, R., Cubasch, U., Ding, Y., Mauritzen, C., Mokssit, A., Peterson, T., and Prather, M.: Historical overview of climate change, in: Climate Change 2007: The Physical Science Basis. Contribution of Working Group I to the Fourth Assessment Report of the

- ¹⁵ Intergovernmental Panel on Climate Change. edited by: Solomon, S., Qin, D., Manning, M., Chen, Z., Marquis, M., Averyt, K. B., Tignor, M., and Miller, H. L., Cambridge University Press, Cambridge, 129 pp., 2007.
 - Levin, Z., Teller, A., and Ganor, E.: On the interactions of mineral dust, sea-salt particles, and clouds: a measurement and modeling study from the Mediterranean Israeli Dust Experiment campaign, J. Geophys. Res., 110, D05207, doi:10.1029/2005JD005810, 2005.
- Mallet, M., Dulac, F., Formenti, P., Nabat, P., Sciare, J., Roberts, G., Pelon, J., Ancellet, G., Tanré, D., Parol, F., di Sarra, A., Alados, L., Arndt, J., Auriol, F., Blarel, L., Bourrianne, T., Brogniez, G., Chazette, P., Chevaillier, S., Claeys, M., D'Anna, B., Denjean, C., Derimian, Y., Desboeufs, K., Di Iorio, T., Doussin, J.-F., Durand, P., Féron, A., Freney, E., Gaimoz, C., Goloub, P., Gómez-Amo, J. L., Granados-Muñoz, M. J., Grand, N., Hamonou, E., Jankowiak, I., Jeannot, M., Léon, J.-F., Maillé, M., Mailler, S., Meloni, D., Menut, L., Momboisse, G., Nicolas, J., Podvin, J., Pont, V., Rea, G., Renard, J.-B., Roblou, L., Schepanski, K., Schwarzen-
- boeck, A., Sellegri, K., Sicard, M., Solmon, F., Somot, S., Torres, B., Totems, J., Triquet, S., Verdier, N., Verwaerde, C., Wenger, J., and Zapf, P.: Overview of the Chemistry-Aerosol
 Mediterranean Experiment/Aerosol Direct Radiative Forcing on the Mediterranean Climate (ChArMEx/ADRIMED) summer 2013 campaign Atmos Cham Phys. Discuss. 15, 19615–
 - (ChArMEx/ADRIMED) summer 2013 campaign, Atmos. Chem. Phys. Discuss., 15, 19615– 19727, doi:10.5194/acpd-15-19615-2015, 2015.



- Moulin, C., Lambert, C. E., Dayan, U., Masson, V., Ramonet, M., Bousquet, P., Legrand, M., Balkanski, Y. J., Guelle, W., Marticorena, B., Bergametti, G., and Dulac, F.: Satellite climatology of African dust transport in the Mediterranean atmosphere, J. Geophys. Res., 103, 13137–13144, 1998.
- Müller, D., Ansmann, A., Mattis, I., Tesche, M., Wandinger, U., Althausen, D., and Pisani, G.: Aerosol-type-dependent lidar ratios observed with Raman lidar, J. Geophys. Res.-Atmos., 112, D16202, doi:10.1029/2006JD008292, 2007.

Nabat, P., Somot, S., Mallet, M., Chiapello, I., Morcrette, J. J., Solmon, F., Szopa, S., Dulac, F., Collins, W., Ghan, S., Horowitz, L. W., Lamarque, J. F., Lee, Y. H., Naik, V., Nagashima, T.,

¹⁰ Shindell, D., and Skeie, R.: A 4-D climatology (1979–2009) of the monthly tropospheric aerosol optical depth distribution over the Mediterranean region from a comparative evaluation and blending of remote sensing and model products, Atmos. Meas. Tech., 6, 1287–1314, doi:10.5194/amt-6-1287-2013, 2013.

Nabat, P., Somot, S., Mallet, M., Michou, M., Sevault, F., Driouech, F., Meloni, D., di Sarra, A.,

¹⁵ Di Biagio, C., Formenti, P., Sicard, M., Léon, J.-F., and Bouin, M.-N.: Dust aerosol radiative effects during summer 2012 simulated with a coupled regional aerosol–atmosphere–ocean model over the Mediterranean, Atmos. Chem. Phys., 15, 3303–3326, doi:10.5194/acp-15-3303-2015, 2015a.

Nabat, P., Somot, S., Mallet, M., Sevault, F., Chiacchio, M., and Wild, M.: Direct and semi-direct

- aerosol radiative effect on the Mediterranean climate variability using a Regional Climate System Model, Clim. Dynam., 44, 1127–1155, doi:10.1007/s00382-014-2205-6, 2015b.
 Nisantzi, A., Mamouri, R. E., Ansmann, A., and Hadjimitsis, D.: Injection of mineral dust into
 - the free troposphere during fire events observed with polarization lidar at Limassol, Cyprus, Atmos. Chem. Phys., 14, 12155–12165, doi:10.5194/acp-14-12155-2014, 2014.
- Papayannis, A., Amiridis, V., Mona, L., Tsaknakis, G., Balis, D., Bösenberg, J., Chaikovski, A., De Tomasi, F., Grigorov, I., Mattis, I., Mitev, V., Müller, D., Nickovic, S., Pérez, C., Pietruczuk, A., Pisani, G., Ravetta, F., Rizi, V., Sicard, M., Trickl, T., Wiegner, M., Gerding, M., Mamouri, R. E., D'Amico, G., and Pappalardo, G.: Systematic lidar observations of Saharan dust over Europe in the frame of EARLINET (2000–2002), J. Geophys. Res., 113, D10204, doi:10.1029/2007JD009028. 2008.
 - Paronis, D., Dulac, F., Chazette, P., Hamonou, E., and Liberti, G. L.: Aerosol optical thickness monitoring in the Mediterranean, J. Aerosol Sci., 29, S671–S672, 1998.



Ramanathan, V. and Vogelmann, A. M.: Greenhouse effect, atmospheric solar absorption and the Earth's radiation budget: from the Arrhenius–Langley era to the 1990s, Ambio, 26, 38–46, 1997.

Ramanathan, V., Crutzen, P. J., Kiehl, J. T., and Rosenfeld, D.: Aerosols, climate, and the hydrological cycle, Science, 294, 2119–2124, doi:10.1126/science.1064034, 2001a.

- drological cycle, Science, 294, 2119–2124, doi:10.1126/science.1064034, 2001a.
 Ramanathan, V., Crutzen, P. J., Lelieveld, J., Mitra, A. P., Althausen, D., Anderson, J., Andreae, M. O., Cantrell, W., Cass, G. R., Chung, C. E., Clarke, A. D., Coakley, J. A., Collins, W. D., Conant, W. C., Dulac, F., Heintzenberg, J., Heymsfield, A. J., Holben, B., Howell, S., Hudson, J., Jayaraman, A., Kiehl, J. T., Krishnamurti, T. N., Lubin, D., McFar quhar, G., Novakov, T., Ogren, J. A., Podgorny, I. A., Prather, K., Priestley, K., Prospero, J. M.,
- Quinn, P. K., Rajeev, K., Rasch, P., Rupert, S., Sadourny, R., Satheesh, S. K., Shaw, G. E., Sheridan, P., and Valero, F. P. J.: The Indian Ocean Experiment: an integrated assessment of the climate forcing and effects of the great Indo–Asian haze, J. Geophys. Res., 106, 28371– 28393, 2001b.
- ¹⁵ Randriamiarisoa, H., Chazette, P., and Megie, G.: Retrieving the aerosol singlescattering albedo from the NO₂ photolysis rate coefficient, Tellus B, 56, 118–127, doi:10.3402/tellusb.v56i2.16408, 2004.

Raut, J.-C. and Chazette, P.: Radiative budget in the presence of multi-layered aerosol structures in the framework of AMMA SOP-0, Atmos. Chem. Phys. Discuss., 8, 12461–12528, doi:10.5194/acpd-8-12461-2008, 2008.

Royer, P., Chazette, P., Lardier, M., and Sauvage, L.: Aerosol content survey by mini N₂-Raman lidar: application to local and long-range transport aerosols, Atmos. Environ., 45, 7487–7495, 2011.

20

30

Salomonson, V. V., Barnes, W. L., Maymon, P. W., Montgomery, H. E., and Ostrow, H.: MODIS:

- ²⁵ advanced facility instrument for studies of the earth as a system, IEEE T. Geosci. Remote, 27, 145–153, 1989.
 - Smirnov, A., Holben, B. N., Eck, T. F., Dubovik, O., and Slutsker, I.: Cloud screening and quality control algorithms for the AERONET database, Remote Sens. Environ., 73, 337–349, 2000.
 Taylor, K. E. and Penner, J. E.: Response of the climate system to atmospheric aerosols and greenhouse gases, Nature, 369, 734–737, doi:10.1038/369734a0, 1994.
 - Tesche, M., Gross, S., Ansmann, A., Müller, D., Althausen, D., and Freudenthaler, V., and Esselborn, M.: Profiling of Saharan dust and biomass-burning smoke with multiwave-



length polarization Raman lidar at Cape Verde, Tellus B, 63, 649–676, doi:10.1111/j.1600-0889.2011.00548.x, 2011.

Thomason, L. W., Pitts, M. C., and Winker, D. M.: CALIPSO observations of stratospheric aerosols: a preliminary assessment, Atmos. Chem. Phys., 7, 5283–5290, doi:10.5194/acp-7-5283-2007, 2007.

5

Vaughan, M. A., Young, S. A., Winker, D. M., Powell, K. A., Omar, A. H., Liu, Z., Hu, Y., and Hostetler, C. A.: Fully automated analysis of space-based lidar data: an overview of the CALIPSO retrieval algorithms and data products, Proc. SPIE Int. Soc. Opt. Eng., 5575, 16– 30, doi:10.1117/12.572024, 2004.

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15, 32723–32757, 2015	
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Figure 1. Location of the Cap d'en Font surface station on Menorca Island, on a map of the MSG/SEVIRI-derived aerosol optical thickness at 550 nm over ocean, averaged over the campaign period (10 June–3 July 2013, daytime).





Figure 2. Result of the lidar inversion on the both the integrated AOT and the BER for the elastic (Klett (1985) algorithm) and N_2 -Raman channels: (a) the nights of 16–17 June, and (b) the nights of 27–28 June. The grey area highlights the gap between the two approaches.





Figure 3. Temporal evolution between 10 June and 3 July 2013, local time, of the aerosol optical thickness at 500 nm (AOT), the Ångström exponent between 440 and 675 nm (Ang) and the fine mode fraction (Frac) as derived from the sunphotometer measurements at Cap d'en Font. The AERONET products are completed by the Microtops II manual sunphotometer measurements on 10 June.





Figure 4. Temporal evolution of **(a)** the backscatter-to-extinction ratio (BER) and **(b)** the relative humidity (RH) between 10 and 29 June 2013. In **(a)** the root mean square error (rmse) is given by the grey area. The main aerosol categories have been identified in the figure as pollution, dust, local, marine and biomass burning aerosols using BER and PDR values (see text). The sunphotometer-derived BER is superimposed in red with its standard deviation (variability over half a day). In **(b)** the RH is calculated from lidar measurements using the thermodynamic temperature given by ECMWF analyses. During daytime the RH in the free troposphere is that of ECMWF; the boundary between the two RH determinations is highlighted by the continuous black line.





Figure 5. Backscatter-to-extinction ratio (BER) retrieved from **(a)** the synergy of the WALI lidar and the sunphotometer during daytime, **(b)** the coupling between the elastic and N_2 -Raman channels during night-time, and **(c)** the synthesis of daytime and night-time results.











Figure 7. (a) Results of aerosol speciation as given by lidar-derived extinction, PDR and BER, with backscatter coefficient coded as saturation (no saturation, white = 0, full saturation = $5 \times 10^{-6} \text{ m}^{-1} \text{ sr}^{-1}$); (b) key for the colours of the above. Night-time: dust-, pollution- and marine-like aerosols coded as red, green and blue respectively. Daytime: PDR coded as the saturation of red (top of the colour key). Intermediate colours and grey thus designate undetermined layers where aerosol mixing may occur.





Figure 8. (a) Scatter plots between SEVIRI and the ground-based sunphotometer of Menorca for the aerosol optical thickness AOT at 550 nm. The dotted line corresponds to the best fit against both retrievals. **(b)** The Ångström exponent for similar spectral ranges. A total of 846 coincident data pairs between 10 and 30 June are available for comparison. To the four aerosol types identified in Fig. 7 are added 3 mixed types which are all distinguished by their colour: pollution (Pol), dust (Dust), marine (March) and biomass burning (BB), mixing of pollution and marine (Pol March), Marine and dust (Mar Dust), dust and pollution (Dust Pol). The solid curve represents the centroid of the distribution.





Figure 9. AOT composition between the MODIS observations over land and SEVIRI over sea. The SEVIRI AOT was corrected by a factor 0.65 as identified from comparisons with the sunphotometer reported in Fig. 8. Top-left panel (a) is for 17 June, top-right (b) for 26 June, bottom-left (c) for 27 June, and bottom-right (d) for 28 June. For 28 June the night-time CALIOP ground track (at about 02:00 UTC) is marked by a continuous grey line.





Figure 10. 27 back trajectories from Menorca (39°51′44″ N, 4°15′30″ E). The back trajectories have been computed using the ensemble mode of the HYSPLIT model (courtesy of NOAA Air Resources Laboratory; http://www.arl.noaa.gov). The end location of the air mass trajectories is at 6 km a.m.s.l. in the plume detected by the WALI lidar on 27 June 2013 over Menorca (see Fig. 6a). The top panel presents the location of each back trajectory superimposed over the MODIS-derived AOT at 550 nm of 24 June 2013, for several orbits. The bottom panel shows the altitude of the back trajectories against time.

