



## Consistency of aerosols above clouds characterisation from A-Train active and passive measurements

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10 **Abstract.** This study presents a comparison between the retrieval of aerosol above clouds (AAC) optical properties from different techniques developed for the A-Train sensors CALIOP and POLDER/PARASOL. The main objective is to analyse the consistency between the results of the active and the passive measurements. We assess the Aerosol Optical Thickness (AOT) of above optically thick clouds (Cloud Optical Thickness (COT) larger than 3) and their Ångström Exponent (AE). These parameters are retrieved with the CALIOP operational method, the POLDER operational polarization method and the  
15 CALIOP-based depolarization ratio method (DRM) – for which we also propose a calibrated version (denominated DRM<sub>SODA</sub>, SODA as Synergized Optical Depth of Aerosols). We analyse six months of data over three distinctive regions characterized by different types of aerosols and clouds. Additionally, for these regions, we select three case studies: a biomass-burning event over the South Atlantic Ocean, a Saharan dust case over the North Atlantic Ocean and a Siberian biomass-burning event over the North Pacific Ocean. 4.5 years of data are studied over the entire globe for distinct situations  
20 where aerosol and cloud layers are in contact or vertically separated. Overall, the regional analysis shows a good correlation between the POLDER and the DRM<sub>SODA</sub> AOTs when the microphysics of aerosols is dominated by fine-mode particles of biomass-burning aerosols from southern Africa (correlation coefficient ( $R^2$ ) of 0.83) or coarse-mode aerosols of Saharan dust ( $R^2$  of 0.82). A good correlation between these methods ( $R^2$  of 0.68) is also observed in the global treatment, when the aerosol and cloud layers are well separated. The analysis of detached layers also shows a mean difference in AOT of 0.07 at  
25 532 nm between POLDER and DRM<sub>SODA</sub>, at a global scale. The correlation between the retrievals decreases when a complex mixture of aerosols is expected ( $R^2$  of 0.37) – as in the East Asia region, and when the aerosol-cloud layers are in contact ( $R^2$  of 0.36). The correlation coefficient between the CALIOP operational method and POLDER is low, as the CALIOP method largely underestimates the aerosol loading above clouds by a factor that ranges from two to four.

Potential biases on the retrieved AOT as a function of cloud properties are also investigated. For different types of  
30 scenes, the retrieval of above-cloud AOT from POLDER and from DRM are compared for different underlying cloud properties (droplet effective radius ( $r_{eff}$ ) and COT retrieved with MODIS). The results reveal that DRM AOT vary with  $r_{eff}$ . When accounting for  $r_{eff}$  in the DRM algorithm, the consistency between the methods increases. The sensitivity study shows



that an additional polarized signal coming from aerosols located *within* the cloud could affect the polarization method, which leads to an overestimation of the AOT retrieved with POLDER algorithm. In addition, the aerosols *attached* or *within* the cloud can potentially impact the DRM retrievals through the modification of the cloud droplet chemical composition and its ability to backscatter light. The next step of this work is to combine POLDER and CALIOP to investigate the impacts of aerosols on clouds and climate when these particles are transported above or within clouds.

## 1 Introduction

By interacting with radiations and by modifying the cloud reflectivity and the cloud formation, aerosols have important impacts on the Earth's radiative budget and water cycle (IPCC, 2014). These atmospheric particles absorb and scatter the sunlight, resulting in the so-called "Direct Radiative Effect" (DRE). Although aerosols always produce a cooling effect at the Earth's surface, the sign and the amplitude of the DRE of aerosols at the top of the atmosphere depends on the aerosol properties but also on the underlying surface. For instance, in cases where absorbing aerosol layers are located above clouds, the DRE of aerosols is predominantly positive as a result of the reduction of the local planetary albedo (Keil and Haywood, 2003). By absorbing sunlight, aerosols also warm the layer of the atmosphere where they reside. This modifies the vertical profile of temperature in the atmosphere, which may affect the process of evaporation and cloud formation. This effect is called the "semi-direct effect" (Hansen et al., 1997; Ramanathan et al., 2001). Aerosols also impact the cloud properties by acting as cloud condensation nuclei and ice nuclei. They may modify the cloud microphysics and cloud brightness with potential impacts on precipitation and cloud lifetime (Rosenfeld, 2000; Twomey, 1974). These effects are referred as "aerosol indirect effects" and tend to cool the Earth.

The lack of knowledge of aerosol properties in cases of aerosols above clouds (AAC) scenes has been recently highlighted as a source of uncertainty for the estimation of the DRE of aerosols (Peers et al., 2016). Different approaches have been developed to quantify the DRE of AAC using satellite observations (Chand et al., 2009; Meyer et al., 2013). But despite recent observational and modelling studies (De Graaf et al., 2014; Peers et al., 2015, 2016; Zhang et al., 2016), the aerosol DRE for AAC remains subject to large discrepancies. In the process of quantification and interpretation of the aerosol impact on climate, the aerosol interactions with clouds constitute the largest uncertainty in global climate models (Myhre et al., 2013a, 2013b). The study of AAC may also contribute to reduce those uncertainties. For instance, in case of absorbing AAC, the warming of the atmosphere occurring above stratocumulus clouds might reduce the strength of the convection and consequently impact the vertical development and the cloud properties. This warming might inhibit the entrainment of dry air at the top of the cloud, preserving the humidity of the cloud and increasing the liquid water content and the persistence of clouds (Johnson et al., 2004; Wilcox, 2010). Evidence of the first indirect effect was also found over the South Atlantic region, where AAC events are frequently observed. **Costantino and Bréon (2012) notably found a strong decrease in the droplet effective radius when the aerosol layers are touching the top of the cloud layer.**



The scientific community is working on better monitoring the load and microphysical properties of AAC in order to assess the influence of those particles on the Earth's radiative budget and clouds. The constellation of satellites called A-Train provides different passive and active sensors for monitoring clouds and aerosols (<http://atrain.nasa.gov/publications/A-TrainFactSheet.pdf>). Passive techniques have a large spatial coverage, but have no direct information of the vertical distribution of particles in the atmosphere. Active methods offer unique capabilities, complementary with the passive methods and are dedicated to the study of the vertical profiles of clouds and aerosols. The main retrieved optical properties for aerosols are the Aerosol Optical Thickness (AOT), the Ångström Exponent (AE), which is a parameter of the particles size, and the aerosol Single Scattering Albedo (SSA) (Torres et al., 2012; Waquet et al., 2016).

The active sensor Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) installed on CALIPSO (Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observation) satellite provides high-resolution vertical profiles of aerosols and clouds (Chand et al., 2008; Winker et al., 2003). CALIOP provides the total attenuated backscatter signal ( $\text{km}^{-1} \text{sr}^{-1}$ ) at 532 nm and 1064 nm. From the backscatter measurements, an operational aerosol algorithm allows for retrieval of the vertical extinction profiles as well as the AOT in clear and cloudy skies by assuming an aerosol lidar ratio (extinction to backscatter) (Vaughan et al., 2009; Winker et al., 2009). Moreover, two orthogonally polarized channels measure the parallel and perpendicular backscatter signal at 532 nm that allows calculating the depolarization ratio (i.e. the ratio of the two orthogonal polarization signals) (Winker et al., 2007). Depolarization measurements are used for discrimination between spherical and non-spherical particles (Sassen, 1991). CALIOP provides exhaustive details on the vertical distribution of optical and microphysical properties of aerosols and clouds, including their shape, and a qualitative classification of aerosol type (via the wavelength dependence of the backscatter) (Winker et al., 2009; Young and Vaughan, 2009).

Alternative CALIOP-based research methods have also been introduced to retrieve Above-Cloud AOT (ACAOT). The depolarization ratio method (DRM) (Hu et al., 2007a) and the color ratio method (CRM) (Chand et al., 2008) use fewer assumptions for the retrieval of aerosol properties. These methods are based on light transmission methods and treat the liquid water clouds situated underneath the aerosol layer as a target. Hu et al. (2007b) have shown that, in the case of opaque water clouds, the layer integrated attenuated backscatter at 532 nm and layer integrated attenuated depolarization ratio at 532 nm can be used to retrieve the aerosol optical depth of the overlying aerosol or optically thin cloud layers. The CRM uses the layer integrated attenuated color ratio, which is the ratio of integrated attenuated backscatter at 1064 nm to 532 nm. Over the visible to near-infrared spectral region, fine-mode absorbing aerosols above clouds exhibit a strong wavelength dependence color ratio (Chand et al., 2008). This makes possible the detection of absorbing biomass-burning aerosols transported above clouds. The color ratio observed in the case of coarse mode particles or purely scattering fine mode aerosols transported above clouds exhibits little or no wavelength dependence and thus, these particles can be less accurately detected with the CRM method.

Passive sensors have also been used to obtain information on aerosols above clouds. For example, Torres et al. (2012) have developed an algorithm to retrieve the ACAOT and the underlying aerosol-corrected cloud optical depth, using radiance measurements performed in the ultra violet (UV) by the Ozone Monitoring Instruments (OMI). The method takes



5 advantage of the ability of biomass burning and mineral dust aerosols to strongly absorb UV radiations. Another method that can retrieve the ACAOT and, simultaneously, the aerosol-corrected COT is the “color ratio” method proposed by Jethva et al. (2013) that employs measurements in visible and shortwave infrared (SWIR) channels from the Moderate Resolution Imaging Spectroradiometer (MODIS). Also, Meyer et al. (2015) developed an algorithm that employs reflectance measurements from six MODIS channels (from the visible to the shortwave infrared) to retrieve the ACAOT, as well as the COT and droplet effective radius ( $r_{eff}$ ) of the underlying cloud.

10 The multi-directional polarization measurements have shown sensitivity to AAC scenes (Waquet et al., 2009, Hasekamp, 2010; Knobelspiesse et al., 2011). Polarization and Directionality of Earth Reflectances (POLDER) instrument is measuring the directionality and polarization of light reflected by the Earth-atmosphere system. The aerosols generate an additional polarized light at forward and side scattering angles ( $70^\circ$  to  $130^\circ$ ) and reduce the polarized signal of the cloud bow (i.e. a strong polarized rainbow feature observed near  $140^\circ$  in scattering angle). Mineral dust particles do not much polarize light, but they strongly minimize the cloud bow magnitude. Based on these effects, Waquet et al. (2009) have developed a method for retrieving the properties of aerosols above clouds that relies on the polarized radiances measured by POLDER. Because polarized radiances are not affected by the optical thickness of the cloud (i.e. the polarized radiance reflected by the cloud is saturated when the cloud is optically thick enough), the method is able to retrieve the scattering ACAOT at two wavelengths (670 nm and 865 nm) without much assumption on cloud properties. An analysis of the global results obtained with the operational algorithm is given in Waquet et al. (2013a). Furthermore, Peers et al. (2015) have developed a complementary method that uses additional total multidirectional radiances measured by POLDER. The method provides the aerosol SSA and the aerosol-corrected cloud optical thickness. So far, the algorithm of Peers et al. (2015) is a research method, only applied for regional studies (Peers et al., 2016).

25 Jethva et al. (2014) performed an intercomparative analysis of the ACAOT retrieved with the aforementioned methods, in order to check the consistency among independent techniques. The results were encouraging and, despite the use of different assumptions and measurements, a close agreement was reported over homogeneous clouds. Similar to this study, our paper will focus on the comparison between collocated active and passive AAC inversion products, improving our understanding of the ACAOT. But, compared to Jethva et al. (2014), who focused only on two study cases, **we perform a global and multi-annual investigation to provide robust statistics results**. The vertical distribution of the aerosol and cloud layer will be also considered. We will concentrate on the following methods: (a) the CALIOP operational method (CALIOP<sub>OM</sub>) because of the numerous studies in which it was used, (b) the DRM developed by Hu et al. (2007a), (c) a calibrated version of the DRM algorithm and (d) the POLDER polarization method. The DRM and POLDER methods were chosen because both are measuring AAC properties above the same type of cloudy scenes (i.e. optically thick and homogeneous liquid water clouds). Moreover, both techniques are sensitive to all types of particles (scattering or absorbing particles, fine or coarse ones), which is not the case for CRM that can operate only for absorbing aerosols. It is also interesting to compare these two approaches since the POLDER method requires a hypothesized aerosol microphysics, while



DRM doesn't require any hypothesis for the aerosols but requires assumptions and an approximate model to estimate the signal backscattered by clouds.

To begin with, we briefly recall the principle of each algorithm and the data selection strategy. The results of AOT intercomparison are presented in sections 3 and 4. We first present a regional comparison and then describe a global comparison for a period of 4.5 years in function of the type of aerosols and AAC scenes (aerosol and cloud layers in contact or well separated). Discussions and conclusions will be drawn in sections 5 and 6.

## 2 Methodology and data selection

### 2.1 POLDER polarisation method

POLDER, an instrument on the PARASOL (Polarization and Anisotropy of Reflectances for Atmospheric Science coupled with Observations from a Lidar) satellite, is a wide-field imaging radiometer/polarimeter (Tanré et al., 2011). This instrument measures the angular and spectral behaviour of the normalized total and polarized radiances (Herman et al., 2005).

The method for retrieving the above-cloud scattering AOT developed by Waquet et al. (2013b) consists of a comparison between polarized radiances measured by POLDER at 670 nm and 865 nm and polarized radiances pre-computed with a Successive Order of Scattering (SOS) code (Deuzé et al., 1989) for seven aerosol models that follow a single lognormal size distribution. Six models correspond to spherical aerosols (fine-mode particles) with radius from 0.06 to 0.16  $\mu\text{m}$ , for which a complex refractive index of  $1.47-0.01i$  is assumed. The seventh model is bimodal and characteristic of non-spherical aerosols (dust) with a refractive index of  $1.47-0.0007i$ . In the search for the best fitting aerosol model, the operational algorithm follows the strategy described by Waquet et al. (2013b). After a first step, the algorithm produces an approximation of the AOT at 865 nm. As a function of this AOT value, a decision tree is applied: if the AOT is larger than 0.1 then the algorithm will search the best fitting model within all the seven models without any angular constraint for the selection of the POLDER data (scattering angle ranging from  $0^\circ$  to  $180^\circ$ ). Next, if the mineral dust model fails to reproduce the data or if the AOT retrieved in the first step is smaller than 0.1, then only fine-mode models are considered in the retrieval scheme and the viewing geometries are restricted to side or forward viewing geometries (scattering angles smaller than  $130^\circ$ ). The AOT threshold of 0.1 at 865 nm is empirical and was introduced since the retrieval of the aerosol type (dust or fine mode particles) becomes difficult for small AOT.

Collocated cloud properties retrieved from MODIS (Moderate Resolution Imaging Spectroradiometer) at high resolution ( $1 \times 1 \text{ km}^2$  at nadir) are used to characterize and to select the cloudy scenes within a POLDER pixel ( $6 \text{ km} \times 6 \text{ km}$  at nadir). We only consider fully covered cloudy pixels associated with optically thick liquid water clouds: the cloud optical thickness retrieved by MODIS has to be larger than three and a cloud phase algorithm is applied to select liquid water clouds (Riedi et al., 2010). Moreover, Waquet et al. (2013b) have introduced a mask to eliminate cirrus above liquid clouds that makes use of the MODIS Brightness Temperature Difference (BTD) between 8.5 and 11  $\mu\text{m}$  wavelength bands as well as



MODIS and POLDER cloud top pressure estimates. Lastly, the AOT retrievals at the high-POLDER pixel resolution are aggregated to  $18 \text{ km} \times 18 \text{ km}$ . The retrieved solution is kept if the number of  $6 \text{ km} \times 6 \text{ km}$  pixels is larger than 5 and if the standard deviation computed for the mean AOT is smaller than 0.1. This latter procedure allows to remove edges of clouds. In our study, we use the version 3.00 of the official output product PARASOL\_PM02-L2 for AAC scenes available at  
5 ICARE website (<http://www.icare.univ-lille1.fr/parasol/products/>).

## 2.2 CALIOP methods

### 2.2.1 Operational method

The CALIPSO lidar (CALIOP) is a frequency-doubled Nd:YAG laser, dual-wavelength, dual-polarization, elastic backscatter lidar (Winker et al., 2003) The lidar returned signal is normalized and range-corrected to provide the total  
10 attenuated backscatter coefficient ( $\text{km}^{-1} \text{sr}^{-1}$ ).

In order to retrieve the attenuated backscatter data and the columnar AOT at 532 nm and 1064 nm, the operational CALIOP algorithm combines the feature and layer detection scheme (Winker al., 2009) with the extinction retrieval algorithm (Young and Vaughan, 2009) that employs assumptions on the extinction-to-backscatter ratio of aerosols. There are several steps involved in the operational data processing: 1) cloud and aerosol layers are detected in the backscattered signal  
15 along with their altitudes, 2) the algorithm determines which layers have cloud or aerosol features, 3) the cloud ice-water phase is estimated and the aerosol lidar-ratio is determined, using assumptions on the aerosol models, and finally, 4) the extinction coefficients and AOT are retrieved at 532 and 1064 nm.

Lidar systems have a limited capability to determine the composition and size of aerosols. Hypotheses are then used on the aerosol phase function at  $180^\circ$  and on the aerosol single scattering albedo (SSA) in order to calculate the aerosol lidar  
20 ratio. In the operational algorithm, the aerosol models consist in a mixture of aerosol components characteristic of a region or an air mass. **It should be noted that an incorrect assumption for the lidar ratio is a source of substantial errors in the AOT retrieved with this method.**

**For our study we use the level 2 version 2.01 of the inversion products**, officially named CAL\_LID\_L2\_05kmALay (ALay) and CAL\_LID\_L2\_05kmCLay (CLay) (that can be found at <http://www.icare.univ-lille1.fr/calipso/products/>), which  
25 provide respectively the aerosol and cloud layer parameters at a nominal horizontal resolution of 5 km. From these products we used the AOTs retrieved at 532 nm and 1064 nm, the aerosol base and top altitudes, the cloud top altitude, the ice-water cloud phase and the feature type. We also use CALIOP level 1 dataset, labelled CAL\_LID\_L1-ValStage1 (link above) that provides the attenuated backscatter coefficient calculated at a vertical resolution of 30 m and 333 m and at a horizontal resolution from -0.5 to 8.2 km altitude (Winker et al., 2007).



### 2.2.1 Depolarisation Ratio Method

For retrieving the optical thickness of a thin high layer (aerosols or clouds) above a lower and optically thick water cloud layer, Hu et al. (2007a) and Chand et al. (2008) describe the depolarization ratio method applied to CALIOP measurements. An opaque cloud with a minimum optical depth of three will attenuate the lidar beam completely. For optically thick clouds, we estimate the optical thickness of the above thin aerosol or cloud layer by treating the opaque cloud as a target and by using the Beer-Lambert law to estimate the direct transmission of light above this cloud layer. We will refer to this product hereafter as  $DRM_{Hu}$ .

The physical properties used in this method are the cloud attenuated backscatter coefficient ( $\gamma'_{water}$ ) integrated from the base to the top of the cloud layer at 532 nm and the integrated attenuated depolarization ratio ( $\delta'$ ) at 532 nm. The definition of  $\gamma'_{water}$  is given by the following equation:

$$\gamma'_{water} = \int_{z_{top}}^{z_{base}} \beta'(z) dz, \quad (1)$$

where  $\beta'$  is the total attenuated backscatter coefficient ( $km^{-1} sr^{-1}$ ).

In situations where the cloud is optically thick and there are no aerosol above the cloud, the lidar equation simplifies to the following definition, expressed as a function of the lidar ratio ( $S_c$ ) and layer effective multiple scattering factor ( $\eta_c$ ):

$$\gamma'_{water,calc} = \int_{z_{top}}^{z_{base}} \beta'(z) dz = (2\eta_c S_c)^{-1}, \quad (2)$$

$S_c$  is narrowly constrained to about 19 sr at a wavelength of 532 nm. This value is typically used for liquid water clouds with droplets smaller than about 50  $\mu m$  (O'Connor et al., 2004; Pinnick et al., 1983).  $\eta_c$ , which takes values between 0 and 1, is strongly related to the cloud depolarization ratio  $\delta'$ , since multiple scattering processes tend to depolarize light. An approximate relation was derived from Monte Carlo simulations (Hu et al., 2006):

$$\eta_c = \left( \frac{1-\delta'}{1+\delta'} \right)^2, \quad (3)$$

After  $\gamma'_{water}$  is corrected for molecular and gaseous absorption, the ratio between  $\gamma'_{water}$  and  $\gamma'_{water,calc}$  should be equal to 1 in the absence of higher aerosol or cloud layer, and with an accurate lidar calibration. Instead, in case of an overlying aerosol or cloud layer, this ratio can be written as

$$\frac{\gamma'_{water}}{\gamma'_{water,calc}} = T^2 = \exp(-2\tau_{top,DR}), \quad (4)$$

where  $T^2$  is the transmission of light after a two-way propagation between the sensor and the targeted cloud, and  $\tau_{top,DR}$  is the higher layer's optical thickness. It follows from Eq. (4) that the optical depth ( $\tau_{top,DR}$ ) is given by:

$$\tau_{top,DR} = \frac{-1}{2} \ln(2S_c \gamma'_{water} \eta_c), \quad (5)$$



**DRM<sub>Hu</sub> differs from the operational method by the fact that it does not rely on assumptions related to aerosol microphysical properties** (aerosol phase function and SSA) to estimate the AOT integrated over the atmospheric column. The main uncertainties of the DRM<sub>Hu</sub> are linked to the calibration of the lidar, which impact the estimate of the parameters in Eq. (5).

5 Providing a robust, self calibrated method at global scale and for the whole CALIPSO dataset is not trivial and in order to improve the estimate of the AOT with the DR method, the creators of the Synergized Optical Depth of Aerosols and ICE clouds (SODA & ICE, available at the ICARE thematic center), Josset et al. (2010; 2012), modified the original formalism of DRM<sub>Hu</sub>. The main reason for these modifications is that the relationship between the multiple scattering factor and the depolarization by the cloud shows a systematic deviation from the theory (see Fig. 2 in Hu, 2007). The multiple  
10 scattering–depolarization relationship has been confirmed by laboratory experiments (Cao et al., 2009). Even if it has to be modified in presence of submicrometer or non-spherical particles, the origin of the discrepancy between theory and observation points towards an instrumental issue. The long transient response of the receiver has been proposed as an explanation and a correction was also proposed (Hu et al., 2007b). There are, however other issues related to the calibration of the polarization channel that could explain the discrepancy. **The low gain/ high gain merging scheme and the day/night  
15 calibration transfer are a significant source of uncertainty (Sassen and Zhu, 2009).**



In order to overcome these difficulties and improve the accuracy of the method, SODA takes advantage of the high number of CALIOP observations of liquid water clouds in the absence of AAC. Practically, the SODA algorithm introduces global scale correction factors in the multiple scattering coefficient to depolarization relationship and a recalibrated value of the liquid water cloud lidar ratio as a function of latitudes. These two corrections come from the fact that, when the liquid  
20 water clouds are optically dense and in absence of AAC, the lidar equation can be reduced to Eq. (2). Over the ocean, the lidar ratio of most liquid water clouds is relatively constant (Hu et al., 2006) and the multiple scattering coefficient can be measured directly if the lidar is well calibrated. This correction follows the original intent of DRM<sub>Hu</sub> (Hu et al., 2007a), which has always been to be a self calibrated method, unaffected by instrumental or geophysical uncertainties (see Eq. (4) of Hu et al. (2007a) and related discussion). However, because the discrepancy between theory and observations is due to an  
25 instrumental artefact linked to the receiver electronics, SODA introduces a clearer separation between the parallel and perpendicular channel than in DRM<sub>Hu</sub> (Hu et al., 2007b). DRM<sub>Hu</sub> relates the total backscatter coefficient to the ratio of perpendicular and parallel backscatter coefficient while SODA links the parallel backscatter coefficient to this ratio. This approach is supported by the theory of light propagation in dense medium where the contribution of multiple scattering to the perpendicular and parallel channel is identical (Xu and Alfano, 2005) and by the analysis of CALIOP data.

30 A preliminary and mandatory step of the calibration procedure is to select optically opaque liquid water clouds with no AAC. The calibration modules of SODA use the following criteria. Note that there is some level of redundancy in order to increase data quality selection.

a) Criteria of optical density:



- the top and bottom of the cloud is given by the 333 m CALIPSO cloud product. This ensures a minimum level of signal strength and the presence of a transparent atmosphere above it.

- the maximum of the lidar signal is above the base of the cloud. This ensures a minimum level of attenuation of the signal from the surface.

5 - the ocean surface integrated attenuated backscatter is below a detectability threshold of  $7.5 \times 10^{-6} \text{ km}^{-1} \text{ sr}^{-1}$  for nighttime data and  $1 \times 10^{-3} \text{ km}^{-1} \text{ sr}^{-1}$  for daytime data. The intent of this threshold is the same as the previous criteria. More specifically, the goal is to use a threshold such that half the shots are below the noise sensitivity of the instrument.

b) Criteria of cloud in liquid phase

10 - the temperature at the top of the cloud is higher than  $0^\circ\text{C}$ . The isotherm is defined by the GMAO (Global Modelling and Assimilation Office) temperature when interpolated on the CALIPSO vertical grid.

- the total cloud liquid water contained in a vertical column of atmosphere retrieved from collocated pixels of AMSR-E/AMSR2 is larger than 0 mm.

c) Criteria of clear air above the cloud

15 - the total 532 nm integrated attenuated backscatter coefficient from 20 km of altitude to the top of the cloud is below the following threshold:

$$\int_{z_{\text{top}}}^{20\text{km}} \beta'(z) dz < \frac{1 - \exp(-2\tau_{\text{air,mol}})}{2^{\frac{8\pi}{3}} 1.5}, \quad (6)$$

where  $\tau_{\text{air,mol}}$  is the optical depth due to air Rayleigh scattering and ozone absorption. The factor 1.5 allows reducing the occurrences of false positives due to noise. It also allows to simplify the formalism as **the King color** (Bates, 1984; King, 1923) can be neglected with no expected impact on the results.

20 As a first step, SODA calibrates the multiple scattering to depolarization relationship for nighttime data on a monthly basis. For all opaque liquid water clouds defined with the above criteria, SODA looks for the least square fit between the data and  $\eta_{\text{calibr}}$  defined as:

$$\eta_{\text{calibr}} = A\eta_c + B\eta_c^2, \quad (7)$$

The data of interest are based on Eq. (2) and can be written as

25 
$$\eta_{\text{calibr}} = \frac{1}{2 \times 19 \times \gamma'_{\text{water,parallel}}}, \quad (8)$$

where  $\gamma'_{\text{water,parallel}}$  is the parallel-integrated backscatter coefficient.

As a second step, SODA calculates the apparent lidar ratio  $S_{c,\text{lat}}$  of all opaque liquid water clouds as a function of latitude. This procedure is done separately for daytime and nighttime data. **The latitudinal dependency is aimed at correcting the calibration inaccuracies of CALIOP, which are dependent on latitude (Powell et al., 2010)** and possible geophysical variations of cloud microphysical properties between the northern and southern hemisphere.

30



$$S_{c,lat} = \frac{1}{2 \times \eta_{calibr} \times \gamma'_{water,parallel}}, \quad (9)$$

All these coefficients are finally integrated in the AOT retrieval equation:

$$\tau_{top,DRcalibr} = -\frac{1}{2} \ln(2S_{c,lat} \eta_{calibr} \gamma'_{water,parallel}), \quad (10)$$

Through this study, we will refer to this product as  $DRM_{SODA}$  and can be found at ICARE data center  
5 (<http://www.icare.univ-lille1.fr/projects/soda/>).

## 2.3 Data selection

### 2.3.1 Collocation

The satellites pass through close orbits within several minutes, providing coincident observations of POLDER,  
MODIS and CALIOP instruments. Using the nearest pixel approximation, CALIOP files are used as a space reference for  
10 sampling POLDER and MODIS products. CALTRACK is the output dataset and can be found at ICARE data and service  
center. It contains coincident data from POLDER at  $18 \text{ km} \times 18 \text{ km}$  and MODIS extracted under the CALIOP track at 5 km  
horizontal resolution. The  $DRM_{Hu}$  and  $DRM_{SODA}$  optical depth retrievals are processed at the CALIOP native resolution of  
333 m and aggregated afterwards at 5 km horizontal resolution. Moreover, for a better consistency of the AOT comparison,  
the POLDER AOT was extrapolated at 532 nm using the AE retrieved with the POLDER algorithm.

15 We also limited the cloud top altitude at 5 km because we are interested in low-level clouds. The maximal aerosol  
top altitude is limited at 10 km. This maximal value should be sufficient, since most of the biomass burning and dust aerosol  
layers are typically observed between 0.5 and 4.0 km over ocean (Torres et al., 2013).

### 2.3.2 Distinction between vertical profiles

20 Additionally, we have employed an approach that is similar to the concept of Costantino and Bréon (2012) to  
classify the type of AAC scenes. The respective positions of the aerosol and cloud layers are defined using the CALIOP  
ALay and CLay products. We classify the AAC scenes into three categories: “attached”, “detached” and “undetermined”.  
The so-called “attached cases” correspond to situations where the aerosol layer touches the top of the beneath cloud layer.  
For these cases, we assume that the vertical distance of the aerosol bottom altitude from cloud top altitude must be lower  
than 100 meters. Inversely, the “detached cases” correspond to aerosol and cloud layers that are considered well separated,  
25 considering a distance higher than 500 m between the aerosol base altitude and the cloud top. Aerosol and cloud layers  
within a distance between 100 and 500 meters are considered too uncertain and are excluded from our study. We also  
removed the situations for which the aerosols are located within or below the cloud layers, assuming that these data are too  
uncertain. Practically, we rejected the CALIOP data for which the aerosol layer penetrates the cloud layer by more than 50



10 meters. The third category, called “undetermined” corresponds to situations for which the respective position of the aerosol  
or cloud layer cannot be determined by CALIOP (i.e. missing data).



We also distinguish the “two layer situation” (i.e. one aerosol layer and one cloud layer) from the “multiple layer  
situations” (more than one aerosol layer and/or more than one cloud layer). These latter situations are filtered in our analysis  
5 for the sake of simplicity (see Sect. 3.4 and Sect. 4).

### 3 Regional analysis and case studies



The results presented in this section were acquired from May to October 2008. We selected **three distinctive regions**  
that are under the influence of various aerosol species and different types of clouds: a) an area that extends from 30° S to 5°  
N and 12° W to 14° E over the South Atlantic Ocean (SAO), b) an area between 10 to 35° N and 10 to 40° W over the North  
10 Atlantic Ocean (NAO) and c) an area located between 35 to 60° N and 140 to 170° E over the North Pacific Ocean (NPO).  
The south of the African continent is the main contributor to biomass-burning aerosols above clouds, originating from man-  
made crops fires (Waquet et al., 2013b). These aerosols are highly absorbing (SSA of approximately 0.84 at 865 nm) and  
associated with high AE values; they mainly contribute to the fine mode. The NAO area is mainly under the influence of  
dust aerosols originating from the Saharan Desert for the time period of interest. These particles are mainly non-spherical  
15 and contribute primary to the coarse mode. They are moderately absorbing at the wavelength of CALIOP (532 nm) and  
almost non-absorbing at 865 nm (SSA of approximately 0.98) (Balkanski et al., 2007; Dubovik et al., 2002; Peers et al.,  
2015). The North Pacific Ocean (NPO) is associated with various types of particles: fine mode aerosols with rather scattering  
properties originating from man-made pollution (Waquet et al., 2013a; Yu et al., 2008), biomass-burning from forest fires  
(Peers et al., 2015; Zhang et al., 2003) and dust originated from the Asian deserts. Potential mixture of these different species  
20 is also possible for this area (Yu et al., 2006).

Cloud **properties** expected to be different in these three regions (Warren et al. 1988). Low-level stratocumulus  
clouds typically cover the SAO, with some occurrences of cumulus and altostratus clouds. Cumulus, altostratus clouds and  
some stratocumulus clouds generally cover the NAO. The cloud cover is generally fractional over this part of the Atlantic  
Ocean. Stratocumulus clouds also frequently cover the NPO. Higher altostratus and cumulus clouds are also often observed  
25 over this area. Cirrus clouds can be frequently found at mid-latitudes and also in the intertropical convergence zone, which  
includes the NPO and the NAO regions.

We studied six months of data over each region to observe the consistency between different techniques for various  
types of aerosols. For this part of the study, we mixed the “two-layer” and “multiple layer” situations and we analysed all the  
data, disregarding the position of the aerosol and cloud layers. A case study was selected for each region in order to show the  
30 spatial variability of the AOT at 532 nm retrieved along the CALIOP transect. The first case is related to a biomass-burning  
event detected off the coast of Namibia on 13 August 2006. The second event concerns Saharan dust lifted above clouds



westwards over the North Atlantic Ocean on 4 August 2008, and the third case concerns Siberian biomass-burning aerosols transported over the Okhotsk Sea, on 3 July 2008.

Figure 1 presents the backscatter profile at 532 nm ( $\text{km}^{-1} \text{sr}^{-1}$ ) of the lidar CALIOP for the three case studies, which directly provides information on the aerosol and cloud vertical distribution. In addition, the AOT and AE values measured by different techniques are presented along the CALIOP track. Additional results for the study cases comparison are shown in Table 1.

Figure 2 shows the regional comparison between the AOT and AE retrieved with POLDER and  $\text{DRM}_{\text{SODA}}$  for a period of six months in 2008. The retrieval of aerosol type becomes difficult at small AOT. Therefore the AE comparison was performed only when the values of POLDER AOT at 865nm and  $\text{DRM}_{\text{SODA}}$  AOT at 532 nm were larger than 0.1. The AE mean value is shown with a dashed blue line. The lateral histograms show the data distribution. For the AOT comparison the color scale represents the POLDER  $\text{AE}_{670/865}$ . In the case of AE comparison, the POLDER  $\text{AOT}_{532\text{nm}}$  was also reported with a color scale. The above-mentioned description is also considered in Fig. 3, which presents the regional comparison between the AOT and AE retrieved with POLDER and  $\text{CALIOP}_{\text{OM}}$  for the same period. Additional results for the regional inter-comparison are reported in Table 2.

### 3.1 African biomass-burning aerosols

According to the CALIOP vertical profile of the biomass-burning case (Fig. 1a), the cloud top is at around 1.5 km and the aerosol layer is located between 3 and 5 km. We observe a thin cirrus cloud between 10° and 12° S that was not filtered, probably since the cirrus is optically too thin (Fig. 1b and 1c). In general, there is an excellent agreement between POLDER,  $\text{DRM}_{\text{Hu}}$  and  $\text{DRM}_{\text{SODA}}$  AOT retrievals with a square correlation  $R^2 = 0.93$  (see Table 1). High values of AOT are retrieved by the different methods, with AOT values as large as 1.5. The retrieved POLDER  $\text{AE}_{670/865}$  is larger than 1.8 (Fig. 1c), which is characteristic for fine mode particles. The  $\text{DRM}_{\text{SODA}}$   $\text{AE}_{532/1064}$  is consistent with the POLDER AE, with values higher than 1.5. AOT values retrieved by  $\text{CALIOP}_{\text{OM}}$  are much lower than the ones retrieved by the three other techniques. The maximal AOT retrieved by  $\text{CALIOP}_{\text{OM}}$  at 532 nm is 0.5. A possible explanation for this potential low bias was proposed by Jethva et al. (2014): in case of optically thick aerosol layer, the sensitivity of the backscattered signal to the altitude of the base of the aerosol layer would be reduced or lost, being strongly attenuated by the two-way transmission term. As a result, the operational algorithm may overestimate the aerosol base altitude and so underestimate the geometrical thickness of the aerosol layer and consequently the AOT. The  $\text{CALIOP}_{\text{OM}}$  mean  $\text{AE}_{532/1064}$  seems quite low for fine mode particles (AE values are lower than 1). The selection of an inappropriate aerosol model might also contribute to the underestimation of the AOT for this case study.

Regional analysis shows that South Atlantic region is mostly characterized by biomass-burning aerosols with large AOT and AE (Fig. 2a and 2b). On average, the cloud top height is located below 1.5 km, while the aerosol layers are frequently located between 2.5 and 4 km (see Table 3). The  $\text{AOT}_{532\text{nm}}$  measured by  $\text{DRM}_{\text{SODA}}$  and POLDER may reach values as large as 1.30 (Fig. 2a), with 80 % of the retrieved AOTs ranging between 0.05 and 0.8. This AOT inter-comparison



shows close correlation between  $DRM_{SODA}$  and POLDER ( $R^2 = 0.83$ ). The mean value of POLDER  $AE_{670/865}$  is 2.05, whereas the mean  $DRM_{SODA} AE_{532/1064}$  is 1.79 (Table 5) (both typical for BBA).  $DRM_{Hu}$  and  $DRM_{SODA}$  give rather same results. From the linear regressions performed (see Table 2) we can observe that the offset is always positive for  $DRM_{Hu}$  and systematically larger than the absolute value for  $DRM_{SODA}$ , when compared to POLDER method. The AOT estimated by POLDER is constantly between  $DRM_{Hu}$  and  $DRM_{SODA}$ .

We do not find a good correlation between the  $CALIOP_{OM}$  and POLDER AOT and AE retrievals. The  $CALIOP_{OM}$  mean  $AOT_{532nm}$  is 0.12 and the mean  $AE_{532/1064}$  is 0.97. Comparing with POLDER and  $DRM_{SODA}$ ,  $CALIOP_{OM}$  is underestimating the ACAOT by a factor of 2.92.

### 3.2 Saharan desert dust aerosols

For the mineral dust case (Fig. 1d), the cloud top altitude is located at approximately 1 km altitude whereas the aerosol layer is located between 2 and 5 km for latitudes between  $18^\circ$  and  $23^\circ$  N. Figure 1e shows that the POLDER,  $DRM_{SODA}$  and  $DRM_{Hu}$   $AOT_{532nm}$  increase up to 0.92, following the same gradient. The correlation coefficients between POLDER parameters and  $DRM_{Hu}$  and  $DRM_{SODA}$  parameters are close (Table 1). The majority of POLDER  $AE_{670/865}$  and  $DRM_{SODA} AE_{532/1064}$  are associated with values lower than 0.4 (Fig. 1f), which indicates that coarse mode particles are predominant. **Except for few retrievals associated with an abrupt change in the AE and AOT measured by  $CALIOP_{OM}$  (around  $21^\circ$  N in latitude), 90 % of the  $CALIOP_{OM} AOT_{532nm}$  is lower than 0.45, being once again underestimated with respect to the other estimates. Most of  $CALIOP_{OM} AE_{532/1064}$  values are underestimated (i.e. overestimation of the particles size) in comparison with the AE retrieved by the two other algorithms.**

A regional study shows similar AOT and AE results over the North Atlantic region (Fig. 2c). On average, the aerosol layers are located between 3 and 4.5 km and the cloud top heights are typically around 1.4 km (see Table 3). The values of  $AOT_{532}$  retrieved from POLDER and  $DRM_{SODA}$  are well correlated ( $R^2 = 0.82$ ), with maximum values of respectively 1.19 and 0.95. Nonetheless, we observe a larger offset between  $DRM_{SODA}$  and POLDER  $AOT_{532nm}$  for this region (-0.09) compared to the South Atlantic Ocean region (-0.03). The use of only one dust model in the LUT algorithm used for POLDER remains a limitation that might explain this larger offset. The introduction of additional dust models with larger or smaller effective radius values may contribute to improve the AOT retrievals for dust ACC events. Regarding the POLDER  $AE_{670/865}$  retrievals, most of the values are lower than 0.4, **which is expected for desert dust aerosols** (Fig. 2c and 2d). However, for AOT values lower than 0.2, the  $AE_{670/865}$  retrieved by POLDER is between 1.4 and 2.2. This is explained by the fact that the selection of the dust model is not permitted in the POLDER algorithm in case of low AOTs. Nonetheless, all three methods are consistent in revealing the predominance of the coarse mode. The mean values for the AE are 0.49 for POLDER, 0.10 for  $DRM_{SODA}$  and -0.19 for  $CALIOP_{OM}$ . The  $AOT_{532nm}$  correlation between  $CALIOP_{OM}$  and POLDER is low, with  $R^2 = 0.42$ .



### 3.3 East Asian mixture of aerosols

The CALIOP transect shows that Siberian biomass-burning case is located between 40° and 52° N, the cloud top altitude is constantly around 1 km, and the base of the aerosol layer decreases from 10 km in the south (at 45° N) to around 2 km in the north (at 54° N) (Fig. 1g). We notice also cirrus clouds at high altitude (around 10 km) between 47° and 51° N, which were efficiently eliminated from the retrievals (Fig. 1h). The maximum POLDER AOT value is as large as 1.9, while DRM reaches 1.3 in AOT. Nonetheless, Table 1 shows that POLDER and DRM methods AOT<sub>532nm</sub> retrievals are consistent ( $R^2 = 0.90$ ). POLDER AE<sub>670/865</sub> values are between 1.7 and 2.3, indicating small particles of smoke, while DRM<sub>SODA</sub> AE<sub>532/1064</sub> has a large range of values (Fig. 1i). The number of sampled ACAOT events by CALIOP<sub>OM</sub> is 4.5 times less than of POLDER and DRM<sub>SODA</sub>. For these, the CALIOP<sub>OM</sub> AOTs are underestimated by a factor of 1.5 compared to ones retrieved by the other methods. Also the correlation coefficient with POLDER is 0.45.

On a regional scale, this area is under the influence of various aerosols (BBA, DDA, pollution) and elevated cirrus clouds are frequent. The mean cloud top altitude is around 1 km and the aerosols are between 2.5 km and 4.0 km. As indicated in Table 3, the maximum aerosol altitude is 9.85 km, which might suggest cirrus misclassification. In some cases, DRM<sub>SODA</sub> gives large values of AOT<sub>532nm</sub> (larger than 1) whereas the POLDER estimates AOT<sub>532nm</sub> smaller than 0.2. These situations could be explained by a misinterpretation of thin cirrus clouds as aerosols. Otherwise, the POLDER mean AOT<sub>532nm</sub> and DRM<sub>SODA</sub> AOT<sub>532nm</sub> are in rather close agreement (0.18 and 0.15, respectively, see Table 4), but the correlation between them is low ( $R^2 = 0.37$ , Table 2). All methods show a large variability for the retrieved AE, with values that correspond to particle size distributions dominated by coarse or fine modes and mixtures (Table 5). As previously mentioned, the algorithm developed for POLDER uses a bimodal aerosol model for dust. However, the possibility of mixing different fine and coarse aerosol models in various proportions is not yet included. This might explain why we found a lower correlation between the POLDER and DRM retrievals for this region. As for above, the CALIOP<sub>OM</sub> and POLDER AOT<sub>532nm</sub> are not correlated ( $R^2 = 0.24$ ).

In general, there is a good agreement between POLDER and DRM<sub>SODA</sub> AOTs, especially when the fine mode or coarse mode dominates the particle size distribution (i.e. BBA and DDA). Overall, DRM<sub>SODA</sub> and DRM<sub>Hu</sub> give similar results. However, the AOTs retrieved with DRM<sub>Hu</sub> are generally larger than those of DRM<sub>SODA</sub> for all the three regions (i.e. 0.37 compared to 0.28 for SAO, see Table 4). While DRM<sub>SODA</sub> has a constant negative offset when compared to POLDER, DRM<sub>Hu</sub> rarely retrieves null AOT values (offsets always larger than 0, see Table 2). This is likely to be a consequence of the calibration performed for the DRM<sub>SODA</sub> method. Also, there is no obvious correlation between the CALIOP<sub>OM</sub> and POLDER AOT<sub>532nm</sub> retrievals for all regions.

Finally, in addition to the six months regional study, we also examined the impact of the vertical aerosol-cloud profiles over the three regions using data acquired from May to October between 2006 and 2010. We systematically found higher correlation coefficients between the DRM<sub>SODA</sub> and POLDER AOTs when the layers were well separated than when



they were in contact (see Table 6). These results have led us to consider the vertical distribution of aerosols and clouds in the global comparison.

#### 4 Global analysis on different types of scenes

##### 4.1 Detached, attached, undetermined



5 Figure 4a shows the global comparison between the  $AOT_{532nm}$  and AE retrieved with POLDER and  $DRM_{SODA}$  for the *detached* cases. The AE comparison was only performed when the POLDER AOT at 865 nm and  $DRM_{SODA}$  AOT at 532 nm are larger than 0.1. The color scales used in Fig. 4 represent either the POLDER  $AE_{670/865}$  for the AOT comparison (Fig 4a) or the POLDER  $AOT_{532nm}$  for the AE comparison (Fig. 4d). Considering the large amount of selected data (85.6 % of the two-layer cases) in terms of both spatial and temporal coverage, the comparison shows a good correlation between the two  
10 methods ( $R^2 = 0.68$ ). A better agreement between the methods is found when the values of  $DRM_{SODA}$  and POLDER AE are larger than 1.8. This is likely due to the fact that the POLDER method is more sensitive to fine mode aerosols, due to polarization measurements, and also because an improved description of the fine mode properties was included in the LUT (i.e. six fine mode aerosol models are used).

Events for which the aerosol layer is *attached* to the cloud top represent 14.4 % of the total number of two-layer  
15 cases. They are associated with lower AOT and the correlation between the two retrievals largely decreases (compare to the detached events). The correlation between the two AOT retrievals also decreases ( $R^2 = 0.36$ , Fig. 4c). The POLDER AOT is larger by a coefficient of 1.7 than the  $DRM_{SODA}$  AOT on average. The AE given by both methods is approximately 1.0 (when considering only AE values associated with  $AOT > 0.1$ ). The lateral histogram shows that the POLDER method identifies AAC events associated with both low and high AE values resulting in a mean AE of about 1.0.

20 The *undetermined* situations correspond to retrievals when CALIOP does not give all the information regarding the layer altitudes. The number of cases is significant (approximately 92 % of the total number of global retrievals) but most of data (95 %) corresponds to  $AOT_{532nm}$  lower than 0.2. This probably explains why the layer detection algorithm has difficulties in estimating the base and top of the aerosol layer. For the *undetermined* cases, we observe that there is not such  
correlation between POLDER and  $DRM_{SODA}$  measurements. On average, the  $DRM_{SODA}$  AOTs are centred around zero for  
25 this category whereas POLDER has a non-negligible low AOT for most cases. In this category, the AE comparison shows a better consistency between the methods for  $AOT_{532nm} > 0.5$  and for AE of approximately 2.0.

Table 7 shows the results of the linear regressions performed between the AOTs retrieved with POLDER and the other active method considered in our study for each category (i.e. detached, attached and undetermined). We recapture the systematically larger offsets of  $DRM_{Hu}$   $AOT_{532nm}$  compared to  $DRM_{SODA}$ , and the underestimation of  $CALIOP_{OM}$  AOT with  
30 respect to the other methods.



## 4.2 Evolution of the above cloud AOT retrievals with cloud properties

In principle, the retrieval of AAC properties from the methods considered in this study should not depend on the properties of the underlying clouds. However, hypotheses and empirical relations used in the retrieval methods to exploit the signal backscattered by the underlying cloud cover have obviously  limitations. In order to understand potential issues linked with diversity of cloud properties, we analyse in this section the difference between the AOT retrievals of POLDER, DRM<sub>SODA</sub> and DRM<sub>Hu</sub> by classes of cloud properties (COT and  $r_{eff}$  retrieved with MODIS). We considered global measurements acquired for 4.5 years of data and used the classification defined in Sect. 2.3.2.

Figure 5 presents POLDER and DRM<sub>SODA</sub> AOT<sub>532</sub> retrievals as a function of the MODIS droplets effective radius ( $r_{eff}$ ), while Fig. 7 displays POLDER and DRM<sub>SODA</sub> AOT<sub>532nm</sub> as a function of the MODIS cloud optical thickness (COT). Histograms of the cloud properties are also reported in Fig. 5, 6 and 7. The results of the POLDER and DRM<sub>Hu</sub> AOT<sub>532</sub> comparison as a function of the effective radius are shown in Fig. 6. DRM<sub>SODA</sub> and DRM<sub>HU</sub> AOT<sub>532nm</sub> generally exhibit rather similar behaviour, at least qualitatively. Therefore, we did not report the results found for the DRM<sub>Hu</sub> AOT<sub>532</sub> as a function of MODIS COT.

### 4.2.1 AOT versus $r_{eff}$

The lateral histograms plotted in Fig. 5 and Fig. 7 show that most of the AAC scenes correspond to cloud droplets effective radius values between 8 and 15  $\mu\text{m}$  (mean  $r_{eff}$  equal to 12  $\mu\text{m}$ ) and COT ranging from 5 to 15 (mean COT of 10). These mean values are expected since most of the of AAC events are generally associated with low-level non-precipitating clouds, such as stratocumulus ones, which typically show rather small droplets (approximately 10  $\mu\text{m}$ ) and optical thickness values of approximately 10.

Figure 5a shows the POLDER and DRM<sub>SODA</sub> AOTs for the *detached* situations. For the two methods, the retrieved AOTs are maximal for the smallest values of  $r_{eff}$  and progressively decrease with  $r_{eff}$ . Same tendencies are observed for the DRM<sub>Hu</sub> (see Fig. 6a). The two curves have however an offset. The histogram of the differences between POLDER and DRM<sub>SODA</sub> AOT ( $\Delta\tau$ ) is presented in Fig. 5d. The mean  $\Delta\tau$  value computed over the entire range of  $r_{eff}$  is equal to 0.073. This offset is not constant and slightly increases with  $r_{eff}$ , suggesting a sensitivity of one of the two methods to the cloud droplets effective radius. The DRM algorithm assumes a constant lidar ratio of 19 sr, independent of the cloud droplet effective radius. In order to evaluate the accuracy of this approximation, we recalculated the DRM<sub>SODA</sub> AOT<sub>532nm</sub> taking into account the dependence of  $S_c$  on  $r_{eff}$ . As defined in Josset et al. (2011),  $S_c$  was computed using a Mie code with the following equation:

$$S_c = \frac{4\pi}{\omega_0 \times p(180^\circ)}, \quad (11)$$

where  $p(180^\circ)$  is the average value of the phase function in the backscatter direction computed over the size distribution.  $\omega_0$  is the Single Scattering Albedo of the particles, defined as the ratio between the mean scattering coefficient and the mean



extinction coefficient computed over the particle size distribution. We used a two-parameter gamma size distribution with an effective variance of 0.088. The real refractive index was set to 1.337. Liquid water droplets do not significantly absorb at 532 nm and the imaginary part of the complex refractive index was set to 0. As shown in Figure 8,  $S_c$  slightly decreases with  $r_{eff}$  from 19.5 to 15.5 as the effective radius values increases from 5 to 40  $\mu\text{m}$ . With this correction, the mean difference between POLDER and the  $\text{DRM}_{\text{SODA}}$   $\text{AOT}_{532\text{nm}}$  ( $\Delta\tau$  corr  $S_c$  in Fig. 5d) decreases from 0.073 to 0.065. We found equivalent results for the *attached* and *undetermined* cases (Fig. 5b and 5c). After correction of  $S_c$ , the difference between POLDER and  $\text{DRM}_{\text{SODA}}$  decreases on average by 0.01, for the *attached* cases, and by 0.019 for the *undetermined* cases. We also observe that most of the negative AOT values retrieved by the  $\text{DRM}_{\text{SODA}}$  shift either to null values or weakly positive values when this correction is included (Fig 5a, 5b and 5c). We are aware that MODIS effective radius may be affected by the presence of aerosols above clouds. For example, biases of  $\pm 2 \mu\text{m}$  for  $r_{eff}$  were found by Haywood et al. (2004) for strong dust events above clouds. However, we consider that the impact of the biases on the retrieved  $r_{eff}$  on our findings and conclusions can be neglected, since the analysis hold for (i) a wide range of droplets effective radius (from 5 to 40  $\mu\text{m}$ ) and (ii) AAC events associated with low aerosol loadings (see the results for the *undetermined* cases), where the impacts of the aerosols on the cloud retrievals are expected to be minimized or negligible.

#### 4.2.2 AOT versus COT

The two methods were developed to detect AAC events in the case of optically thick and homogeneous liquid water clouds. In the following, we only discuss results obtained for large values of COT (larger than 5). If the clouds are optically thinner, the two methods are potentially less accurate since they become sensitive to the surface contribution. For COTs ranging between 5 and 30 and for *detached* cases, the POLDER AOTs are almost constant and reach 0.3 on average at 532 nm (see Fig. 7a). Most of the associated COT values are then ranging between 5 and 10. For these cases,  $\text{DRM}_{\text{SODA}}$  and POLDER AOTs are offset by around 0.07 on average, as noted above. However, the  $\text{DRM}_{\text{SODA}}$  AOT progressively increases with the COT, which is not observed for the POLDER AOT. Consequently, the differences in AOT between the two methods become almost negligible for the largest (and less frequent) values of COT (larger than 20). For COTs larger than 3, the polarized signal reflected by the cloud is saturated and the POLDER method should be insensitive to COT.  $\text{DRM}_{\text{SODA}}$  is sensitive to the multiple scattering processes occurring within the cloud layers and might be impacted by the COT since multiple scattering increases with the optical thickness. The measured depolarization ( $\delta'$ ) and the multiple scattering factor ( $\eta_{\text{calibr}}$ ) plotted as a function of the COT are shown in Fig. 7d. As expected, the depolarization and the multiple scattering factor respectively increase and decrease as COT increases. The increase in the  $\text{DRM}_{\text{SODA}}$  AOT observed at large COTs might be due to an increase in the multiple scattering. We recall that  $\text{DRM}_{\text{SODA}}$  uses a relationship to connect the depolarization and the multiple scattering factor and that this relation is calibrated based on CALIOP data. The calibration might be less accurate in case of AAC events associated with clouds for which the properties are statistically less representative. Again, we presume that our conclusions are not impacted by the fact that the MODIS COTs can be



potentially biased in case of AAC events since the tendencies we observed hold for a large range of variability in COT (5 to 30) and also for AAC events associated with low AOT above clouds (see the results for the *undetermined* cases).

## 5 Discussion

In the first part of this section, we quantify and discuss the overall differences found between the active and passive methods in terms of the retrieved AOT. In the second part, we address more specifically the *attached* cases and make hypotheses regarding the meaning of these results.

On average, the difference between POLDER and DRM<sub>SODA</sub> AOTs at 532 nm is equal to 0.073 for the *detached* cases and 0.087 for the *undetermined* cases. These differences slightly decrease to 0.065 and 0.068, respectively, when we account for the dependency of the cloud droplets lidar ratio ( $S_c$ ) to  $r_{eff}$  in Eq. 10. The POLDER AOTs are systematically smaller than the ones retrieved with DRM<sub>Hu</sub>. On average, these differences between these two methods are equal to -0.039 and -0.057, for the *detached* cases, and reach -0.036 and -0.048 for the *undetermined* cases, respectively without and with corrections for  $S_c$ . Thereby, the POLDER AOT estimates range, on average, between the DRM<sub>Hu</sub> and DRM<sub>SODA</sub> ones. The differences in AOTs found between the POLDER method and the two DRM ones could be set to zero by modifying the lidar calibration by roughly  $\pm 10\%$ . One another main difference between the three methods is their different responses in terms of AOT when the atmosphere above the clouds becomes pristine. The majority of AOT (94 %) is lower than 0.1 at 865 nm for the *undetermined* cases. For these cases, the POLDER algorithm retrieves a mean AOT of about 0.04 at 865 nm. The accuracy of the POLDER AOT product is in the same order of magnitude. For an AOT<sub>865nm</sub> of 0.2, the error for a real refractive index uncertainty of  $\pm 0.06$  would be about 0.05; for an imaginary refractive index uncertainty of  $\pm 0.01$ , the error would be of 0.02 (Peers et al., 2015). The impact of the assumed refractive index is lower at smaller AOT (especially for an AOT of 0.04). The background reaches 0.09 in AOT at 532 nm. This latter value is only reported for the sake of comparison with the two other methods since the Ångström exponent retrieved by POLDER, (and consequently the AOT extrapolated at 532 nm) cannot be accurately retrieved for low AOTs. DRM<sub>SODA</sub> found a mean AOT of about 0.005 at 532 nm for the *undetermined* cases (see Fig. 5c). The result is likely due to the re-calibration process since DRM<sub>Hu</sub> found a background even larger than the POLDER one, of about 0.12 at 532 nm. It is difficult to assess the truthfulness of this background, considering the given level of accuracy of the POLDER method and the uncertainties associated with the lidar calibration. We assume that these background values are not physical and could be due to some inherent limitations of the retrieval methods. From our data, however, we cannot exclude the possibility that there is always a background loading of particles above clouds (e.g. aerosols or fine droplets in formation). Nevertheless, the main result of our investigation is that POLDER and DRM methods compare well for most situations with a mean difference of about  $\pm 0.07$  in AOT at 532 nm.

Although the number of cases is small, the results of the *attached* cases are interesting. They suggest that the lidar CALIOP and POLDER could be affected by layers of aerosols that physically and locally interact with the upper part of the cloud. In order to understand how the vertical profiles differ from one situation to another, we compared the CALIOP



attenuated backscatter coefficient for *attached* and *detached* cases. We considered the period 2006–2010 and used data acquired over the entire globe. We only select the *attached* and *detached* cases where the cloud top altitude is below 1.5 km and the  $\text{DRM}_{\text{SODA}} \text{AOT}_{532\text{nm}}$  is larger than 0.1. These criteria allow for selection of data that corresponds to AAC events associated with similar cloud vertical extents and with significant AOTs. For these cases, we computed the average and median of the CALIOP level 1 attenuated backscatter coefficients at 532 nm. Figure 9 presents these results and some information concerning the mean and median values of CALIOP level 2 products: cloud top altitude, aerosol layer's base and top altitudes. The mean and median values computed for the AOTs retrieved by POLDER and  $\text{DRM}_{\text{SODA}}$  and the numbers of sampled events are also reported. Two different types of profiles can be observed for the *detached* and *attached* situations. For the *detached* cases, the aerosol and cloud backscattering profiles can be easily distinguished in both the median and mean profiles. The strong peaks in the backscatter profiles at around 1 km correspond to the top of the clouds, whereas the increase in the lidar backscatter signal observed between 2 and 4 km in altitude comes from the aerosols. For the *attached* situations, the backscatter profiles are noisier, which is likely due to the fact that the number of detected events is smaller compared to *detached* cases. The top of the cloud layer is still clearly visible in the mean and median backscattered lidar signals, but two maxima can be observed. We assume that  sampled two different regimes of clouds. In addition, there is a continuous transition in the backscatter signal between the top of the cloud and the above molecular atmosphere that is most clearly visible in the median profiles. This signal doesn't appear for the *detached* cases. This signal could explain the non-negligible above-cloud AOTs retrieved by POLDER and  $\text{DRM}_{\text{SODA}}$  for the *attached* cases (see Fig. 4). It is difficult to assess the origin of this signal. This might be due to aerosols layers that penetrate the cloud layers at the top of the clouds. Natural aerosol or fine droplets in formation, commonly present in the vicinity of the clouds, might also create this additional signal. 

Another hypothesis that could explain the low AOT correlation for the *attached* cases is that the aerosols located within the cloud layer could affect the polarized radiances measured by POLDER. Since the operational algorithm developed for POLDER assumes that the entire aerosol layer is located above the clouds, an additional polarized signal coming from aerosol located within the cloud would lead to an overestimation of the above cloud AOT retrieved from POLDER. To test this assumption, we modelled the polarized radiance measured by POLDER for AAC scenes, considering different vertical locations of the aerosol layer (Fig. 10). We used the Successive orders of scattering (SOS) radiative transfer code (Lenoble et al., 2007) for this simulation. We considered a liquid water cloud located between 0 and 1 km. The particles (aerosol and cloud) are vertically homogeneously mixed. The COT is equal to 10 and the effective radius and variance are equal to 10  $\mu\text{m}$  and 0.08, respectively. The aerosol layer is characterized by an AOT of 0.25 at 865 nm, a refractive index of  $m = 1.47 - 0.01i$  and an effective radius of 0.15  $\mu\text{m}$ . Fig. 9 shows the typical polarized feature for AAC events in case of *detached* situations (i.e. aerosols located between 1.25 and 1.75 km): a creation of polarization is observed at side and forward scattering angles, whereas the cloud bow magnitude decreases. For the *attached* case (aerosols between 0.75 and 1.25 km), the amount of polarization created at forward scattering angles decreases and the cloud bow attenuation is less significant in comparison with the *detached* scenario. When the aerosol layer is located within the upper part of the cloud layer (between



0.5 and 1 km), we still observe a weak polarized signal created at forward scattering angles. When the aerosol layer is located in the lower part of the cloud layer, the effects of the aerosols disappear since the polarized radiance scattered by the aerosols is lost due to multiple scattering occurring within the clouds. These simulations were processed with the POLDER algorithm (Waquet et al., 2013b). We recall that the LUTs used in this algorithm were built for *detached* situations. The algorithm retrieved an AOT of 0.09 at 865 nm when the aerosols are located within the upper part of the cloud layer. This demonstrates that polarized radiances are sensitive to aerosols situated *within* the clouds for the *attached* cases.

The DRM methods might also be impacted by the presence of aerosols within the clouds. Aerosols is solution within the cloud droplets (i.e. internal mixture) might impact the chemical composition of the droplets and modify their ability to backscatter light. Fig. 8 shows lidar ratio computed for absorbing cloud droplets. We used an imaginary part of 0.0001 for the complex refractive index of the droplets. This might simulate, for instance, the properties of brown clouds contaminated by absorbing aerosols. We observe a drastic increase of  $S_c$  with  $r_{eff}$  (from 21.7 sr at 5  $\mu\text{m}$  to 50 sr at 40  $\mu\text{m}$ ) when the water droplet is weakly absorbing. In the case of an external mixture, we assume that the presence of aerosols at the top of cloud might also modify the value of  $S_c$ . Any deviation from the 19 sr value assumed for the droplets lidar ratio in Eq. (10), will necessarily impact the retrieved AOT and the differences observed between the AOT estimates provided by the POLDER and DRM methods.

## 6 Conclusions

In this study, we compared and analysed the consistency of the AOT and AE retrievals above clouds from different passive and active techniques. We used the CALIOP operational algorithm (Winker et al., 2009) the POLDER polarisation method (Waquet et al., 2013b), and the CALIOP-based depolarisation ratio method ( $\text{DRM}_{\text{Hu}}$ ) (Hu et al., 2007a) – for which we proposed a re-calibrated version of the DRM algorithm ( $\text{DRM}_{\text{SOD}\lambda}$ ). The observations were made for: a) three case studies corresponding to an African biomass-burning event, a Saharan dust event and a Siberian biomass-burning event; b) a regional scale analysis, over South Atlantic Ocean, North Atlantic Ocean and North Pacific Ocean for a period of six months in 2008 and c) a global scale analysis for different vertical layer distributions for the period 2006–2010.

In the regional analyse, we observed that POLDER method and DRM are in good agreement when the microphysics of aerosols is dominated by fine-mode particles of biomass-burning aerosols (in the South Atlantic region,  $R^2 = 0.83$ ) or coarse-mode aerosols of dust (in the North Atlantic region,  $R^2 = 0.82$ ). A good correlation between these methods ( $R^2 = 0.68$ ) is also noticed in the global treatment, when the aerosol and cloud layers are well separated. The CALIOP operational method largely underestimates the AOT above clouds in all situations, with respect with the other methods.

The differences between the DRM and POLDER retrievals increase when a complex mixture of aerosols is expected (such as in the East Asia region). This is probably due to the fact that the current algorithm developed for POLDER uses a limited number of microphysical models of aerosols. Also, the relative position of the aerosol layer above the cloud impact the AOT retrievals from both active and passive measurements: the correlation decreases when the layers are in



contact ( $R^2 = 0.36$ ), suggesting that aerosols at the top or within the cloud can affect the AOT retrievals. One hypothesis is that an additional polarized signal coming from aerosol located *within* the cloud could affect the polarization signal and method, which leads to an overestimation of the AOT retrieved with POLDER algorithm. The aerosols *attached with* or *within* the cloud also have the potential to impact the DRM retrievals, by modifying the lidar ratio (and consequently the  
5 AOT) as a result of internal or external mixture.

Furthermore, we investigated potential biases in the retrieved AOT measured by POLDER and  $DRM_{SODA}$  as a function of MODIS cloud properties (i.e. droplet effective radius ( $r_{eff}$ ) and cloud optical thickness (COT)). The tendencies show an increase in the difference between the two methods for larger  $r_{eff}$ , suggesting sensitivity to the cloud droplet effective radius. For this reason, we recalculated the  $DRM_{SODA}$   $AOT_{532nm}$  taking into account the dependence of lidar ratio on  
10  $r_{eff}$ , as this method assumes a constant lidar ratio regarding the droplet effective radius. By doing so, we observed a decrease in the difference between POLDER and DRM methods and a shift of the DRM AOT values from negative to positive. For a better accuracy of DRM retrievals in future studies, this correction should be taken into account. The results show also that the multiple scattering processes, which are more pronounced in optically thick clouds, could also affect the DRM technique.

All of the aforementioned situations have revealed that  $DRM_{Hu}$  has larger mean AOT than that of  $DRM_{SODA}$ . This is  
15 likely to be a consequence of the re-calibration performed for the  $DRM_{SODA}$  method. Actually, POLDER  $AOT_{532nm}$  values are consistently smaller than the ones of  $DRM_{Hu}$  and larger than those of  $DRM_{SODA}$ . The primary conclusion of our investigation is that POLDER and DRM techniques are comparable for the majority of cases, with a mean difference of about  $\pm 0.07$  in AOT at 532 nm, depending on lidar calibration.

Given the fact that each method relies upon different physical concepts, applied to different sensors and  
20 measurements, the high value of the correlation obtained for the AOT retrievals is a remarkable result that highlights the coherence between active and passive methods for aerosols above clouds. Nonetheless, more efforts have to be done to increase the accuracy of the methods, in order to better understand aerosols above clouds and their related effects. A first perspective is to improve the POLDER algorithm by introducing additional dust or mixture models with larger or smaller effective radii values in the LUT. This would definitely improve the AOT and AE retrievals in more complex situations  
25 (such as East Asia region). Also, our results suggest that a combination of POLDER and DRM methods has the potential to detect aerosols *within* clouds. It is very relevant to study these situations, since they can affect the retrievals and provide important information regarding the cloud processes. A second perspective would be to exploit the synergy between CALIOP and POLDER to infer the direct aerosol radiative forcing, aerosol heating rates and the semi-direct effect of absorbing aerosols located above clouds.

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## References

- Balkanski, Y., Schulz, M., Claquin, T. and Guibert, S.: Reevaluation of Mineral aerosol radiative forcings suggests a better agreement with satellite and AERONET data, *Atmos. Chem. Phys.*, 7(1), 81–95, doi:10.5194/acp-7-81-2007, 2007.
- Bates, D. R.: Rayleigh scattering by air., 1984.
- 5 Cao, X., Roy, G., Roy, N. and Bernier, R.: Comparison of the relationships between lidar integrated backscattered light and accumulated depolarization ratios for linear and circular polarization for water droplets, fog oil, and dust, *Appl. Opt.*, 48(21), 4130–4141, doi:10.1364/AO.48.004130, 2009.
- Chand, D., Anderson, T. L., Wood, R., Charlson, R. J., Hu, Y., Liu, Z. and Vaughan, M.: Quantifying above-cloud aerosol using spaceborne lidar for improved understanding of cloudy-sky direct climate forcing, *J. Geophys. Res.*, 113(D13), D13206, doi:10.1029/2007JD009433, 2008.
- 10 Chand, D., Wood, R., Anderson, T. L., Satheesh, S. K. and Charlson, R. J.: Satellite-derived direct radiative effect of aerosols dependent on cloud cover, *Nat. Geosci.*, 2(3), 181–184, doi:10.1038/ngeo437, 2009.
- Costantino, L. and Bréon, F.-M.: Aerosol indirect effect on warm clouds over South-East Atlantic, from co-located MODIS and CALIPSO observations, *Atmos. Chem. Phys. Discuss.*, 12(6), 14197–14246, doi:10.5194/acpd-12-14197-2012, 2012.
- 15 Deuzé, J. L., Herman, M. and Santer, R.: Fourier series expansion of the transfer equation in the atmosphere–ocean system, *J. Quant. Spectrosc. Radiat. Transf.*, 41(6), 483–494, doi:10.1016/0022-4073(89)90118-0, 1989.
- Dubovik, O., Holben, B., Eck, T. F., Smirnov, A., Kaufman, Y. J., King, M. D., Tanré, D., Slutsker, I., Sciences, G. E. and Directorate, E. S.: Variability of Absorption and Optical Properties of Key Aerosol Types Observed in Worldwide Locations, *J. Atmos. Sci.*, 59(3), 590–608, doi:10.1175/1520-0469(2002)059<0590:VOAAOP>2.0.CO;2, 2002.
- 20 De Graaf, M., Bellouin, N., Tilstra, L. G., Haywood, J. and Stammes, P.: Aerosol direct radiative effect of smoke over clouds over the southeast Atlantic Ocean from 2006 to 2009, *Geophys. Res. Lett.*, 41(21), 7723–7730, doi:10.1002/2014GL061103, 2014.
- Hansen, J., Sato, M. and Ruedy, R.: Radiative forcing and climate response, *J. Geophys. Res. Atmos.*, 102(D6), 6831–6864, doi:10.1029/96JD03436, 1997.
- 25 Hasekamp, O. P.: Capability of multi-viewing-angle photo-polarimetric measurements for the simultaneous retrieval of aerosol and cloud properties, *Atmos. Meas. Tech.*, 3(4), 839–851, doi:10.5194/amt-3-839-2010, 2010.
- Haywood, J. M., Osborne, S. R. and Abel, S. J.: The effect of overlying absorbing aerosol layers on remote sensing retrievals of cloud effective radius and cloud optical depth, *Q. J. R. Meteorol. Soc.*, 130(598), 779–800, doi:10.1256/qj.03.100, 2004.
- Herman, M., Deuzé, J. L., Marchand, A., Roger, B. and Lallart, P.: Aerosol remote sensing from POLDER/ADEOS over the ocean: Improved retrieval using a nonspherical particle model, *J. Geophys. Res. D Atmos.*, 110(10), 1–11, doi:10.1029/2004JD004798, 2005.
- 30 Hu, Y.: Depolarization ratio-effective lidar ratio relation: Theoretical basis for space lidar cloud phase discrimination, *Geophys. Res. Lett.*, 34(11), L11812, doi:10.1029/2007GL029584, 2007.



- Hu, Y., Liu, Z., Winker, D., Vaughan, M., Noel, V., Bissonnette, L., Roy, G. and McGill, M.: Simple relation between lidar multiple scattering and depolarization for water clouds, *Opt. Lett.*, 31(12), 1809, doi:10.1364/OL.31.001809, 2006.
- Hu, Y., Vaughan, M., Liu, Z., Powell, K. and Rodier, S.: Retrieving Optical Depths and Lidar Ratios for Transparent Layers Above Opaque Water Clouds From CALIPSO Lidar Measurements, *IEEE Geosci. Remote Sens. Lett.*, 4(4), 523–526, doi:10.1109/LGRS.2007.901085, 2007a.
- 5 Hu, Y., Vaughan, M., Liu, Z., Lin, B., Yang, P., Flittner, D., Hunt, B., Kuehn, R., Huang, J., Wu, D., Rodier, S., Powell, K., Trepte, C. and Winker, D.: The depolarization - attenuated backscatter relation: CALIPSO lidar measurements vs. theory, *Opt. Express*, 15(9), 5327–5332, doi:10.1364/OE.15.005327, 2007b.
- Jethva, H., Torres, O., Remer, L. A. and Bhartia, P. K.: A color ratio method for simultaneous retrieval of aerosol and cloud optical thickness of above-cloud absorbing aerosols from passive sensors: Application to MODIS measurements, *IEEE Trans. Geosci. Remote Sens.*, 51(7), 3862–3870, doi:10.1109/TGRS.2012.2230008, 2013.
- 10 Jethva, H., Torres, O., Waquet, F., Chand, D. and Hu, Y.: How do A-train sensors intercompare in the retrieval of above-cloud aerosol optical depth? A case study-based assessment, *Geophys. Res. Lett.*, 41(1), 186–192, doi:10.1002/2013GL058405, 2014.
- 15 Johnson, B. T., Shine, K. P. and Forster, P. M.: The semi-direct aerosol effect: Impact of absorbing aerosols on marine stratocumulus, *Q. J. R. Meteorol. Soc.*, 130(599), 1407–1422, doi:10.1256/qj.03.61, 2004.
- Josset, D., Hu, Y., Pelon, J., Zhai, P., Tanré, D., Rogers, R., Lucker, P., Trepte, C., Powell, K., Rodier, S., Pascal, N. and Team, T. I.: Advances in Research Products From CALIPSO: Optical Depth Direct Retrieval Over Ocean, Water Clouds and Land, in International Coordination Group on Laser Atmospheric Studies ( ICLAS ). 25th International Laser Radar Conference (ILRC), St. Petersburg, Russia. Curran Associates, Inc., pp. 1280–1283, Curran Associates, Inc., 2010.
- 20 Josset, D., Rogers, R., Pelon, J., Hu, Y., Liu, Z., Omar, A., Zhai, P.-W., Fernald, F. G., Herman, B. M., Reagan, J. A., Winker, D. M., Pelon, J., Coakley, J. A., Ackerman, S. A., Charlson, R. J., Colarco, P. R., Flamant, P., Fu, Q., Hoff, R., Kittaka, C., Kubar, T. L., LeTreut, H., McCormick, M. P., Megie, G., Poole, L., Powell, K., Trepte, C., Vaughan, M. A., Wielicki, B. A., Stephens, G. L., Vane, D. G., Tanelli, S., Im, E., Durden, S., Rokey, M., Reinke, D., Partain, P., Mace, G.
- 25 G., Austin, R., Haynes, J., Lebsock, M., Suzuki, K., Waliser, D., Wu, D., Kay, J., Gettelman, A., Wang, Z. and Marchand, R.: CALIPSO lidar ratio retrieval over the ocean, *Opt. Express*, 19(19), 18696, doi:10.1364/OE.19.018696, 2011.
- Josset, D., Pelon, J., Hu, Y., Rogers, R., Liu, Z., Omar, A., Vaughan, M., Zhai, P. and Team, I.: Global scale lidar ratio retrieval over the ocean, in 26th International Laser Radar Conference (ILRC 26), Porto Heli, Greece., 2012.
- Keil, A. and Haywood, J. M.: Solar radiative forcing by biomass burning aerosol particles during SAFARI 2000: A case study based on measured aerosol and cloud properties, *J. Geophys. Res. Atmos.*, 108(D13), n/a-n/a, doi:10.1029/2002JD002315, 2003.
- King, L. V: On the Complex Anisotropic Molecule in Relation to the Dispersion and Scattering of Light, *Proc. R. Soc. London. Ser. A, Contain. Pap. a Math. Phys. Character*, 104(726), 333–357, 1923.
- Knobelspiesse, K., Cairns, B., Ottaviani, M., Ferrare, R., Hair, J., Hostetler, C., Obland, M., Rogers, R., Redemann, J.,



- Shinozuka, Y., Clarke, A., Freitag, S., Howell, S., Kapustin, V. and McNaughton, C.: Combined retrievals of boreal forest fire aerosol properties with a polarimeter and lidar, *Atmos. Chem. Phys.*, 11(14), 7045–7067, doi:10.5194/acp-11-7045-2011, 2011.
- Lenoble, J., Herman, M., Deuzé, J. L., Lafrance, B., Santer, R. and Tanré, D.: A successive order of scattering code for solving the vector equation of transfer in the earth's atmosphere with aerosols, *J. Quant. Spectrosc. Radiat. Transf.*, 107(3), 479–507, doi:10.1016/j.jqsrt.2007.03.010, 2007.
- Meyer, K., Platnick, S., Oreopoulos, L. and Lee, D.: Estimating the direct radiative effect of absorbing aerosols overlying marine boundary layer clouds in the southeast Atlantic using MODIS and CALIOP, *J. Geophys. Res. Atmos.*, 118(10), 4801–4815, doi:10.1002/jgrd.50449, 2013.
- 5 Meyer, K., Platnick, S. and Zhang, Z.: Simultaneously inferring above-cloud absorbing aerosol optical thickness and underlying liquid phase cloud optical and microphysical properties using MODIS, *J. Geophys. Res. Atmos.*, 120(11), 5524–5547, doi:10.1002/2015JD023128, 2015.
- Myhre, G., Shindell, D., Bréon, F.-M., Collins, W., Fuglestedt, J., Huang, J., Koch, D., Lamarque, J.-F., Lee, D., Mendoza, B., Nakajima, T., Robock, A., Stephens, G., Takemura, T. and Zhang, H.: Anthropogenic and Natural Radiative Forcing, *Clim. Chang. 2013 Phys. Sci. Basis. Contrib. Work. Gr. I to Fifth Assess. Rep. Intergov. Panel Clim. Chang.*, 659–740, doi:10.1017/CBO9781107415324.018, 2013a.
- 15 Myhre, G., Samset, B. H., Schulz, M., Balkanski, Y., Bauer, S., Bernsten, T. K., Bian, H., Bellouin, N., Chin, M., Diehl, T., Easter, R. C., Feichter, J., Ghan, S. J., Hauglustaine, D., Iversen, T., Kinne, S., Kirkev??g, A., Lamarque, J. F., Lin, G., Liu, X., Lund, M. T., Luo, G., Ma, X., Van Noije, T., Penner, J. E., Rasch, P. J., Ruiz, A., Seland, Skeie, R. B., Stier, P.,
- 20 Takemura, T., Tsigaridis, K., Wang, P., Wang, Z., Xu, L., Yu, H., Yu, F., Yoon, J. H., Zhang, K., Zhang, H. and Zhou, C.: Radiative forcing of the direct aerosol effect from AeroCom Phase II simulations, *Atmos. Chem. Phys.*, 13(4), 1853–1877, doi:10.5194/acp-13-1853-2013, 2013b.
- O'Connor, E. J., Illingworth, A. J. and Hogan, R. J.: A Technique for Autocalibration of Cloud Lidar, *J. Atmos. Ocean. Technol.*, 21(5), 777–786, doi:10.1175/1520-0426(2004)021<0777:ATFAOC>2.0.CO;2, 2004.
- 25 Peers, F., Waquet, F., Cornet, C., Dubuisson, P., Ducos, F., Goloub, P., Szczap, F., Tanré, D. and Thieuleux, F.: Absorption of aerosols above clouds from POLDER/PARASOL measurements and estimation of their direct radiative effect, *Atmos. Chem. Phys.*, 15(8), 4179–4196, doi:10.5194/acp-15-4179-2015, 2015.
- Peers, F., Bellouin, N., Waquet, F., Ducos, F., Goloub, P., Mollard, J., Myhre, G., Skeie, R. B., Takemura, T., Tanr??, D., Thieuleux, F. and Zhang, K.: Comparison of aerosol optical properties above clouds between POLDER and AeroCom models over the South East Atlantic Ocean during the fire season, *Geophys. Res. Lett.*, 43(8), 3991–4000, doi:10.1002/2016GL068222, 2016.
- 30 Pinnick, R. G., Jennings, S. G., Chýlek, P., Ham, C. and Grandy, W. T.: Backscatter and extinction in water clouds, *J. Geophys. Res.*, 88(C11), 6787, doi:10.1029/JC088iC11p06787, 1983.
- Powell, K., Vaughan, M., Winker, D., Lee, K. P., Pitts, M., Trepte, C., Detweiler, P., Hunt, W., Lambeth, J. and Lucker, P.:



- Cloud–Aerosol LIDAR Infrared Pathfinder Satellite Observations (CALIPSO), Data Manag. Syst. Data Prod. Cat. Doc. No PC-SCI-503, Release, 3, 2010.
- Ramanathan, V., Crutzen, P. J., Kiehl, J. T. and Rosenfeld, D.: Aerosols, climate, and the hydrological cycle., *Science*, 294(5549), 2119–24, doi:10.1126/science.1064034, 2001.
- 5 Riedi, J., Marchant, B., Platnick, S., Baum, B. A., Thieuleux, F., Oudard, C., Parol, F., Nicolas, J. M. and Dubuisson, P.: Cloud thermodynamic phase inferred from merged POLDER and MODIS data, *Atmos. Chem. Phys.*, 10(23), 11851–11865, doi:10.5194/acp-10-11851-2010, 2010.
- Rosenfeld, D.: Suppression of Rain and Snow by Urban and Industrial Air Pollution, *Science* (80-. ), 287(5459), 1793–1796, doi:10.1126/science.287.5459.1793, 2000.
- 10 Sassen, K.: The Polarization Lidar Technique for Cloud Research: A Review and Current Assessment, *Bull. Am. Meteorol. Soc.*, 72(12), 1848–1866, doi:10.1175/1520-0477(1991)072<1848:TPLTFC>2.0.CO;2, 1991.
- Sassen, K. and Zhu, J.: A global survey of CALIPSO linear depolarization ratios in ice clouds: Initial findings, *J. Geophys. Res. Atmos.*, 114(21), doi:10.1029/2009JD012279, 2009.
- Tanré, D., Bréon, F. M., Deuzé, J. L., Dubovik, O., Ducos, F., François, P., Goloub, P., Herman, M., Lifermann, A. and  
15 Waquet, F.: Remote sensing of aerosols by using polarized, directional and spectral measurements within the A-Train: the PARASOL mission, *Atmos. Meas. Tech.*, 4(7), 1383–1395, doi:10.5194/amt-4-1383-2011, 2011.
- Torres, O., Jethva, H. and Bhartia, P. K.: Retrieval of Aerosol Optical Depth above Clouds from OMI Observations: Sensitivity Analysis and Case Studies, *J. Atmos. Sci.*, 69(3), 1037–1053, doi:10.1175/JAS-D-11-0130.1, 2012.
- Torres, O., Ahn, C. and Chen, Z.: Improvements to the OMI near-UV aerosol algorithm using A-train CALIOP and AIRS  
20 observations, *Atmos. Meas. Tech.*, 6(11), 3257–3270, doi:10.5194/amt-6-3257-2013, 2013.
- Twomey, S.: Pollution and the planetary albedo, *Atmos. Environ.*, 8(12), 1251–1256, doi:10.1016/0004-6981(74)90004-3, 1974.
- Vaughan, M. A., Powell, K. A., Winker, D. M., Hostetler, C. A., Kuehn, R. E., Hunt, W. H., Getzewich, B. J., Young, S. A., Liu, Z. and McGill, M. J.: Fully Automated Detection of Cloud and Aerosol Layers in the CALIPSO Lidar Measurements, *J.*  
25 *Atmos. Ocean. Technol.*, 26(10), 2034–2050, doi:10.1175/2009JTECHA1228.1, 2009.
- Waquet, F., Riedi, J., Labonnote, L. C., Goloub, P., Cairns, B., Deuzé, J.-L. and Tanré, D.: Aerosol Remote Sensing over Clouds Using A-Train Observations, *J. Atmos. Sci.*, 66, 2468–2480, doi:10.1175/2009JAS3026.1, 2009.
- Waquet, F., Peers, F., Ducos, F., Goloub, P., Platnick, S., Riedi, J., Tanré, D. and Thieuleux, F.: Global analysis of aerosol properties above clouds, *Geophys. Res. Lett.*, 40(21), 5809–5814, doi:10.1002/2013GL057482, 2013a.
- 30 Waquet, F., Cornet, C., Deuzé, J.-L., Dubovik, O., Ducos, F., Goloub, P., Herman, M., Lapyonok, T., Labonnote, L. C., Riedi, J., Tanré, D., Thieuleux, F. and Vanbauce, C.: Retrieval of aerosol microphysical and optical properties above liquid clouds from POLDER/PARASOL polarization measurements, *Atmos. Meas. Tech.*, 6(4), 991–1016, doi:10.5194/amt-6-991-2013, 2013b.
- Waquet, F., Péré, J.-C., Peers, F., Goloub, P., Ducos, F., Thieuleux, F. and Tanré, D.: Global detection of absorbing aerosols



- over the ocean in the red and near-infrared spectral region, *J. Geophys. Res. Atmos.*, 121(18), 10,902–10,918, doi:10.1002/2016JD025163, 2016.
- Warren, S. G., Hahn, C. J., London, J., Chervin, R. M. and Jenne, R. L.: Global Distribution of Total Cloud Cover and Cloud Amounts over the Ocean, NCAR Tech. Note, NCAR/TN-27, 42, doi:10.2172/5415329, 1988.
- 5 Wilcox, E. M.: Stratocumulus cloud thickening beneath layers of absorbing smoke aerosol, *Atmos. Chem. Phys.*, 10(23), 11769–11777, doi:10.5194/acp-10-11769-2010, 2010.
- Winker, D. M., Pelon, J. R. and McCormick, M. P.: The CALIPSO mission: spaceborne lidar for observation of aerosols and clouds, in *Third International Asia-Pacific Environmental Remote Sensing Remote Sensing of the Atmosphere, Ocean, Environment, and Space*, edited by U. N. Singh, T. Itabe, and Z. Liu, pp. 1–11, International Society for Optics and
- 10 Photonics., 2003.
- Winker, D. M., Hunt, W. H. and McGill, M. J.: Initial performance assessment of CALIOP, *Geophys. Res. Lett.*, 34(19), L19803, doi:10.1029/2007GL030135, 2007.
- Winker, D. M., Vaughan, M. A., Omar, A., Hu, Y., Powell, K. A., Liu, Z., Hunt, W. H. and Young, S. A.: Overview of the CALIPSO Mission and CALIOP Data Processing Algorithms, *J. Atmos. Ocean. Technol.*, 26(11), 2310–2323,
- 15 doi:10.1175/2009JTECHA1281.1, 2009.
- Xu, M. and Alfano, R. R.: Random Walk of Polarized Light in Turbid Media, *Phys. Rev. Lett.*, 95(21), 213901, doi:10.1103/PhysRevLett.95.213901, 2005.
- Young, S. A. and Vaughan, M. A.: The Retrieval of Profiles of Particulate Extinction from Cloud-Aerosol Lidar Infrared Pathfinder Satellite Observations (CALIPSO) Data: Algorithm Description, *J. Atmos. Ocean. Technol.*, 26(6), 1105–1119,
- 20 doi:10.1175/2008JTECHA1221.1, 2009.
- Yu, H., Kaufman, Y. J., Chin, M., Feingold, G., Remer, L. A., Anderson, T. L., Balkanski, Y., Bellouin, N., Boucher, O., Christopher, S., DeCola, P., Kahn, R., Koch, D., Loeb, N., Reddy, M. S., Schulz, M., Takemura, T. and Zhou, M.: A review of measurement-based assessments of the aerosol direct radiative effect and forcing, *Atmos. Chem. Phys.*, 6(3), 613–666, doi:10.5194/acp-6-613-2006, 2006.
- 25 Yu, H., Remer, L. A., Chin, M., Bian, H., Kleidman, R. G. and Diehl, T.: A satellite-based assessment of transpacific transport of pollution aerosol, *J. Geophys. Res.*, 113(D14), D14S12, doi:10.1029/2007JD009349, 2008.
- Zhang, Y. H., Wooster, M. J., Tutubalina, O. and Perry, G. L. W.: Monthly burned area and forest fire carbon emission estimates for the Russian Federation from SPOT VGT, *Remote Sens. Environ.*, 87(1), 1–15, doi:10.1016/S0034-4257(03)00141-X, 2003.
- 30 Zhang, Z., Meyer, K., Yu, H., Platnick, S., Colarco, P., Liu, Z. and Oreopoulos, L.: Shortwave direct radiative effects of above-cloud aerosols over global oceans derived from 8 years of CALIOP and MODIS observations, *Atmos. Chem. Phys.*, 16(5), 2877–2900, doi:10.5194/acp-16-2877-2016, 2016.



Table 1: Linear regressions of AOT calculated between different methods for three case studies: African biomass-burning aerosols (BBA), Saharan desert dust aerosols (DDA) and Siberian biomass-burning aerosols.  $R^2$  represents the coefficient of determination (COD) between the two sets of data.

Linear regressions		African BBA (13.08.2006)	Saharan DDA (04.08.2008)	Siberian BBA (03.07.2008)
DRM <sub>SODA</sub> vs. POLDER	Slope	0.89±0.01	0.74±0.04	0.56±0.01
	Intercept	0.04±0.01	0.01±0.02	0.07±0.009
	$R^2$ (COD)	0.93	0.79	0.90
DRM <sub>Hu</sub> vs. POLDER	Slope	0.91±0.01	0.74±0.03	0.60±0.01
	Intercept	0.11±0.01	0.15±0.02	0.23±0.009
	$R^2$ (COD)	0.93	0.82	0.89
CALIOP <sub>OM</sub> vs. POLDER	Slope	0.19±0.01	0.86±0.11	0.47±0.08
	Intercept	0.05±0.01	-0.16±0.07	-0.04±0.08
	$R^2$ (COD)	0.35	0.41	0.45



Table 2: Linear regressions of AOT calculated between different methods for data acquired over six months (May to October 2008), over three different regions: South Atlantic Ocean (SAO), North Atlantic Ocean (NAO) and North Pacific Ocean (NPO).

Linear regressions		SAO	NAO	NPO
DRM <sub>SODA</sub> vs. POLDER	Slope	0.89±0.004	0.81±0.009	0.76±0.01
	Intercept	-0.03±0.001	-0.09±0.004	-0.03±0.003
	R <sup>2</sup> (COD)	0.83	0.82	0.37
DRM <sub>Hu</sub> vs. POLDER	Slope	0.90±0.004	0.86±0.01	0.76±0.01
	Intercept	0.05±0.001	0.04±0.004	0.13±0.003
	R <sup>2</sup> (COD)	0.82	0.82	0.44
CALIOP <sub>OM</sub> vs. POLDER	Slope	0.34±0.004	0.52±0.02	0.28±0.02
	Intercept	-0.04±0.002	-0.01±0.01	0.01±0.01
	R <sup>2</sup> (COD)	0.43	0.42	0.24
CALIOP <sub>OM</sub> vs. DRM <sub>SODA</sub>	Slope	0.34±0.002	0.62±0.01	0.35±0.01
	Intercept	-0.01±0.002	0.04±0.006	0.01±0.007
	R <sup>2</sup> (COD)	0.42	0.48	0.28



Table 3: Regional analysis using CALIOP measurements over six months (May to October 2008), over South Atlantic Ocean (SAO), North Atlantic Ocean (NAO) and North Pacific Ocean (NPO): mean cloud altitude for altitudes smaller than 5 km; mean aerosol base and top altitudes for altitudes smaller than 10 km.

	SAO	NAO	NPO
Mean cloud top altitude [km]	1.24±0.43	1.35±0.5	1.09±0.84
	Min: 0.30	Min: 0.20	Min: 0.05
	Max: 4.95	Max: 3.25	Max: 5.0
Mean aerosol top altitude [km]	3.83±0.093	4.50±1.03	2.74±1.68
	Min: 0.50	Min: 0.44	Min: 0.47
	Max: 6.73	Max: 6.67	Max: 9.85
Mean aerosol base altitude [km]	2.90±0.97	2.97±1.12	3.48±1.78
	Min: 0.02	Min: 0.02	Min: 0.05
	Max: 5.80	Max: 5.74	Max: 9.31



Table 4: Calculated mean, minimum value and maximum value of  $AOT_{532nm}$  over six months in 2008, for three regions (SAO, NAO, NPO) and for different methods.

$AOT_{532nm}$		SAO	NAO	NPO
POLDER	Mean	$0.35 \pm 0.23$	$0.39 \pm 0.21$	$0.18 \pm 0.21$
	Min	0.005	0.005	0.005
	Max	1.27	1.19	2.17
DRM <sub>SODA</sub>	Mean	$0.28 \pm 0.22$	$0.23 \pm 0.19$	$0.15 \pm 0.38$
	Min	-0.13	-0.16	-0.16
	Max	1.30	0.95	3.26
DRM <sub>Hu</sub>	Mean	$0.37 \pm 0.23$	$0.38 \pm 0.20$	$0.32 \pm 0.40$
	Min	-0.07	-0.06	-0.06
	Max	1.50	1.17	3.68
CALIOP <sub>OM</sub>	Mean	$0.12 \pm 0.11$	$0.23 \pm 0.18$	$0.14 \pm 0.23$
	Min	0.001	0.005	0.001
	Max	1.88	2.38	2.01



Table 5: Mean value of AE over six months in 2008, for three regions (SAO, NAO, NPO) for different methods after filtering the POLDER  $AOT_{865nm} > 0.1$  and  $DRM_{SODA} AOT_{532nm} > 0.1$ , respectively CALIOP<sub>OM</sub>  $AOT_{532nm} > 0.1$ .

		SAO	NAO	NPO
POLDER	Mean $AE_{670/865}$	$2.05 \pm 0.27$	$0.49 \pm 0.27$	$1.67 \pm 0.50$
	Min	0.36	0.36	0.36
	Max	2.56	2.03	2.39
$DRM_{SODA}$	Mean $AE_{532/1064}$	$1.79 \pm 0.58$	$0.10 \pm 0.27$	$1.47 \pm 0.84$
	Min	-1.15	-1.14	-1.21
	Max	4.19	1.43	3.93
CALIOP <sub>OM</sub>	Mean $AE_{532/1064}$	$0.97 \pm 0.51$	$-0.19 \pm 0.32$	$0.41 \pm 0.72$
	Min	-2.27	-1.62	-2.63
	Max	3.16	1.27	4.41



Table 6: Linear regression calculated between  $DRM_{SODA} AOT_{532nm}$  and  $POLDER AOT_{532nm}$  for situation when the aerosol layer is attached to the cloud top and when the aerosol layer is well separated from the cloud over three regions (South Atlantic Ocean, North Atlantic Ocean and North Pacific Ocean) and for a period of 4.5 years.

	SAO		NAO		NPO	
	Attached	Detached	Attached	Detached	Attached	Detached
Slope	$0.60 \pm 0.02$	$0.77 \pm 0.003$	$0.63 \pm 0.07$	$0.59 \pm 0.01$	$0.78 \pm 0.12$	$0.80 \pm 0.02$
Intercept	$0.04 \pm 0.006$	$0.02 \pm 0.001$	$-0.005 \pm 0.02$	$-0.011 \pm 0.006$	$-0.04 \pm 0.02$	$-0.015 \pm 0.007$
$R^2$ (COD)	0.54	0.715	0.39	0.57	0.19	0.435



Table 7: Linear regressions calculated between different methods for data acquired over June 2006 to December 2010, on a global scale above the ocean in the case of aerosol *attached* to the cloud top, *detached* from the cloud and *undetermined* situations for AOT smaller than 1.5.

Linear regressions		Detached	Attached	Undetermined
DRM <sub>SODA</sub> vs. POLDER	Slope	0.84±0.003	0.59±0.01	0.24±0.001
	Intercept	-0.03±0.001	-0.02±0.002	-0.02
	R <sup>2</sup> (COD)	0.68	0.36	0.03
DRM <sub>Hu</sub> vs. POLDER	Slope	0.78±0.002	0.55±0.001	0.28±0.001
	Intercept	0.10±0.001	0.12±0.002	0.09
	R <sup>2</sup> (COD)	0.68	0.36	0.05
CALIOP <sub>OM</sub> vs. POLDER	Slope	0.17±0.002	0.12±0.007	0.06±0.008
	Intercept	0.013	0.02±0.001	0.14±0.002
	R <sup>2</sup> (COD)	0.15	0.047	0.003
CALIOP <sub>OM</sub> vs. DRM <sub>SODA</sub>	Slope	0.17±0.002	0.1±0.007	0.21±0.01
	Intercept	0.029	0.04±0.001	0.14±0.001
	R <sup>2</sup> (COD)	0.15	0.03	0.01

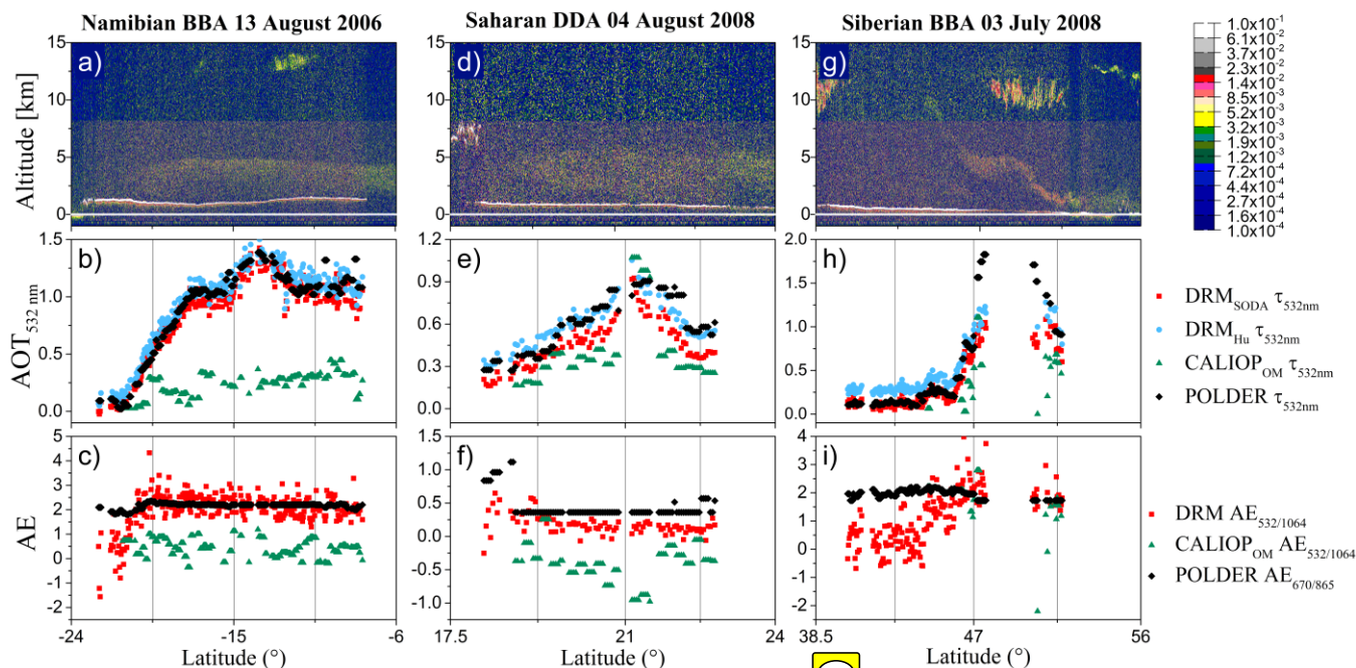
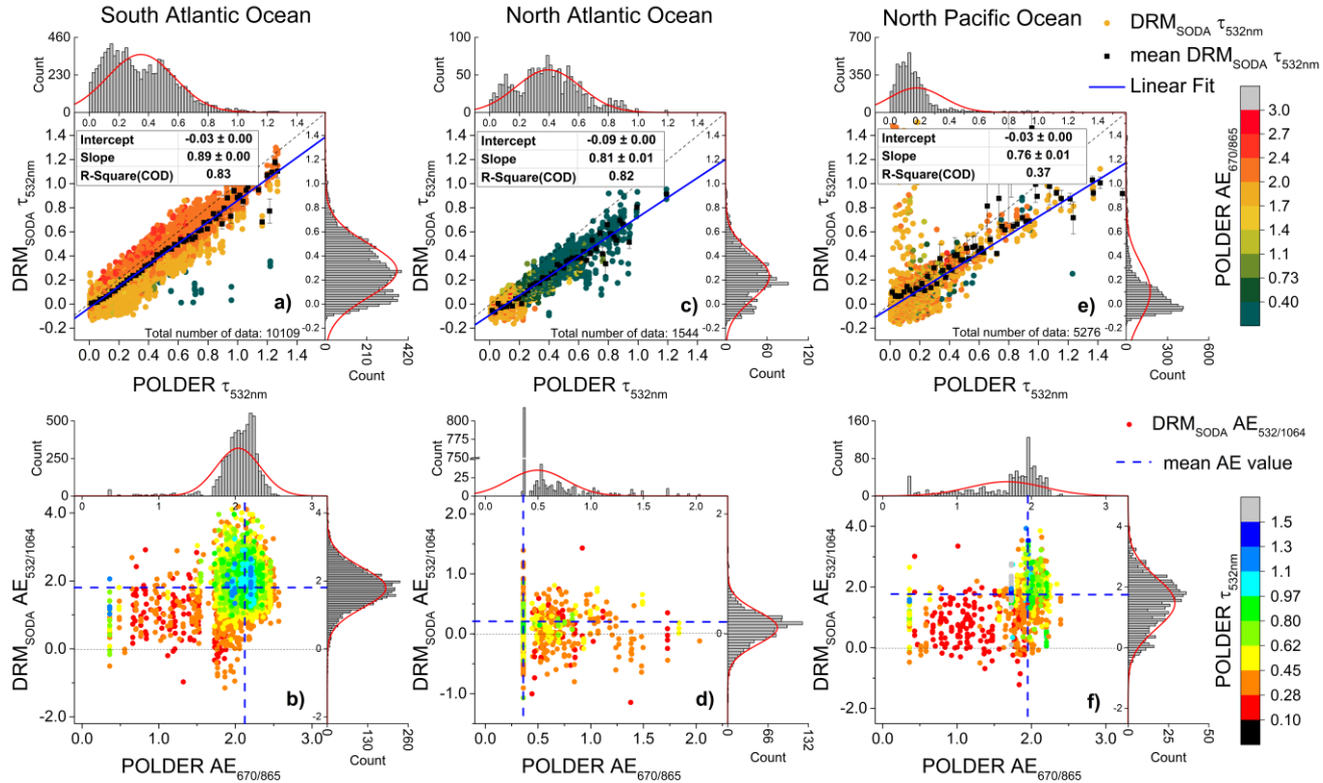


Figure 1: The first row of the panel shows the lidar CALIOP backscatter profile at 532 nm (km<sup>-1</sup> sr<sup>-1</sup>) for three case studies: African biomass-burning (BBA) aerosols above clouds on 13 August 2006 ((a), (b), (c)), Saharan dust (DDA) on 4 August 2008 ((d), (e), (f)) and Siberian biomass-burning aerosols over the Okhotsk Sea on 3 July 2008 ((g), (h), (i)). For these cases, the above-cloud AOT at 532 nm and the Ångström exponent (AE) as a function of latitude, measured with several techniques are displayed.

5



**Figure 2:** The first row of the panel shows the comparison of AOT at 532 nm retrieved from DRM<sub>SODA</sub> and POLDER methods, with the corresponding POLDER AE color scale, computed between 670 and 865 nm. The second row presents the Ångström exponent comparison for AOTs larger than 0.1, retrieved from DRM<sub>SODA</sub> and POLDER methods, with the corresponding POLDER AOT at 532 nm color scale. The measurements were made over a period of six months (May to October 2008) and over three distinctive regions: South Atlantic Ocean - between 30° S to 5° N and 12° W to 14° E ((a) and (b)), North Atlantic Ocean - between 10 to 35° N and 10 to 40° W ((c) and (d)) and North Pacific Ocean - between 35 to 60° N and 140 to 170° E ((e) and (f)). The histograms present the data distribution. The error bars in figures (a), (c) and (e) represent the standard error of the mean (SEM).

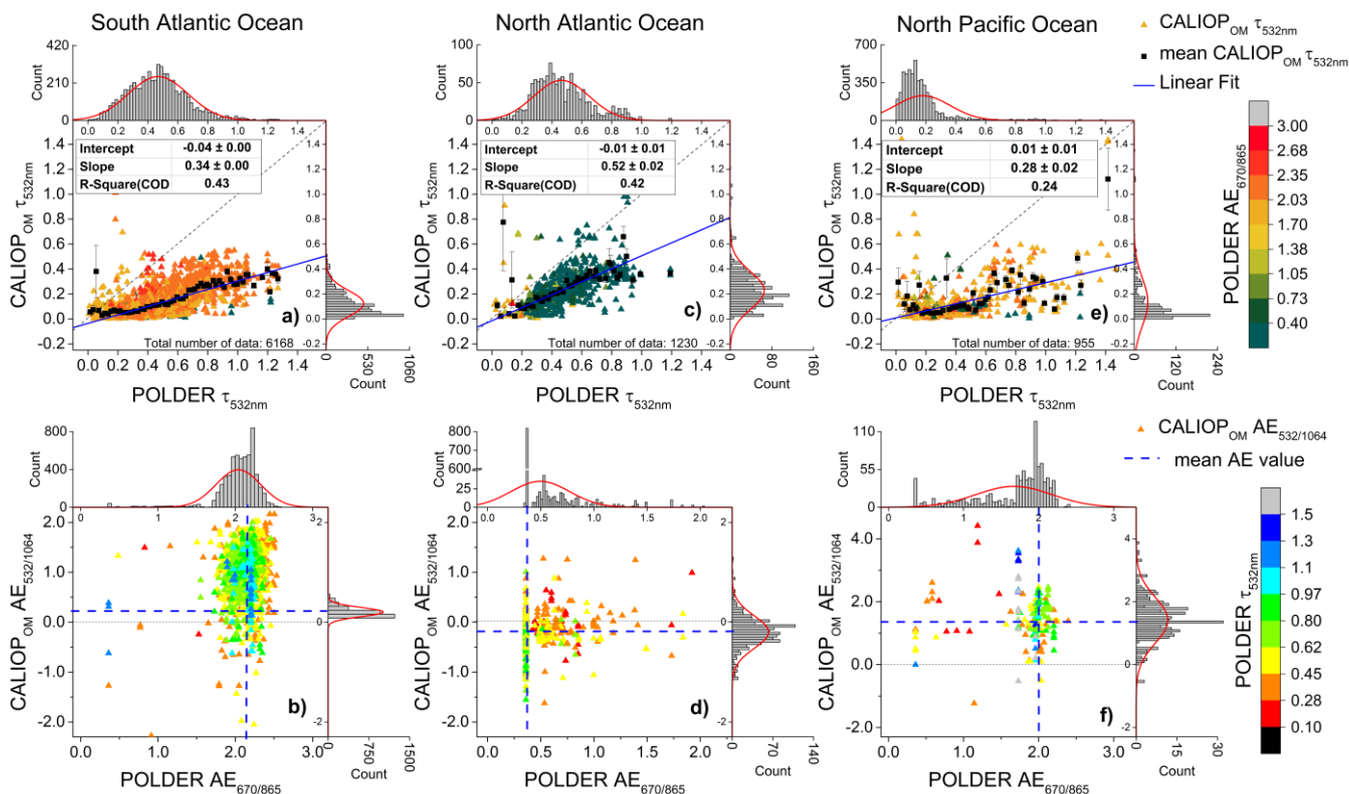
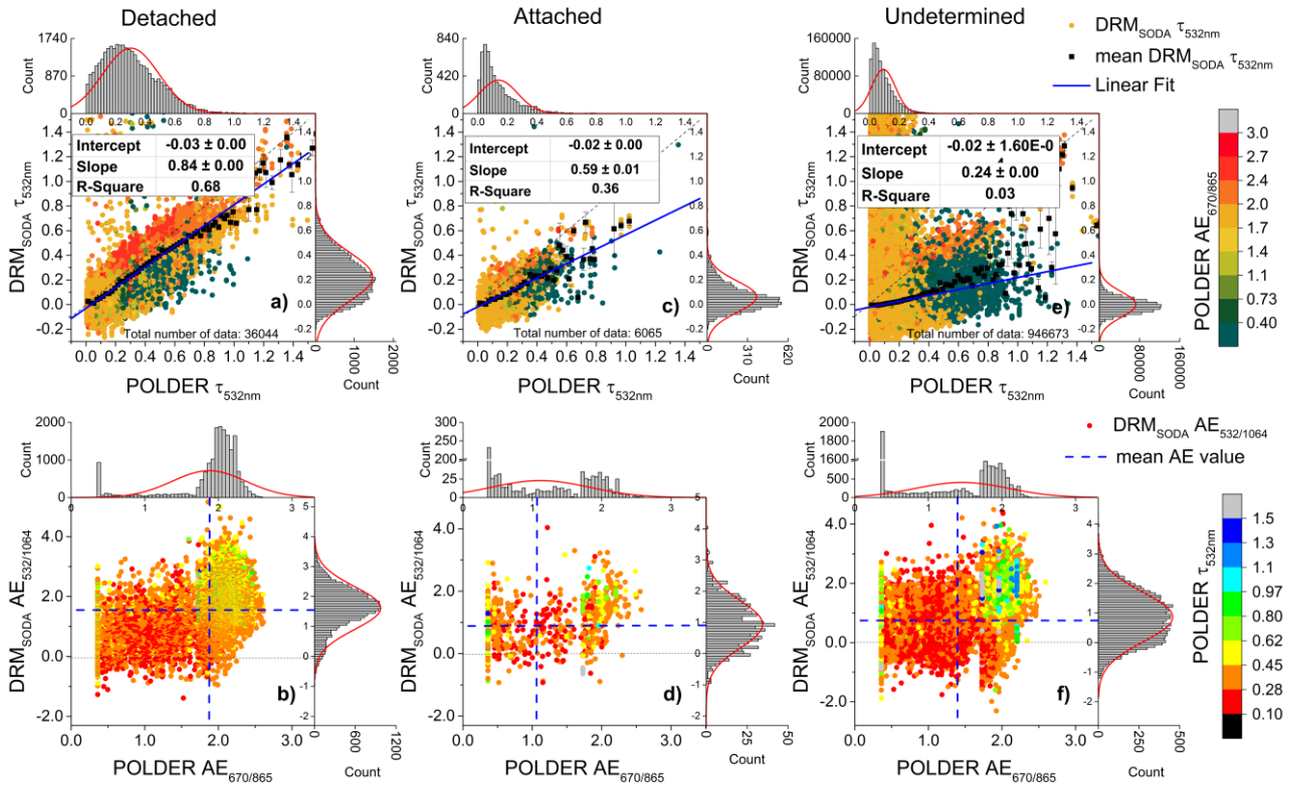


Figure 3: Same as Fig. 2, retrieved from CALIOP operational method and POLDER method.



**Figure 4:** Global comparison over a period of 4.5 years (June 2006 to December 2010) for situations with aerosol layer well separated from the cloud top - *detached* ((a) and (b)), for cases where the aerosol layer is in contact with the cloud – *attached* ((c) and (d)) and for *undetermined* situations ((e) and (f)). The comparison of AOT at 532 nm retrieved from DRM<sub>SODA</sub> and POLDER methods is shown in the first row. The color scale represents the corresponding POLDER AE computed between 670 and 865 nm. The second row presents the Ångström exponent for AOTs larger than 0.1, with a POLDER AOT at 532 nm color scale. The histograms present the data distribution. The error bars in figures (a), (c) and (e) represent the standard error of the mean (SEM).

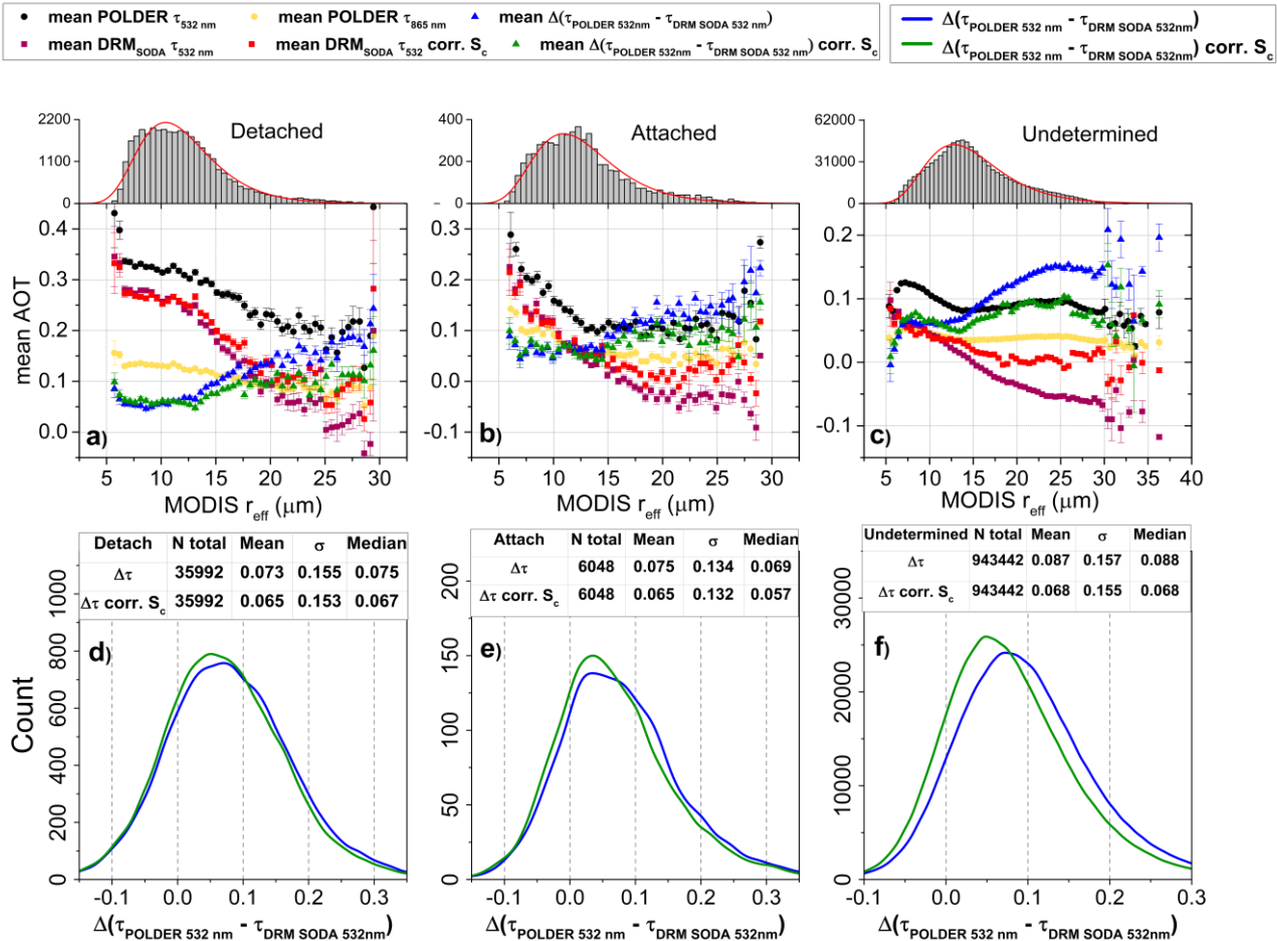


Figure 5: 4.5 years of global study on the evolution of POLDER and  $\text{DRM}_{\text{SODA}}$  above cloud AOT retrievals as a function of MODIS effective radius ( $r_{eff}$ ,  $\mu\text{m}$ ) for situations where: the aerosol layer is *detached* from the cloud top ((a) and (d)), for cases where the aerosol layer is *attached* to the cloud top ((b) and (e)) and for *undetermined* situations (c) and (f). The histograms from figures (a), (b) and (c) represent the distribution of  $r_{eff}$ . The histograms in figures (d), (e) and (f) present the difference between POLDER and  $\text{DRM}_{\text{SODA}}$  mean AOTs, before the correction of  $\text{DRM}_{\text{SODA}}$  AOT with  $r_{eff}$  ( $\Delta\tau$ ) and after this correction ( $\Delta\tau \text{ corr. } S_c$ ). The associated tables indicate the number of cases, mean, standard deviation ( $\sigma$ ) and median values of these differences. The error bars in figures (a), (b) and (c) represent the standard error of the mean (SEM).

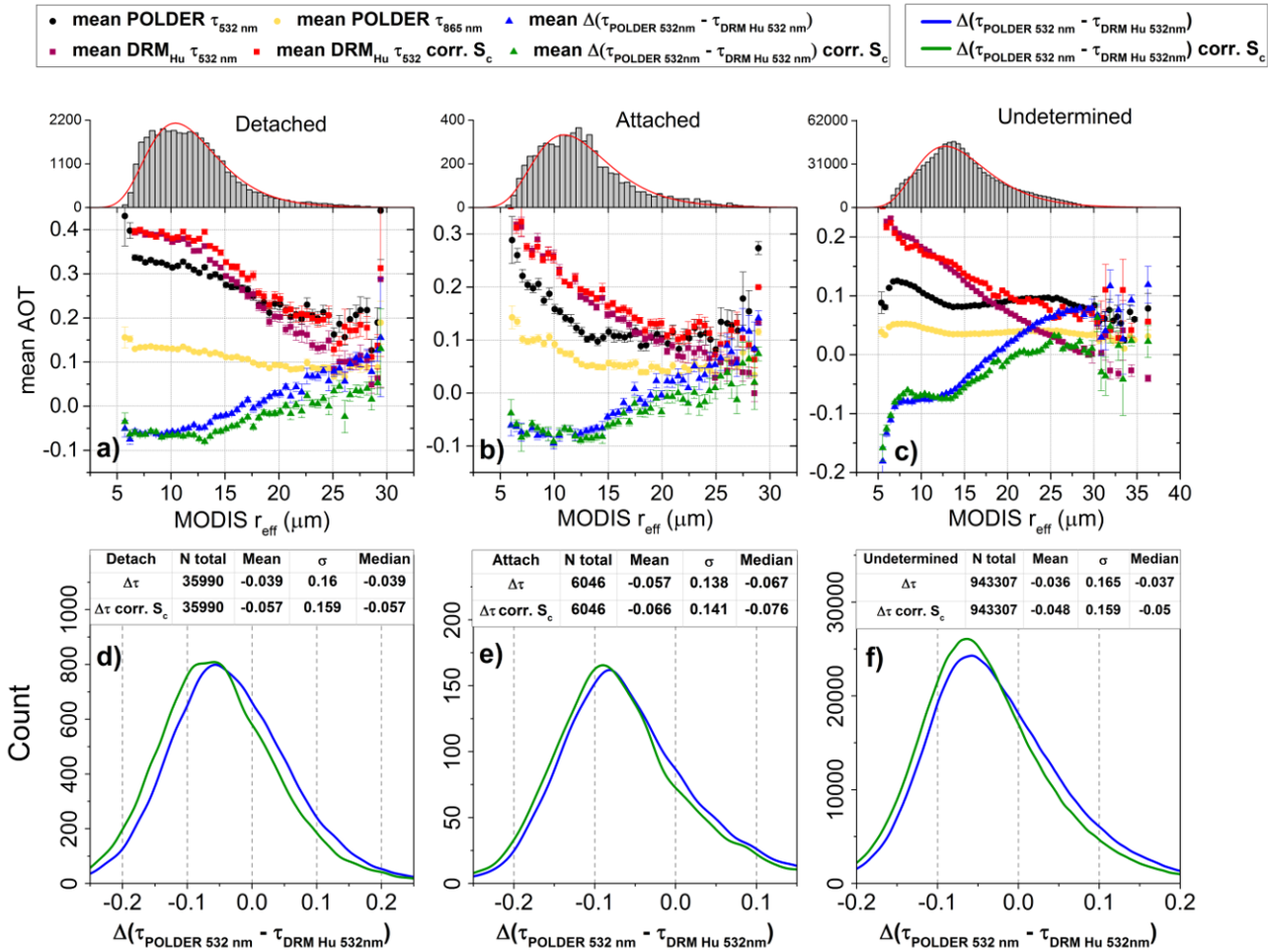
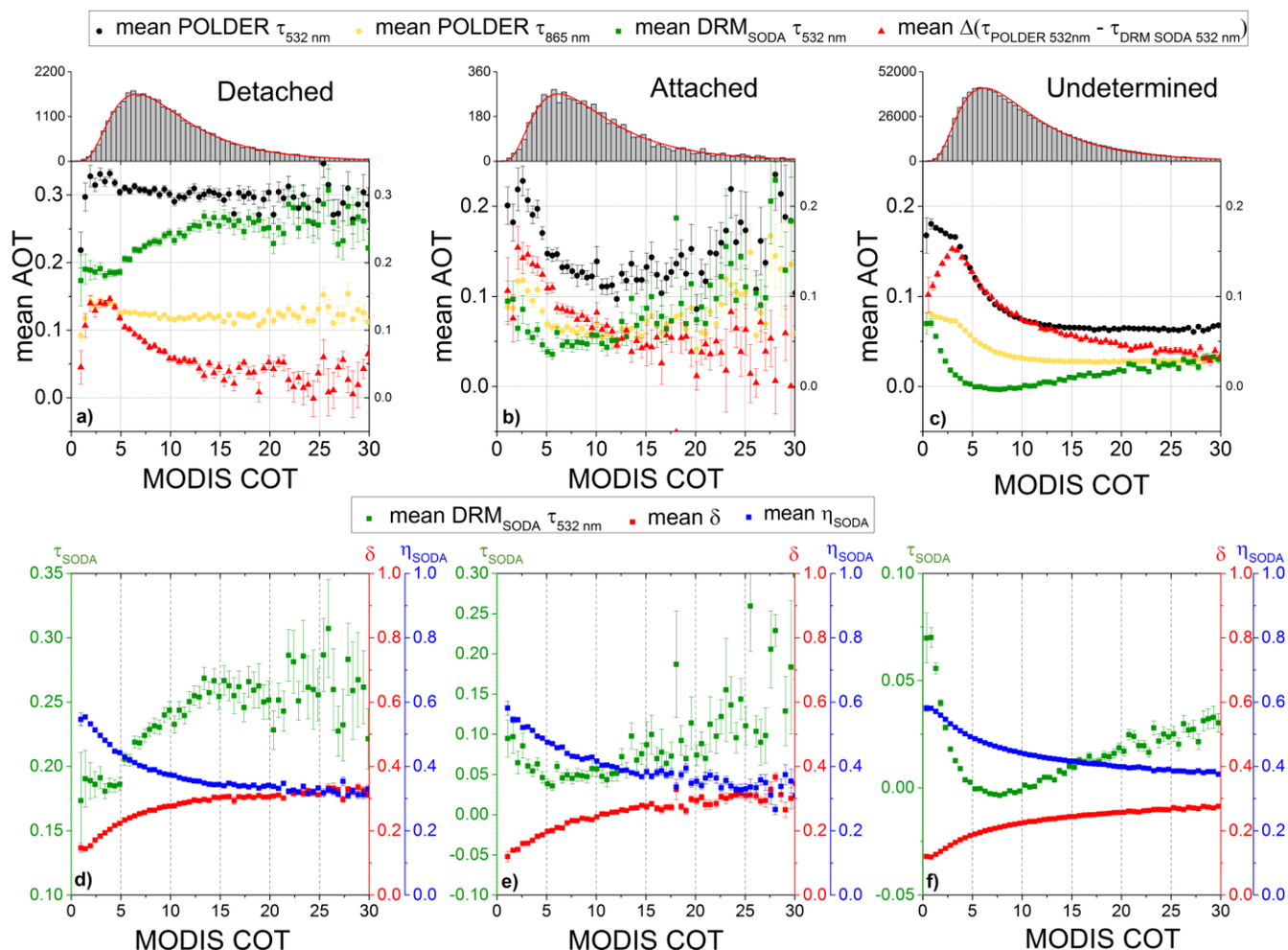


Figure 6: Same as Fig. 5, POLDER and DRM<sub>Hu</sub> above cloud AOT retrievals as a function of MODIS effective radius ( $r_{\text{eff}}$ ,  $\mu\text{m}$ ).



**Figure 7:** 4.5 years of global study of the evolution of POLDER and DRM<sub>SODA</sub> above cloud AOT retrievals, as well as the difference of these two methods as a function of MODIS cloud optical thickness (COT), for situations where: the aerosol layer is *detached* from the cloud top (a), for cases where the aerosol layer is *attached* to the cloud top (b) and for *undetermined* situations (c). The histograms represent the distribution of COT. The error bars show the standard error of the mean (SEM). Figures (d), (e) and (f) display the evolution of DRM<sub>SODA</sub> AOT ( $\tau_{SODA}$ ), depolarization ratio ( $\delta$ ) and multiple scattering factor ( $\eta_{SODA}$ ) as a function of MODIS COT, for the abovementioned situations.

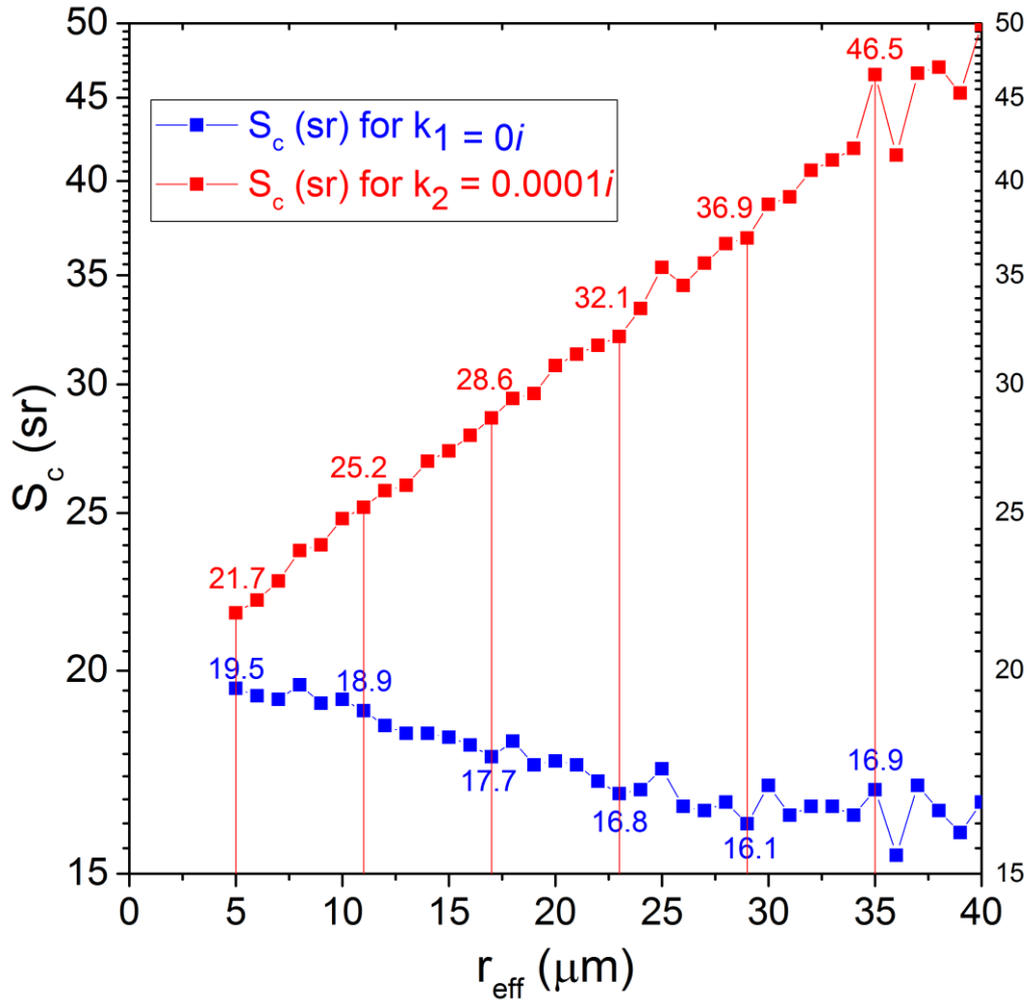
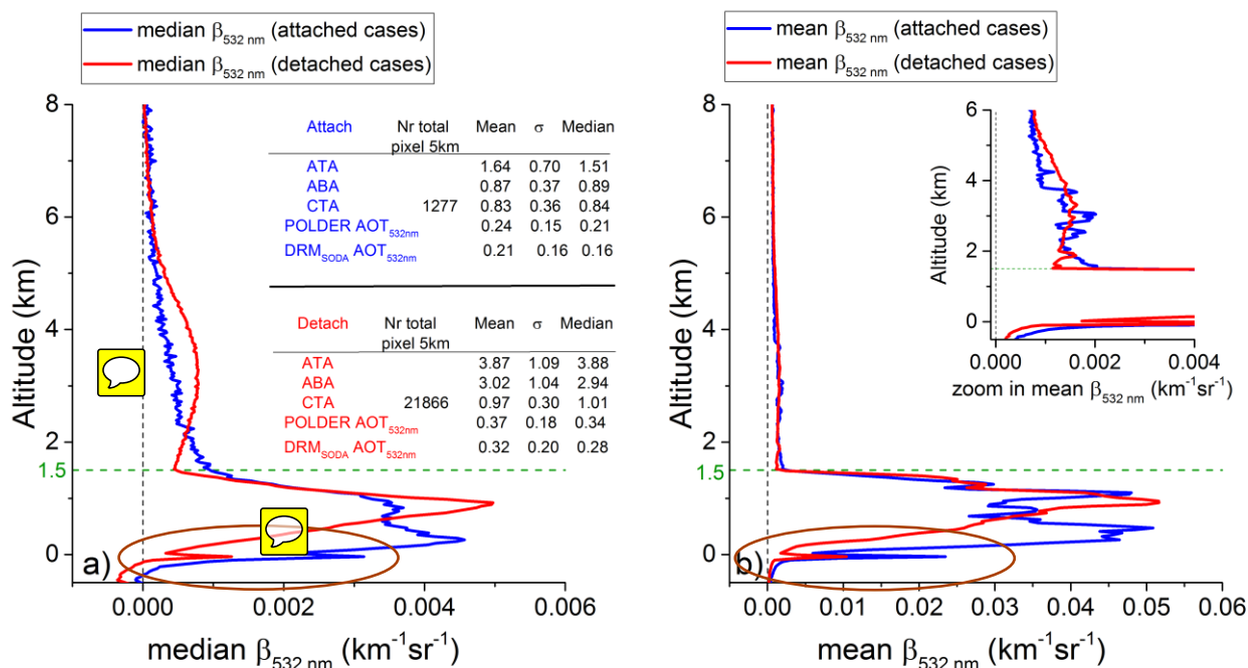


Figure 8: Sensitivity study of lidar ratio ( $S_c$ , sr) as a function of the cloud droplets effective radius, using a two-parameter Gamma size distribution in Mie code. The effective variance,  $v_{\text{eff}}$  is set to 0.088. The real part of the refractive index is fixed to 1.337, while the imaginary part,  $k$ , was set to 0 (blue) and to  $0.0001i$  (red).



**Figure 9: Median (a) and averaged (b) backscatter profiles ( $\text{km}^{-1}\text{sr}^{-1}$ ) for aerosol layer detached from the cloud layer (red) and aerosols attached to the top of the cloud (blue), for a period of 4.5 years on the global scale. The data was filtered for a cloud top altitude lower than 1.5 km and for a DRM<sub>SODA</sub> AOT at 532 nm larger than 0.1. The number of 5 km horizontal resolution pixels is also shown. The mean, standard deviation ( $\sigma$ ) and median of aerosol top altitude (ATA), aerosol base altitude (ABA) and cloud top altitude (CTA) are given for each situation. Same values are shown for POLDER AOT at and DRM<sub>SODA</sub> AOT at 532 nm.**

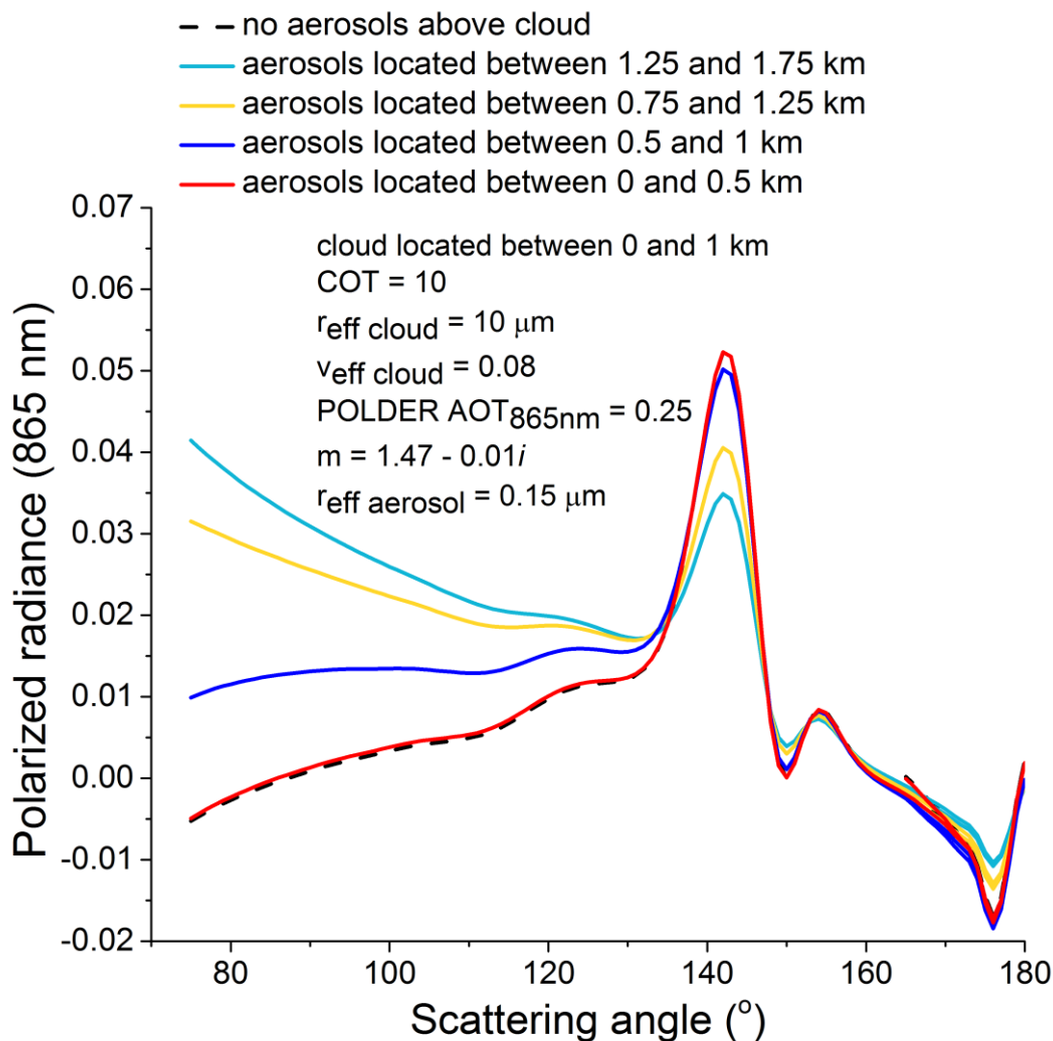


Figure 10: Sensitivity study of polarized radiance at 865 nm to the relative position of the aerosol layer above the cloud. Simulation performed for a cloud layer located between 0 and 1 km and aerosol layers varying at different altitudes. The cloud droplet effective radius is fixed to 10  $\mu\text{m}$  and the effective variance is 0.08. The aerosol layer is characterized by an AOT of 0.25 at 865 nm, a refractive index of 1.47-0.01*i* and an aerosol effective radius of 0.15  $\mu\text{m}$ .