How much of the global aerosol optical depth is found in the boundary layer and free troposphere?

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Abstract. The global aerosol extinction from the CALIOP space lidar was used to compute aerosol optical depth (AOD) over a 9-year period (2007–2015) and partitioned between the boundary layer (BL) and the free troposphere (FT) using BL heights obtained from the ERA-Interim archive. The results show that the vertical distribution of AOD does not follow the diurnal cycle of the BL but remains similar between day and night highlighting the presence of a residual layer during night. The BL and FT contribute 69 and 31 %, respectively, to the global tropospheric AOD during daytime in line with observations obtained in Aire sur l’Adour (France) using the Light Optical Aerosol Counter (LOAC) instrument. The FT AOD contribution is larger in the tropics than at mid-latitudes which indicates that convective transport largely controls the vertical profile of aerosols. Over oceans, the FT AOD contribution is mainly governed by long-range transport of aerosols from emission sources located within neighboring continents. According to the CALIOP aerosol classification, dust and smoke particles are the main aerosol types transported into the FT. Overall, the study shows that the fraction of AOD in the FT – and thus potentially located above low-level clouds – is substantial and deserves more attention when evaluating the radiative effect of aerosols in climate models. More generally, the results have implications for processes determining the overall budgets, sources, sinks and transport of aerosol particles and their description in atmospheric models.

1 Introduction

Aerosols influence the radiative budget of the Earth by absorbing or scattering solar radiation and by changing microphysical properties of clouds. Overall, aerosol particles are short-lived (about a week) compounds in the atmosphere due to the efficient removal by dry and wet deposition in the boundary layer (BL) where a majority of the emission sources are located (e.g., Stier et al., 2005; Dentener et al., 2006). As a consequence, most of the aerosol mass is expected to be found in the BL and the aerosol optical depth (AOD), i.e., the aerosol extinction integrated over the atmospheric column, is expected to be dominated by the BL contribution. Indeed, AOD roughly corresponds to aerosol mass concentration because particles need to be optically active (larger than the wavelength of the incident light) to scatter light. Furthermore, if most of the aerosol mass is found within the BL, then BL aerosols might be expected to have the largest climate effect. However, aerosols advected to the
free troposphere (FT) have a much longer residence time (typically a few weeks) than those in the BL, potentially inducing a more long-term effect on climate and even acting as a source for particles back to the BL. Light-absorbing aerosols may in addition cause enhanced absorption, and thereby a climate warming effect, if they are located above low-level reflective clouds (Zarzycki and Bond, 2010; Zhang et al., 2014), and they may thus counteract the radiative effect of Twomey cloud brightening (Bender et al., 2016). Light-absorbing aerosols above clouds may also modify cloud formation and evolution, e.g., delay the transition of stratocumulus to cumulus clouds (Yamaguchi et al., 2015; Wilcox et al., 2016). Therefore, accurately characterizing the vertical and regional distribution of aerosols – and in particular discriminating between particles in the BL and FT – is crucial in order to properly assess the role of aerosols in the climate system. Furthermore, understanding the transport processes between BL and FT is a necessity for constraining the overall sources, sinks and budgets of atmospheric aerosol constituents.

The BL is the atmospheric layer directly influenced by the underlying surface and it can in general be assumed to be chemically well mixed. The BL is governed by a strong diurnal cycle over land and adjusts to surface forcings within a timescale of an hour (Stull, 1988). The continental BL is typically stable and shallow during the night (< 500 m), and becomes unstable during the day, mixing with the air masses above and extending up to a few kilometers. Processes within the BL control exchange of momentum, heat, moisture and trace gases between the surface and the FT. In addition, the height of the BL – together with the amount of emitted particles – influences the concentration of aerosols near the surface and consequently governs air quality (Liu et al., 2013; Yang et al., 2015). During severe pollution events, a high concentration of particles may also enhance the stability of the BL, reducing its thickness and further increasing the concentration of particles within the BL (Petäjä et al., 2016). Kafle and Coulter (2013) found evidence that the vertical distribution of AOD does not seem to vary with the diurnal cycle of the BL evolution. Although some recent studies have looked at the vertical distribution of aerosols (e.g., Koffi et al., 2012, 2016; Toth et al., 2016), none has to our knowledge specifically determined the partitioning of AOD between the BL and FT on a global scale.

The parameterization of vertical transport of aerosols in models is crucial for the residence time, long-range transport, dynamic processes and the radiative effect of aerosols (e.g., Textor et al., 2006). Nevertheless, large uncertainties still remain regarding the processes controlling the vertical transport. Several studies have pointed out convective transport and in-cloud scavenging as the main processes determining the vertical distribution of aerosols (e.g., Cui and Carslaw, 2006; Ekman et al., 2006; Bourgeois and Bey, 2011; Weigel et al., 2011; Kipling et al., 2016). However, other processes such as condensation, biomass burning injection height, degradation and BL mixing have also been reported as important (Kipling et al., 2016; Peers et al., 2016). In addition, pyro-convection and orographic lifting are two regional processes that can transport aerosols from the surface to the FT (e.g., Fromm et al., 2006; Yumimoto et al., 2009; Bourgeois et al., 2015). Although Veira et al. (2015) showed that only a small fraction (4–5 %) of biomass burning plumes reach the FT, Devasthale and Thomas (2011) indicated that smoke plumes are frequently found above clouds over the ocean near intense biomass burning regions. Peers et al. (2016) reported that many state-of-the-art aerosol–climate models failed to reproduce an observed event where a large smoke plume was transported above clouds over the eastern tropical Atlantic Ocean. It is thus essential to better constrain atmospheric models in terms of AOD below and above clouds in order to accurately predict the radiative effect of aerosols.

Although atmospheric aerosols have been the subject of intense research for the last decades, aerosol observations covering the whole atmospheric column are still relatively sparse. To fill this gap in knowledge and data availability, the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) satellite instrument, dedicated to measuring vertically resolved attenuated backscatter, was launched in space in June 2006 (Winker et al., 2009). In this study, CALIOP observations are used together with the BL height product from the ERA-Interim archive (Berrisford et al., 2011; Dee et al., 2011) in order to discriminate the amount of AOD present in the BL and FT (Sect. 2). The main objective of this study is to evaluate the vertical distribution of AOD in the BL and FT during day and night, respectively (Sect. 3). Finally, results are discussed and summarized in Sect. 4.

2 Data and methods

The partitioning of the AOD in the BL (from the surface to the top of the boundary layer) and FT (from the boundary layer top to the tropopause) has been computed from the CALIOP aerosol extinction retrievals and the BL height from the ERA-Interim archive. The final output is monthly averaged AOD values over a 2 × 2° grid from January 2007 to December 2015. Results for the Antarctica region are likely spurious and thus excluded because high aerosol values are reported by CALIOP despite the region being known to be pristine.

2.1 CALIOP observations

The CALIOP instrument onboard the Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observation (CALIPSO) satellite is a two-wavelength polarization lidar (532 and 1064 nm) that provides vertically resolved attenuated backscatter between the surface and 40 km for both daytime and nighttime along its orbit (Winker et al., 2009). Depending on the altitude, profiles are sampled at a vertical reso-
were considered, and negative aerosol extinction values and a control flag value of 0, 1, 18, or 16 and cloud-free scenes were included when selecting CALIOP retrievals. With the A VD feature flag, an Atmospheric V olume Description (A VD) product containing Version 4.10 Level 2 532 nm aerosol extinction data and the aerosol extinction type is indicated for each aerosol layer and each profile bin. Note that although the solar background illumination decreases the signal-to-noise ratio during daytime leading to a better aerosol extinction detection sensitivity during the night. The detection threshold of the aerosol extinction also depends on the nature of the particles. Overall, the CALIOP aerosol extinction detection sensitivity varies between about 0.01 and 0.07 km$^{-1}$, inducing possible misdetections and resulting in a potential underestimate of the lowest extinction values especially at high latitudes and in the FT (Winker et al., 2013; Rogers et al., 2014; Toth et al., 2018). Although uncertainties remain regarding retrieved aerosol extinction values in the troposphere, CALIOP algorithms have been improved over the years, leading to a realistic and representative view of global aerosols (Winker et al., 2013). A comparison of CALIOP data with MODIS retrievals is performed below and with in situ measurements in Sect. 3.3.

In this study, we use 9 years (2007–2015) of CALIOP Version 4.10 Level 2 532 nm aerosol extinction data and the Atmospheric Volume Description (AVD) product containing feature classification flags. With the AVD feature flag, an aerosol type is indicated for each aerosol layer and each profile bin. Note that although the solar background illumination decreases the signal-to-noise ratio during daytime, both daytime and nighttime aerosol extinction data are here analyzed in a similar manner. To minimize the uncertainties associated with the aerosol extinction, the data were screened following the recommendation by Winker et al. (2013): only aerosol extinction values with a cloud aerosol discrimination confidence inclusive of $-100$ to $-20$, with an extinction quality control flag value of 0, 1, 18, or 16 and cloud-free scenes were considered, and negative aerosol extinction values and “clear air” (meaning air without aerosols) below an aerosol layer base lower than 2.46 km were removed. The 2.46 km threshold was chosen not to bias low the aerosol extinction near the surface where the layer detection algorithm sometimes places the aerosol layer base above the surface (Winker et al., 2013). Finally, aerosol extinction profiles are averaged only for ten or more profiles per grid box which excludes 1 % of data but leads to more robust statistics. With help of the AVD product, the aerosol type of each aerosol extinction data point is determined before the calculation of the AOD. Therefore, AODs are computed for each of the seven aerosol types considered by CALIOP, for both the BL and FT as further described in the Sect. 2.2.

This screening procedure leads to the global AOD averages over land and ocean listed in Table 1. Our calculated AOD values are in good agreement with those previously derived using data from CALIOP (Winker et al., 2013) as well as the MODIS-Aqua instrument (Collection 6; Levy et al., 2013). The slightly larger values in the present study compared to Winker et al. (2013) are most likely due to the use of V4.10 CALIOP data instead of CALIOP V3.

Overall, more than 1.1 million CALIOP vertical profiles are used per month; 55 % (45 %) of them are retrieved during daytime (nighttime) and 40 % (60 %) of them are retrieved over land (ocean).

### 2.2 ERA-Interim data

The European Centre for Medium-range Weather Forecasts Re-Analysis-Interim (ERA-Interim) archive is based on data assimilation of observations, in particular from satellites, and numerical weather prediction modeling (Berrisford et al., 2011; Dee et al., 2011). The boundary layer height product is diagnosed using the parcel lifting method (or bulk Richardson method) proposed by Troen and Mahrt (1986). The product has been evaluated and showed good agreement with other methods to determine the BL height (von Engeln and Teixeira, 2013). In this study, the ERA-Interim boundary layer height product was used at a horizontal resolution of 1° and with a 3 h time step. Since the CALIOP spatial and temporal resolution (5 km and 6 s, respectively) is higher than ERA-Interim, the height of the BL is determined for each
CALIOP vertical profile by using the corresponding ERA-Interim grid box and the closest time step. This value is used to separate the BL and FT AOD before averaging AOD values over a $2 \times 2^\circ$ grid. The different spatial and temporal resolutions between CALIOP retrievals and ERA-Interim data likely leads to spatial and temporal sampling errors (Schutgens et al., 2016); however, an increase in the horizontal resolution of ERA-Interim data from 1 to $0.5^\circ$ changed our AOD results by only 0.3–0.4 %. Since the spatial sampling error does not vary significantly, we used $1^\circ$ horizontal resolution assimilation data for computational efficiency.

As expected, the diurnal cycle of the ERA-Interim BL is more pronounced over land than over ocean. The global and annual average of the BL height varies from about 200 m during night to 1400 m during day above land, while it remains around 700 m over ocean both during day and night (Fig. 1). Sensitivity tests have been performed on the BL height to evaluate its influence on the AOD partitioning between the BL and FT. They showed that a global increase (decrease) of the BL height by 120 m increases (decreases) the global contribution of the BL AOD by about 5 %.

### 3.1 Daytime analysis

The seasonally averaged global distribution of BL and FT daytime AODs over 9 years (2007–2015) are shown in Fig. 2. High values of AOD are observed both in the BL and FT over central (all seasons) and southern (JJA and SON) Africa, the Arabian Peninsula, India and eastern China. This distribution is expected as these regions correspond to the main continental sources of aerosol mass (Ginoux et al., 2001; Giglio et al., 2013; Janssens-Maenhout et al., 2015). Typical features such as transport of aerosols over the Bay of Bengal and the Indo-Gangetic plain (Höpner et al., 2016) or long-range transport of dust aerosols from the Sahara over the Atlantic Ocean (e.g., Generoso et al., 2008) are also observed in the CALIOP data, both in the BL and in the FT. Overall, the global daytime AOD is 0.147 in the troposphere, where 0.102 is found in the BL and 0.045 in the FT (Table 2), corresponding to 69 and 31 %, respectively, of the total AOD. Although the global AOD over land (0.182) and ocean (0.129) is very different, the AOD partitioning between the BL and FT is different.
Figure 2. Seasonal BL (a: DJF; b: MAM; c: JJA; d: SON) and FT (e: DJF; f: MAM; g: JJA; h: SON) daytime AOD average over 9 years (2007 to 2015) of CALIOP data.

Table 2. Seasonal and annual daytime BL and FT AOD average over 9 years (2007 to 2015) of CALIOP data. Standard deviation is smaller than 0.01. Numbers in brackets show the contribution percentages of AOD.

<table>
<thead>
<tr>
<th></th>
<th>Land</th>
<th>Ocean</th>
<th>Global</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>BL</td>
<td>FT</td>
<td>BL</td>
</tr>
<tr>
<td>DJF</td>
<td>0.113 (71)</td>
<td>0.046 (29)</td>
<td>0.093 (71)</td>
</tr>
<tr>
<td>MAM</td>
<td>0.139 (73)</td>
<td>0.050 (27)</td>
<td>0.086 (65)</td>
</tr>
<tr>
<td>JJA</td>
<td>0.138 (68)</td>
<td>0.065 (32)</td>
<td>0.086 (64)</td>
</tr>
<tr>
<td>SON</td>
<td>0.126 (72)</td>
<td>0.050 (28)</td>
<td>0.085 (70)</td>
</tr>
<tr>
<td>Year</td>
<td>0.129 (71)</td>
<td>0.053 (29)</td>
<td>0.087 (68)</td>
</tr>
</tbody>
</table>

similar over land (71 and 29 %, respectively) and ocean (68 and 32 %, respectively).

Figures 3 and 4 show that the largest contribution of the FT to the total AOD is found in the polar regions (> 60 %) where aerosol extinction values are very low and near the
Table 3. Annual daytime average contribution of each aerosol type to total AOD in BL and FT, and fraction of each aerosol type present in the FT. Data are retrieved from CALIOP for the 2007–2015 time period. The standard deviation is less than 2 %.

<table>
<thead>
<tr>
<th>Aerosol compound</th>
<th>Contribution to total BL AOD (%)</th>
<th>Contribution to total FT AOD (%)</th>
<th>Fraction present in the FT (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Land</td>
<td>Ocean</td>
<td>Global</td>
</tr>
<tr>
<td>Marine</td>
<td>2</td>
<td>61</td>
<td>36</td>
</tr>
<tr>
<td>Dusty marine</td>
<td>2</td>
<td>29</td>
<td>18</td>
</tr>
<tr>
<td>Dust</td>
<td>43</td>
<td>5</td>
<td>21</td>
</tr>
<tr>
<td>Polluted dust</td>
<td>26</td>
<td>0</td>
<td>11</td>
</tr>
<tr>
<td>Polluted continental/smoke</td>
<td>22</td>
<td>4</td>
<td>11</td>
</tr>
<tr>
<td>Elevated smoke</td>
<td>5</td>
<td>1</td>
<td>3</td>
</tr>
</tbody>
</table>

Figure 3. Global and annual averaged daytime FT contribution to the total AOD for 9 years (2007 to 2015) of CALIOP data.

Figure 4. Latitudinal distribution of seasonally averaged daytime FT contribution to the total AOD for 9 years (2007 to 2015) of CALIOP data.

which has been extensively reported in the literature (e.g., Cui and Carslaw, 2006; Ekman et al., 2006; Weigel et al., 2011; Kipling et al., 2016).

Figure 5 shows the global average seasonal variation in AOD in BL and FT over land and ocean. The maximum BL and FT AOD is found in MAM and JJA, whereas the minimum occurs in DJF. Note that the last 3 years of the study (2013–2015) show lower BL AOD over land compared with the earlier years (2007–2012). According to our analysis of CALIOP observations, this is mostly due to large AOD decreases in northern Africa and eastern China during these years, which is in agreement with findings by Ridley et al. (2014) and Toth et al. (2016), respectively. As indicated by these studies, the AOD decrease in northern Africa may be due to a reduction in surface winds over the Saharan dust source regions and the AOD decrease in eastern China may be due to a decrease in aerosol loading after the 2008 Olympic Games. Overall, the seasonal cycle of the BL AOD over land is correlated with both the FT AOD over land and ocean with a $R^2$ correlation coefficient of 0.45. In contrast, no correlation is found with the BL AOD over ocean. This result suggests that FT aerosols found over land are often directly related to local aerosol sources while FT aerosols found over ocean are a result of long-range transport (e.g., long-range transport of dust aerosols over the Atlantic Ocean).

As described in Sect. 2.1, seven different aerosol types are discriminated with help of the AVD flag: marine, dusty marine (mixture of dust and marine particles), dust, polluted dust (meaning dust mixed with smoke or other non-depolarizing particles), clean continental, polluted continental/smoke and elevated smoke. In the following, clean continental aerosol data are not shown because they only contribute to 0.2 % of the global AOD (less than 1 % in terms of data fraction). The annual contribution of aerosol types in the BL and FT are summarized in Table 3. It should be noted that some of the CALIOP aerosol typing will lean toward the BL or FT by design. For instance, an aerosol layer will be classified as clean marine if the layer base is below 2.5 km, over ocean, strongly scattering and weakly depolarizing. Simi-
larly, the dusty marine classification occurs for layers with bases below 2.5 km over ocean and moderately depolarizing. The elevated smoke classification occurs for layers having tops above 2.5 km, whereas layers having similar measured optical properties but have tops below 2.5 km are classified as polluted continental/smoke. In Table 3, a tiny fraction of particles classified as clean marine are found over land because CALIOP data have been averaged over a 2 × 2° grid that can be partly land and ocean (see “Data and methods” section).

Overall, natural aerosols (marine, dusty marine and dust) contribute about 71% of the global AOD, while anthropogenic aerosols (polluted continental/smoke and elevated smoke) contribute about 16%. The remaining 13% of aerosols are a mix of natural and anthropogenic aerosols (polluted dust). Natural aerosols contribute more to the BL than FT AOD (75 versus 68%) while anthropogenic particles provide a larger contribution to the FT than BL AOD (14 versus 18%). However, within the natural aerosol category, the AOD contribution of pure dust is smaller in the BL than in the FT (21 versus 31%), while the AOD contribution of marine particles is much larger in the BL than in the FT (36 versus 20%). Overall, marine particles are the aerosol type that is the least efficiently transported into the FT. Indeed, 80% (20%) of the marine particles are found in the BL (FT). In contrast, 60% (40%) of the dust is found in the BL (FT). Smoke aerosols are also efficiently transported into the FT. Although their emission is much smaller than marine and dust aerosols, smoke particles contribute at least 10% of the FT AOD. This can partly be explained by the location of the emission sources (near the convective regions) and aerosol properties such as hygroscopicity, which is likely lower for smoke than sulfate-containing pollution (e.g., Carrico et al., 2010). Sea salt particles are hydrophilic and efficient cloud condensation nuclei. They are found in pristine areas with very few other aerosol sources, which makes them efficiently activated and removed by marine clouds. As a consequence, sea salt aerosols have a much shorter residence time than dust and smoke (less than a day versus several days) (e.g., Stier et al., 2005) and mostly remain in the BL. In contrast, dust and smoke are often emitted in dry convective regions (over desert and/or during the dry season), resulting in less efficient scavenging and more efficient vertical transport. It should be noted that while the AOD can provide a rough measure of total particulate mass, particle residence time and cloud interactions depend strongly on the particle size distribution. In agreement with Fig. 5, BL and FT AOD are correlated for each aerosol type over land ($R^2 \geq 0.7$ for dust, polluted continental/smoke and elevated smoke) but not correlated over ocean (not shown).

### 3.2 Nighttime analysis

During nighttime, the global contribution to the AOD from the BL and FT is 38 and 62%, respectively. In contrast to daytime, most of the AOD is found in the FT due to the low altitude of the BL height at night (about 200 m over land; see Sect. 2.2). However, if we consider the residual layer height (represented by the BL height of the previous day) instead of the nighttime BL defined by ERA-Interim, we find that the nighttime BL and residual layer both contribute about 62% of the total AOD. This is less than the relative contribution of the daytime BL (69%) to the AOD. However, in terms of absolute values, the daytime BL and the nighttime BL and residual layer AOD are similar. This indicates that the larger FT AOD during night than during day is likely due to the lower daytime signal-to-noise ratio of CALIOP measurements which can affect the fidelity of feature detection for weakly scattering layers, resulting in a potential daytime underestimate of the aerosol extinction and AOD in the FT (see Sect. 2). In terms of altitude, the vertical distribution of the AOD is similar during day and night with 58–63% of the global AOD average between 0 and 1 km, 21% between 1 and 2 km, 8–11% between 2 and 3 km, 4–6% between 3 and 4 km, 2% between 4 and 5 km, and less than 1% per kilometer above. This result implies that, overall, the vertical distribution of AOD does not follow the diurnal cycle of the
BL, corroborating the results of Kafle and Coulter (2013) and indicating that a large fraction of the aerosol mass remains in the residual layer during night, which may then even act as a source of aerosols to the BL forming during the following day (Sun et al., 2013; Curci et al., 2015). This feature can be explained by the dynamics related to the formation and evolution of the boundary and residual layers: the height of the BL is affected by the size of the largest turbulent eddies and thus the spatial scale of efficient mixing, which is in turn determined by the input of solar energy. As the BL height increases, emissions from the surface are mixed with air from higher altitudes, resulting in a well-mixed BL up to a few kilometers. In contrast, the aerosol concentration does not follow the decreasing BL height in the evening, but instead remains in the less dynamic residual layer, where the removal processes are much slower than within the BL.

3.3 Comparison with the LOAC in situ observations

Several studies showed that low aerosol extinction values are underestimated by CALIOP due to limitations in the detection of weakly scattering aerosols, leading to a potential underestimate of aerosols, particularly in the daytime due to solar noise. Kacenelenbogen et al. (2011) showed a case study in which the CALIOP retrieval underestimates an AOD plume of about 0.05–0.06 measured with other instruments by about 40%. Kim et al. (2017) found that undetected layer AOD accounts for about 0.031 (0.036 during day and 0.025 during night). Toth et al. (2018) concluded that CALIOP screened AOD values (i.e., those that do not pass the requirement tests) have a primary mode of 0.03–0.05. According to Rogers et al. (2014), AOD in the FT may be underestimated by 0.02. In terms of aerosol extinction, Winker et al. (2013) indicated that true aerosol extinction values of 0.001 km\(^{-1}\) or less are underestimated by CALIOP, while Sheridan et al. (2012) showed that 95% of aerosol extinction values larger than 0.05 km\(^{-1}\) are detected, 50% of aerosol extinction values lower than 0.02 km\(^{-1}\) are undetected and almost none of them are detected below 0.01 km\(^{-1}\). Although these studies used CALIOP v3 data, our results with CALIOP v4.1 data do not show a different behavior. Indeed, the study of Sheridan et al. (2012) has been repeated with v4.1 data and we arrived at the same conclusion with regards to extinction at the limits of CALIOP detection (i.e., an underestimate of the lowest aerosol extinction values). As a consequence, the global statistics of the study are very likely biased low, in particular in the free troposphere, as they are more likely to occur in the FT than in the BL (i.e., further away from sources). However, according to the studies previously cited, CALIOP underestimates the AOD by about 0.03–0.05, in which 0.02 would be due to the FT. This indicates that about half of the CALIOP underestimate is due to the BL and the other half to the FT. Therefore, the AOD contribution of the FT may be underestimated and could reach a global value larger than 31%.

4 Discussion and conclusion

This paper examines the vertical distribution of the aerosol optical depth (AOD) through its partitioning between the
boundary layer (BL) and free troposphere (FT). For this purpose, the CALIOP aerosol extinction product is combined with estimates of BL height from ERA-Interim over a 9-year period (2007–2015). The BL and FT contribute 69 and 31 %, respectively, of the global AOD during the day. It should be noted that although CALIOP data have often been reported as underestimated, in particular in the FT, in previous studies due to detection limitations, the case study over Aire sur l’Adour shows a good agreement between the LOAC in situ measurements and CALIOP data.

The daytime annual average FT contribution ranges from about 10 % at mid-latitudes to 50 % within the tropics, likely indicating a difference in vertical transport efficiency. Convection – occurring more frequently and strongly within the tropics – is often considered as the main transport pathway of chemical compounds from the surface to the FT. This process may be responsible for the difference in the FT contribution to the total AOD between the mid-latitudes and the tropics. The FT contribution to AOD is larger at northern mid-latitudes (25 %) than at southern mid-latitudes (13 %), which is very likely a consequence of the larger land fraction (and thus emission sources and convective activity) in the Northern than Southern Hemisphere. Continental particles, in particular dust and smoke, are the aerosol types transported the most efficiently to the FT (> 40 %), while marine particles mostly remain in the marine BL (80 %). This implies that continental aerosols in the FT can be transported over long distances towards remote/pristine regions, where they can act as a source of aerosols for the BL (Clarke et al., 2013; Bourgeois et al., 2015), impacting the climate far away from their sources.

In contrast to daytime, the BL and FT contribute 38 and 62 %, respectively, of the global AOD during the night. However, when considering the residual layer of the previous day, and including the residual layer in the BL contribution, the contribution of the BL and FT to the total AOD becomes 62 and 38 %, respectively. This indicates that a large fraction of the aerosol mass remains in the residual layer during the night (24 % in terms of AOD) which may then act as a source of aerosols to the BL forming the following day.

To conclude, this paper shows that a large fraction of the global AOD is found in the FT. This result may be crucial for the calculation of the radiative forcing at the top of the atmosphere in global climate models. Therefore, it is necessary to better constrain models in terms of AOD in the BL and FT. Furthermore, the distribution of aerosol particles between BL and FT affects their atmospheric residence time and thus the overall budgets of various particulate-phase pollutants. In other words, the results presented here have potential implications not only for calculating the radiative impact of aerosols but also for modeling of atmospheric chemistry, pollutant transport, air quality and aerosol–cloud interactions.

Code availability. All codes that have contributed to the results reported in this study are available on request.

Data availability. CALIOP data have been retrieved through the ICARE Data Services and Center (http://www.icare.univ-lille1.fr, last access: April 2018) and the ERA-Interim Archive is available on the ECMWF website (http://ecmwf.int/en/research/climate-reanalysis/era-interim, last access: September 2017).

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

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Atmos. Chem. Phys., 18, 7709–7720, 2018


The rest of the text continues with various references and discussions related to aerosol properties, models, and measurements in the boundary layer and free troposphere.


Atmos. Chem. Phys., 18, 7709–7720, 2018 www.atmos-chem-phys.net/18/7709/2018/