Overshooting of clean tropospheric air in the tropical lower stratosphere as seen by the CALIPSO lidar

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Abstract. The evolution of aerosols in the tropical upper troposphere/lower stratosphere between June 2006 and October 2009 is examined using the observations of the space borne CALIOP lidar aboard the CALIPSO satellite. Superimposed on several volcanic plumes and soot from an extreme biomass-burning event in 2009, the measurements reveal the existence of fast-cleansing episodes in the lower stratosphere to altitudes as high as 20 km. The cleansing of the layer, which extends from 14 to 20 km, takes place within 1 to 4 months during the southern tropics convective season that transports aerosol-poor tropospheric air into the lower stratosphere. In contrast, the convective season of the Northern Hemisphere summer shows an increase in the particle load at the tropopause consistent with a lofting of air rich with aerosols. These aerosols can consist of surface-derived material such as mineral dust and soot as well as liquid sulfate and organic particles. The flux of tropospheric air during the Southern Hemisphere convective season derived from CALIOP observations is shown to be 5 times at 16 km and 20 times at 19 km larger, respectively, than that associated with flux caused by slow ascent through radiative heating. These results suggest that convective overshooting is a major contributor to troposphere-to-stratosphere transport with concomitant implications for the Tropical Tropopause Layer top height, the humidity, the photochemistry and the thermal structure of the layer.

1 Introduction

Most natural and human-made short- and long-lived trace gases controlling stratospheric chemistry and radiative balance are emitted at the surface and rapidly lifted up in the tropics by convective systems. They reach the maximum outflow of convective systems near 14 km within a few hours (Gettelman and Forster, 2002), from where they are transported upward across the Tropical Tropopause Layer (TTL) into the stratosphere and distributed globally by the Brewer-Dobson circulation (Holton et al., 1995). Two mechanisms contribute to the transport across the TTL: a slow ascent on a time scale of 6 to 9 months up to 20 km by radiative heating of air masses (Holton, 1995; Corti et al., 2005; Yang et al., 2008; Fueglistaler et al., 2008) and fast convective overshooting updrafts above land thunderstorms as proposed by Danielsen (1982, 1993). Convection associated with the Asian monsoon during the Northern Hemisphere summer may also contribute to the latter mechanism bypassing tropical areas (Dunkerton, 1995; Gettelman et al., 2004; Randel et al., 2010; Fu et al., 2006).

Due to the time required to ascend through the TTL by slow radiative heating, very short-lived chemically active species should be deactivated and the air is expected to be dehydrated at the cold point tropopause. Although recognized to occur episodically (Danielsen, 1982, 1993), fast convective overshooting of adiabatically cooled air is generally believed to occur too infrequently to be significant in the overall mass flux (Fueglistaler et al., 2008 and references therein). However, recent observations adjacent to or above continental tropical convective systems during recent European balloon and high altitude aircraft campaigns,
HIBISCUS (Pommereau et al., 2011) and TROCCINOX (Chaboureau et al., 2007) in Brazil, SCOUT-O3 in northern Australia (Brunner et al., 2009) and SCOUT-AMMA in West Africa (Cairo et al., 2010), have shown that convective overshoots are more common than previously thought. They can reach altitudes as high as 19 km and thus strongly impact the thermal structure of the TTL (Pommereau et al., 2011; Cairo et al., 2010). Moreover, during these field campaigns, the air was not found to be dehydrated as often assumed, but instead associated with water vapor enhancements after evaporation of the injected ice crystals (Nielsen et al., 2007; Chaboureau et al., 2007; Corti et al., 2008; Khaykin et al., 2009), a process relatively well captured by cloud resolving models (CRM) (Jensen et al., 2007; Chaboureau et al., 2007; Grosvenor et al., 2007).

Although the frequent occurrence of such events over tropical continents is becoming more widely accepted, their contribution to troposphere-to-stratosphere transport at global scale is still debated. There are indications however that it might be more significant than previously thought. For instance, the Tropical Rainfall Measurements Mission—Precipitation Radar (TRMM-PR) indicates that at least 1 % of storm turrets reach the TTL (Liu and Zipser, 2005), particularly over Africa (Zipser et al., 2006; Liu and Zipser 2009). Since TRMM-PR are only sensitive to large hydrometeors produced in the core of deep convective systems, this percentage could be significantly underestimated.

Trace gases measurements in the UT/LS such as CH₄ concentration from HALOE, N₂O from ODIN-SMR, N₂O and CO from AURA-MLS, show maxima in the lower stratosphere over tropical continents (Ricaud et al., 2007; Schoeberl et al., 2006), indicative of fast vertical transport over these areas. Finally, the high-resolution aerosols profiles of the CALIOP lidar observations show an apparent aerosol cleansing of the TTL and lower stratosphere during the Northern Hemisphere winter that may be linked to convective overshooting (Vernier et al., 2009).

Except for infrequent strong volcanic eruptions (McCormick et al., 1995), the sources of aerosols in the tropical Upper Troposphere and Lower Stratosphere are poorly understood. In situ particle measurements have shown that the production of Aitken nuclei (r < 50 nm) is very high in the tropics compared to the mid-latitudes due to gas-to-particles conversion of SO₂ and other gas precursors (e.g. DMS) (Brock et al., 1995). High altitude aircraft particle composition and volatility measurements have indicated the presence of aerosols, largely made of a sulfate-organic mixture, and, to a smaller extent, black carbon and other carbonaceous materials (Froyd et al., 2009; Spackman et al., 2011; Bormann et al., 2010). But CALIPSO and SAGE II have also demonstrated that even small volcanic eruptions contribute significantly to the sulfate load of the tropical lower stratosphere (Vernier et al., 2011b). The lifetime of aerosols in the upper troposphere is controlled by convection by which they can be removed and precipitated to the ground (i.e. “washout”). Since at least 1 % of tropical convective systems reach the TTL (Liu and Zipser, 2005), the process might be an important sink of aerosols below the tropopause in this region (Hamill et al., 1997).

The objective of this paper is to further examine the CALIOP clean air events from the beginning of the mission in June 2006 until October 2009, and their possible relation with convection in the tropics. Section 2 provides a description of CALIOP aerosols observations in the UT/LS, followed in Sect. 3 by a comprehensive analysis of the vertical and latitudinal clean air propagation during the Northern Hemisphere (NH) fall-winter-spring in 2007–2008. This period is the optimum period as it is least affected by volcanic or fire events of any period in the CALIOP record. The origin of the clean air is investigated in Sect. 4. Section 5 provides an estimate of tropospheric air mass fluxes at the global scale derived from the observed cleansing compared to the contribution of slow ascent by radiative heating, and Sect. 6 discusses alternative microphysical and dynamical processes which could also contribute to the observations. The conclusions are summarized in Sect. 7.

2 Cleansing of the UT/LS as seen by CALIOP

The CALIPSO mission, a collaboration between the National Aeronautics and Space Administration (NASA) and the Centre National d’Etudes Spatiales (CNES), is dedicated to the study of clouds and aerosols from the troposphere to the stratosphere (Winker et al., 2010). As a part of the A-Train constellation, the satellite is in polar orbit at 705 km, providing measurements at 01:30 and 13:30 LT with a repeat cycle of 16 days. Since June 2006, the CALIOP lidar measures backscatter profiles at 532 nm and 1064 nm, and depolarization at 532 nm with a vertical resolution of around 200 m in the stratosphere (Hostetler et al., 2006). Because of the low signal-to-noise ratio on individual profiles, the detection of aerosols in the stratosphere requires averaging. A method for deriving the Scattering Ratio (SR) (Eq. 1) has been developed by Vernier et al. (2009). The procedure is based on averaging 532 nm nighttime-only measurements in one degree latitude bands (~300 profiles) and further arranged into a regular grid of 1° latitude × 2° longitude and 200 m in altitude for the CALIPSO 16-day repeat cycle. To rectify a known CALIOP calibration deficiency that is particularly important in the UT/LS, all data are corrected using the recalibration technique described in Vernier et al. (2009). Artifacts in the CALIOP SR time series created by sudden changes in the GEOS-5 temperature at 36–39 km, the aerosol free altitude layer used for adjusting the calibration, have been removed by replacing the GEOS-5 temperatures with those of the ECMWF ERA-INTERIM reanalysis found to better match the temperatures observed by MLS/Aura (Schwartz et al., 2008). In addition, a mask for the South Atlantic Anomaly (SAA) has been applied with which all 532 nm
profiles showing dark noise greater than 100 photons in this area have been discarded (Hunt et al., 2009). Finally, clouds below 20 km have been filtered on the same grid by removing all pixels for which the mean volume depolarization ratio \(\delta\) is greater than 5 % using Eq. (2).

\[
\text{SR} = \frac{\beta_{\text{part}} + \beta_{\text{mol}}}{\beta_{\text{mol}}} \quad \text{(1)}
\]

\[
\delta = \frac{\beta_{\|}}{\beta_{\perp}} \quad \text{(2)}
\]

where \(\beta_{\text{mol}}\) and \(\beta_{\text{part}}\) represent the molecular and particulate backscatter (note that the sum of both corresponds to the total backscatter as reported by the CALIOP lidar). \(\beta_{\perp}\) is the perpendicular backscatter at 532 nm measured by CALIOP which is sensitive to aspherical particles such as ice crystals. \(\beta_{\|}\) is the parallel backscatter at 532 nm.

Shown in Fig. 1 is the resulting evolution of the CALIOP zonal mean SR between 14–30 km within the 20° N–20° S tropical band (after removing the SAA) from the beginning of the mission in June 2006 until October 2009. This is complemented by Fig. 2, which represents the time/latitude cross sections of the same quantity at four potential temperature levels: 380 K (17 km), 420 K (18.5 km), 460 K (20 km) and 500 K (21.5 km). The blank section in February–March 2009 on both figures is due to an interruption of the CALIOP measurements.

Several volcanic events with a Volcanic Explosivity Index (VEI) of 4, listed in Table 1, can be seen in the tropics in Fig. 1 as well as at mid-latitudes (Fig. 2) with SR values greater than 1.2. The largest feature is associated with the eruption of the Soufrière Hills volcano (index 1 in Figs. 1 and 2) on Montserrat Island in the Caribbean on 20 May 2006. The plume remained in the tropics at 18–20 km (460 K) for 4–5 months before starting to ascend into the tropical stratosphere eventually reaching 25 km more than a year later. A second volcanic feature, less intense, is seen in October–December 2006 after the eruption of the Tavurvur (2) cone of the Rabaul volcano in Papua-New Guinea on 7 October. The lifetime of this plume is much shorter than the first one as it is rapidly transported to mid-latitudes at isentropic levels from 360 to 440 K (Fig. 2) and replaced by air of low aerosol concentration in the tropics. Other volcanic plumes can be seen during the summers 2008 and 2009 in the Northern Hemisphere (NH) after the eruption of Kasatochi (5) in Alaska (US) on 7 August 2008 (Bourassa et al., 2010) and Sarychev (7) in Kamchatka (Russia) on 12 June 2009. Both plumes are further transported to the northern tropics, but more rapidly at lower (380 K) than at higher levels (460 K) (Fig. 2), creating an apparent ascent in the tropics (Fig. 1). The remaining feature seen at 480–520 K (21 km) in March–April 2009 after the data gap (Figs. 1 and 2), with a SR between 1.10–1.20, is not the signature of a volcano but of soot particles (6) from an extreme biomass-burning episode near Melbourne in Victoria (southern Australia) on 7 February 2009 (Trepte et al., 2009).

In addition to these events, aerosol layers (index M in Fig. 2) are also present at 360–400 K from June to September, more intense in the Northern Hemisphere subtropics and at mid-latitudes, when no volcanic events could be identified in 2006 and 2008. These aerosol layers appear every year during the Asian monsoon period, extending from Western Asia to Eastern Mediterranean (Vernier et al., 2011a). The monthly mean latitude-height cross-sections between June and September in 2007 over Asia (not shown), indicates that they do not come from an old volcanic plume transported southward across the weak barrier afforded by the NH summer jet stream (Chen, 1995; Dunkerton, 1995), but rather from the deep convective outflow of the Asian monsoon.

Besides those features, the “clean air” events are particularly noteworthy. These events have a scattering ratio less than 1.04 that form near the base of the TTL (~14 km) and propagate rapidly upward. Since the CALIOP lidar at 532 nm
is primarily sensitive to particles with radii larger than 30 nm, a “clean air” event refers to air masses with low densities of such particles. The first event in early 2007 can be seen from 14 to 18 km (360–440 K), more pronounced in the tropics and that rapidly replaced the Tavurvur plume. A similar episode occurred during the same season in 2008, a year undisturbed by volcanoes and fires, the lower stratosphere being cleansed up to 19–20 km and remained relatively aerosol-free until the following year between 18 and 21 km (440–520 K). The vertical propagation of the clean air during the NH winter 2007–2008 is compared in Fig. 2 to the theoretical ascent of the lowest bulb of clean air from 380 K to 500 K using Yang et al. (2008) estimations. The observed propagation is much more rapid than the one deduced from the radiative ascent model.

Though of higher resolution and free of clouds interference in the CALIOP data, the cleansing of the lower stratosphere during the NH winter-spring, resulting in an aerosol annual cycle, was observed by SAGE II at 18 km between 1998–2005 (Thomason et al., 2008). It is also confirmed by independent balloon measurements in the tropics within the SCOUT-O3 project shown in Fig. 3 displaying: (i) the SR derived from the particle size distribution of an optical particle counter (OPC, Deshler et al., 2003) during a flight in May 2008 from Teresina in Brazil at 5° S, (ii) the mean SR profile measured by backscatter sondes (BKS, Rosen and Kjome, 1991) flown in September 2008 at 12° N in Niamey, Niger. Both are compared to the mean CALIOP SR within a box of ±7° latitude ±70° longitude centered at the location of the balloon flight.

Fig. 2. Latitude-time cross-sections of CALIOP SR within 40 K (1.5 km) thick layers centred at 380 K (17 km), 420 K (18.5 km), 460 K (20 km) and 500 K (21.5 km) from June 2006 until October 2009. The black dotted curve shows the timing of the uplift of clean air if it was due to slow ascent by radiative heating as calculated by Yang et al. (2008). Letter M corresponds to the aerosol feature associated with the Asian monsoon.

Fig. 3. Left: SR (red) derived from the aerosol size distribution of a balloon-borne Optical Particle Counter at 5° S in Brazil in May 2008; right: mean scattering ratio reported by BKS sondes at 12° N in Niger in September 2008; both compared to CALIOP SR within a box of ±7° latitude ±70° longitude centred at the location of the balloon flight.
Fig. 4. Latitude-height cross-sections of mean CALIOP SR from September 2007 to August 2008 every two months. Shown by white lines are temperature potential levels at 380, 420, 460 and 500 K and by red lines the mean cloud top height (zonal mean depolarization ratio = 5%).

3 Location and vertical propagation of the cleansing

Since the low signal-to-noise ratio of CALIOP does not allow constructing SR maps between 15 and 20 km with a temporal resolution of less than 2 months, and the Quasi Biennial Oscillation (QBO) was in its easterly phase during the winter 2008 with zonal wind speed of around $-15 \text{ m s}^{-1}$ at 50 hPa (www.cpc.ncep.noaa.gov/data/indices/qbo.u50.index), an air mass in the tropical lower stratosphere was circumnavigating twice during this two-month period, making it difficult to determine a possible longitudinal dependence of the cleansing.

We then choose to look at two-month cross-sections shown in Fig. 2 for October 2007 through September 2008 annotated with the mean zonal cloud top in red in Fig. 4. The zonal cloud top is determined using the mean zonal depolarization ratio ($\delta$), derived from the statistics of cloudy/aerosols pixels separation as described in Sect. 2. From this figure, it can be seen that the cleansing of the UTLS starts in October/November 2007 and continues in December/January with SR values falling below 1.04 between 14 km to 17 km. The low aerosol region propagates between 20° S–5° N in phase with the latitude of cloud tops. The cleanest period occurs in February–March where low SR values reach altitudes as high as 20 km, 2–3 km higher than cloud tops, making the whole TTL and the lower stratosphere virtually free of aerosol. Later in April–May, an isolated
A bulb of clean air is seen at 18–20 km between 10°N–20°S above cloud tops, which remains at this level until September 2008. Also seen in April–May is the presence of an aerosol layer at 16–17 km, between 10–30°N, reinforced and extended to higher latitude in June–July, before the arrival of the Kasatochi plume.

To shed light on the relationship between cleansing episodes and deep convection, Fig. 5 shows the latitudinal evolution of the SR between 14–17 km with the location of the Inter Tropical Convergence Zone (ITCZ) at 14 km as inferred from the TRMM Mesoscale Convective Systems (MCS) position (red) and the CALIPSO cloud cover (white). White arrows show the mean meridional wind every 16 days derived from ECMWF at 100 hPa. The TRMM MCSs are selected by considering contiguous precipitating areas greater than 2000 km² of radar signal larger than 20 dBZ reaching at least 14 km, with a minimum of 2 flashes as observed by the Lightning Imaging Sensor (LIS) also on TRMM. The flash discrimination is chosen to select only MCSs with strong updrafts inducing hail formation and lightning. All MCSs are extracted with those four conditions from the TRMM database (http://trmm.chpc.utah.edu/) from June 2006 to February 2009 and sorted into 16-day bins comparable to CALIPSO. The limits of the ITCZ (red line) from these observations are derived using the 10th and 90th percentile of latitudinal position of MCSs. The CALIPSO cloud area is derived from the depolarization channel as described above (Fig. 4). As above, the bounds of clouds are defined by the 10th and 90th percentile latitudinal location of cloudy pixels. The term “cloudy pixels” represents all types of clouds observed by CALIPSO including storms, anvils and cirrus. During the NH winter, the clouds extend to higher latitude than the ITCZ defined by the MCSs location likely due to the persistent thick cirrus cloud layer up to 10°N in the West Pacific where the tropopause is the highest and coldest (Sassen et al., 2009). In contrast, the ITCZ area extends to higher latitudes than the clouds at 14–17 km in the summer of both hemispheres and particularly in the North. This is likely due to the less frequent clouds during CALIPSO overpasses at 01:40 a.m. and p.m. over land convective regions (monsoon in the north) whereas cloud cover exhibits a large diurnal cycle with a strong maximum around 16:00–17:00 h local (Liu and Zipser, 2005).

Aside from volcanic plumes during the NH winter 2006 (Tavurvur, 0–20°S) and the NH summers of 2008 and 2009 (Kasatochi, Sarychev, 20–50°N), the aerosols in the TTL show an annual cycle. The maximum cleansing of the 14–17 km layer is located in the middle of the ITCZ area in the Southern Hemisphere (5°S–25°S) during the northern winter but extends further north and south than the limits of the ITCZ area. As shown by the meridional wind arrows, this altitude level corresponds to the upper branch of the trade wind Hadley cell which is associated with the advection of air masses from the ITCZ to the subtropics where there is no more convection. In contrast, it is the opposite in the Northern Hemisphere summer, where a maximum aerosols load is observed between 20–40°N in the northern part of the ITCZ area. Thus, the impact of convection on the TTL aerosols load may depends on the amount of aerosols available at the surface.
Fig. 6. Bimonthly maps from October 2007 to September 2008 of TRMM Mesoscale Convective Systems (MCS > 14 km) tagged with MODIS Aerosol Optical Depth. The nearest AOD value of each MCS was extracted from MODIS AOD grids with a temporal and spatial resolution of 2 days and 3° lon × 3° lat. MCS that could not be tagged where no AOD value was available are in black. The AOD fields have been constructed using the MODIS Level 3 daily joint aerosol/water vapor/cloud product of the Terra space borne platform that cross the equator at 10:30 a.m. LT.

4 Modulation of the aerosol loading in the TTL by deep convection

As previously shown, the cleansing of the lower stratosphere is correlated with the southern tropics convective season, but if not anticorrelated with the Northern Hemisphere season. We evaluate the convective transport of low versus high aerosol loading of tropospheric air masses in the TTL by studying the geographical position of MCSs as seen by the TRMM radar relative to the amount of aerosols in the troposphere. Since deep convection is known to show a pronounced diurnal cycle of peak intensity in late afternoon (Liu and Zipser, 2005), we have used the Aerosol Optical Depth (AOD) from MODIS on the TERRA satellite platform at 10:30 a.m. LT. Figure 6 represents bimonthly maps from October 2007 to September 2008 of MCS locations reaching at least 14 km tagged with the nearest value of MODIS AOD in space and time. Since more than one day of observations is required to obtain a complete spatial coverage, two days mean AOD maps have been constructed with a spatial resolution of 3° lon × 3° lat from the MODIS level 3 daily joint aerosol/water vapor and cloud product. As shown by Liu and Zipser (2005), the maps indicate that deep convection is dominant over land during the local summer convective season. The statistics of tagged-AOD MCSs is given in Table 2 for the different maps of Fig. 6. In October–November, the maximum of potential aerosols lofting occurs over East Amazonia next to the arid N-E Brazil, where a maximum CO in the lower stratosphere was also reported by Schoeberl et al. (2006). In December–January 2008, the convective activity is greatest above the continents of the Southern Hemisphere, in South America, South Africa, Indonesia and northern Australia. MCS occurred in relative clean environment (AOD < 0.4) 73 % of the time except in Central Africa where convection coincides with the beginning of biomass burning season as shown by the number of fires associated with CO emission (http://earthobservatory.nasa.gov/GlobalMaps/view.php?d1=MODAL2_M_AER_OD,&d2=MOD14A1_M_FIRE).

Later in the winter season (February–March 2008), the ITCZ and land convection have moved northward but still remain in clean tropospheric conditions with 76 % of MCS with AOD < 0.4. In early spring, in April–May, the aerosol loading associated with convection increases (MCS of AOD > 0.4) by 10 % compared to previous periods. This polluted convection occurs primarily in Western and Central
African regions of biomass burning and Saharan dust, but also in South East Asia when the monsoon onset is reaching the Bay of Bengal and Thailand. The highest level of aerosol associated with convection occurs during the NH summer (June–July 2008 and August–September 2008), where 25% of MCSs are associated with AOD > 0.4 and 65% of MCS tagged with extreme AOD values (>0.6) are located over Southeast Asia (NE China and NW India). As shown by Dey et al. (2010), polluted aerosols with dust dominate the population of particles especially in northeast India and along the Indo-Ganges basin. Deep convection occurring over those regions could transport aerosols upwards, confirming the possible convective origin of an elevated layer observed during the Asian monsoon reported by Vernier et al. (2011a).

To better quantify the impact of convection on the TTL aerosol load, we have created an Aerosol Index (AI, Eq. 3) that is the average of the tagged MCS-AOD occurring within a 16-days period in a particular latitude band. It is given by

\[
AI(t) = \frac{1}{N} \sum_{n=1}^{N} \text{MCS}(n,t) \times \text{AOD}(n,t),
\]

where \( n \) is the index of each MCS (\( N \) the total) occurring in the time and space resolution as described below. The AI has been computed from TRMM and CloudSat MCSs extracted for the later one from the University of Utah database (http://trmm.chpc.utah.edu) by selecting convective events reaching 14 km of at least 500 km horizontal extension. Figure 7 shows a comparison between this index and the mean CALIOP SR between 14 and 17 km. To emphasize the effect of deep convection on the TTL aerosols, we have calculated the mean CALIOP SR within the ITCZ from the 10th and 90th percentile CALIPO cloudy pixels (latitude band within the white line in Fig. 5). Since volcanic plumes from Tavurvur and Sarychev contaminate the years of 2006 and 2009, respectively (Figs. 1, 2 and 5), we focus on the 2007–2008 period. The annual cycle of TTL aerosols from CALIPSO is well matched by the AI deep convection-AOD model without significant phase lag using TRMM as well as CloudSat. However, the strong summer time CALIPSO SR peak is not seen on the AI that rather shows a slight maximum expending from April to November.

The relationship between AI and CALIPSO SR is reasonable, indicating that air masses transported by deep convection from the relatively clean surfaces of the southern tropics (Congo, Amazonia and Indonesian Islands) may be indeed responsible for the cleansing of the TTL during the NH winter. The contrast between the convective cleansing observed in the South and the reinforcement of aerosols in the TTL in the North is consistent with the increased AI during the NH summer by convective lofting of natural and anthropogenic aerosols during the Asian and African monsoon seasons (Fig. 7).

However, the rapid cleansing of the TTL up to 19–20 km during the SH convective season shows significantly higher vertical propagation than of the aerosol enhancement in the NH convective season that is limited to 16–17 km. This might be related to the influence of the aerosols on convection intensity. Rosenfeld et al. (2008) suggested that the vigorous convection as inferred from the Convective Available Potential Energy is maximum with AOD close to 0.25. This value is remarkably close to the lowest AI values reach (that represent also a mean AOD) in the Southern tropics during the NH winter when the cleansing is more intense, reaching as high as 19–20 km.

Table 2. Percentage (number) of Mesoscale Convective Systems (MCS) that was tagged with: AOD < 0.2 (column 1), 0.2 < AOD < 0.4 (column 2) and AOD >= 0.4 (column 3). The last column shows the percentage of events that could not be tagged.

<table>
<thead>
<tr>
<th>Period/AOD</th>
<th>AOD &lt; 0.2</th>
<th>0.2 &lt; AOD &lt; 0.4</th>
<th>AOD &gt;= 0.4</th>
<th>NA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oct–Nov-07</td>
<td>34% (377MCS)</td>
<td>39 (342)</td>
<td>14 (157)</td>
<td>11 (128)</td>
</tr>
<tr>
<td>Dec–Jan-08</td>
<td>34 (362)</td>
<td>42 (445)</td>
<td>9 (98)</td>
<td>12 (134)</td>
</tr>
<tr>
<td>Feb–Mar-08</td>
<td>35 (365)</td>
<td>38 (392)</td>
<td>13 (141)</td>
<td>12 (131)</td>
</tr>
<tr>
<td>Apr–May-08</td>
<td>31 (333)</td>
<td>36 (383)</td>
<td>21 (226)</td>
<td>10 (111)</td>
</tr>
<tr>
<td>Jun–Jul-08</td>
<td>29 (325)</td>
<td>30 (341)</td>
<td>25 (289)</td>
<td>14 (165)</td>
</tr>
<tr>
<td>Aug–Sep-08</td>
<td>31 (354)</td>
<td>33 (371)</td>
<td>24 (276)</td>
<td>10 (121)</td>
</tr>
</tbody>
</table>

5 Impact of overshooting updrafts at the global scale

The mass flux of tropospheric air required for explaining the cleansing of the TTL during the SH convective season was tentatively derived from the drop of aerosol observed during the period November 2007–February 2008 (Fig. 5). Figure 8 illustrates the conceptual model used to calculate the time constant of dilution and the total mass flux between the troposphere and the stratosphere corresponding to the updrafts. As detailed in Appendix A, a mass conservation flux is applied to the rectangle on the top of the anvil to estimate the amount of clean tropospheric air required at a level \( z \) to cleanse up the lower stratosphere. Assuming a constant size distribution and similar composition for the aerosols populating the UTLS, the CALIOP (SR-1) (SR-1 = \( \beta_{\text{part}}/\beta_{\text{mol}} \) can
Table 3. Average mass flux in $10^8$ kg s$^{-1}$ and time constant ($\tau$) in month, required for a (SR-1) decrease by 63% in the TTL between 20°S–0°S between 15 km–20 km. Values are derived from the overshoot dilution model (Appendix A) and compared to radiative heating calculations by Corti et al. (2005) on yearly average basis and by Yang et al. (2008) during the NH winter only.

<table>
<thead>
<tr>
<th>Altitude</th>
<th>Mass flux from CALIPSO ($10^8$ kg s$^{-1}$)</th>
<th>From Corti et al. (2005)</th>
<th>From Yang et al. (2008)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15 km</td>
<td>780 ± 390 ($\tau = 1.5 \pm 0.5$ months)</td>
<td>139</td>
<td></td>
</tr>
<tr>
<td>16 km</td>
<td>670 ± 340 ($\tau = 1.5 \pm 0.5$)</td>
<td>102</td>
<td>56</td>
</tr>
<tr>
<td>17 km</td>
<td>430 ± 160 ($\tau = 2 \pm 0.5$)</td>
<td>65</td>
<td>74</td>
</tr>
<tr>
<td>18 km</td>
<td>240 ± 60 ($\tau = 3 \pm 0.5$)</td>
<td>19</td>
<td>28</td>
</tr>
<tr>
<td>19 km</td>
<td>200 ± 50 ($\tau = 3 \pm 0.5$)</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>20 km</td>
<td>120 ± 20 ($\tau = 4 \pm 0.5$)</td>
<td>9</td>
<td>9</td>
</tr>
</tbody>
</table>

be considered to be proportional to an aerosol mixing ratio. After applying the mass and aerosol mixing ratio flux conservation, the aerosol mixing ratio is shown to follow an e-folding law, where $t$, the time constant required for (SR-1) to decrease by 63%, is directly proportional to the mass flux. The time $t$ required for such reduction in CALIOP observations ranges from 1.5 ± 0.5 months at 15 km, to 3 ± 0.5 months at 18 km, and 4 ± 0.5 months at 20 km (Table 3). The fluxes corresponding to the cleansing of each layer at global scale shown in Table 3, are $780 \pm 390 \times 10^8$ kg s$^{-1}$ at 15 km and $240 \pm 60 \times 10^8$ kg s$^{-1}$ at 18 km. Below 18 km, they are 5 to 12 times larger than those derived from radiative heating calculations by Corti et al. (2005) and Yang et al. (2008). At the higher levels, at 18–20 km, they are even larger (up 20 times), showing that convective overshooting could be a large contributor to the cleansing of the lower tropical stratosphere.

There are, however, some limitations in the validity of the method used in the above calculations. The conversion of the Scattering Ratio into an aerosol-mixing ratio required a constant phase function in time and altitude meaning that aerosol composition and size distribution remained the same. However, even if, as measured by Froyd et al. (2009) over Costa Rica, the aerosols above the tropopause are for 75% made of sulphate-organic particles, 20% of them were of carbonaceous material. Since organics particles are known to be more absorbing than scattering, this would imply that the phase function used in the conversion would not be conservative below the tropopause. In addition the calculation applies only if aerosols are long lifetime tracers, insensitive to changes by microphysical processes. But since sulfate-organic particles are dominants in the convective SH region and microphysical processes have little impact on these as discussed below, the above limitations are expected to have little impact on the conclusions.

![Fig. 7. CALIPSO mean SR between 14–17 km (black line) within the “ITCZ area” delimited by the 10th and 90th percentile of the CALIPSO cloudy pixel product derived from the depolarization (Fig. 5). The Aerosol Indices are derived from TRMM (red solid line) and CloudSat (dotted red line) MCSs located between 20°N–20°S.]

6 Discussion

Although there is clear indication of the surface origin of the clean air and its transport in the lower stratosphere by convective updraft, other processes could potentially contribute for explaining them and the difference between the northern and Southern Hemispheres. They are briefly discussed below.

6.1 Cleansing mechanisms

Several mechanisms can be invoked which could potentially contribute to the cleansing of the aerosols in the TTL: (i) a decrease of the particles size by microphysical processes making them less efficient to scattering light; (ii) the capture of the aerosols by cirrus clouds followed by sedimentation; and (iii) a fast sedimentation of the particles.

6.1.1 Particle size reduction

A significant fraction of particles in the TTL seems to be composed of sulfate-organic mixing (Froyd et al., 2009).
Fig. 8. Schematic of the TTL aerosol-cleansing model by overshooting towers.

has been shown by Steel and Hamill (1981) that sulfate particles can grow significantly in the presence of water vapor at low temperature and thus result in SR-1 enhancements. The amplitude of the effect depends on the initial concentration of the particles and the amount of nitric acid (HNO₃) available. Assuming an initial concentration of sulfate of 10 particles cm⁻³ and using water vapor, temperature and HNO₃ observations from MLS/Aura, the Mie SR-1 increase was shown to not exceed 10% in the worst case, mainly because of the very low HNO₃ concentration near the tropical tropopause. The change in retrieved aerosols profiles is not significant.

6.1.2 Capture by cirrus clouds

Another process could be the capture of aerosol particles in cirrus clouds. Indeed, if aerosols served as condensation nuclei for cirrus formation, they could be removed by the further sedimentation of cirrus ice crystals. However, as shown by high altitude aircraft sampling above Costa-Rica during the Pre-AVE campaign (Froyd et al., 2009), the aerosols above the tropopause associated to Amazonian convective systems are mainly sulfate-organic particles of <0.1 µm diameter resulting in very limited freezing (Froyd et al., 2010). However, this could be different in the Northern Hemisphere, since, in contrast, mineral dust and black carbon particles can act as efficient heterogeneous ice nuclei forming ice crystals (Froyd et al., 2010 and references therein). In conclusion, if such process contributes to the 16–17 km altitude limitation of the aerosols in Northern Hemisphere, it cannot explain the cleansing in the Southern Hemisphere during the convective season.

6.1.3 Aerosol sedimentation

Sulfate or sulfate-organic particles of 0.1–0.2 µm radius, as observed in the tropical lower stratosphere by the optical particle counter over Northern Brazil (Fig. 3) or Costa-Rica (Froyd et al., 2010), are falling at a rate of less than 0.1 km month⁻¹ at 18 km altitude. About 40 months would be required for them to drop from 18 to 14 km, which is far too slow for explaining the cleansing. However, this is not true for larger mineral dust particles of ∼1 µm diameter above the Asian sector (Anderson et al., 2003), which will be falling by 1 km month⁻¹ at 18 km, and thus could explain the limited altitude of the aerosols layer in the Northern Hemisphere during the monsoon season.

In conclusion, apart from the freezing and the sedimentation of large mineral dust particles which might explain the absence of aerosols enhancements above 17 km during the Northern Hemisphere summer, there is no indication that the above processes could contribute significantly to the cleansing of the lower stratosphere in the Southern Hemisphere.

6.2 Vertical transport processes

Aside from deep convective updraft, other potential contributing processes include slow ascent by radiative heating, vertical turbulent diffusivity and gravity waves. As already shown, the ascent by radiative heating at a rate of less than 0.4 mm s⁻¹ or 1 km month⁻¹ (Yang et al., 2008) will require 6–9 months for an air mass to reach 20 km from the tropopause, that is far too slow for explaining the observations. It cannot contribute for more than 10% to the zonal mean flux. The same conclusion applies to the average vertical turbulent diffusivity of 0.02 m² s⁻¹ derived by Mote et al. (1998) from the total hydrogen profiles of the Halogen Occultation Experiment which would require about 5 yr for a one kilometre rise. Even with the upper bound equivalent diffusivity of 0.5 m² s⁻¹ derived by Pinso and Legras (2008) from balloon ozone and water vapor profiles measurements during the HIBISCUS campaign in Brazil, thus including turbulent diffusivity and gravity waves, it would take 2.5 months for a one-kilometer uplift. The only remaining explanation is thus frequent injections of clean tropospheric air by convective overshooting that would progressively dilute the aerosols and clean the lower stratosphere. Since the overshooting frequency is known to decrease at increasing altitude (Liu and Zipser, 2005), the amplitude of dilution will also decrease with altitude, which is what is observed. This scheme is consistent with the observations of geyser-like injections of ice particles in the lower tropical stratosphere up to 19 km over land convective regions (Nielsen et al., 2007; Corti et al., 2008; Khaykin et al., 2009) implying a hydration process of the lower stratosphere, as well as the injection of adiabatically cooled air modifying its thermal structure (Pommereau et al., 2011). It would also be consistent with the larger N₂O, CH₄ and CO concentrations in the lower stratosphere over.
tropical continents in contrast to oceans, as well as their seasonal variation reported by Ricaud et al. (2007, 2009) or Schoeberl et al. (2008). It will imply a top TTL, the region sharing tropospheric and stratospheric characteristics, around 20 km, higher than generally assumed (Fueglistaler et al., 2008). Finally, it will result in the fast transport of chemically reactive Very Short-Lived Species (VSLs) in the lower stratosphere with their possible impact on ozone chemistry. However, if such deep overshooting events and the resulting hydration of the lower stratosphere are well captured by non-hydrostatic meso-scale model simulations (e.g. Chaboureau et al., 2007; Grosvenor et al., 2007; Jenessen et al., 2007), they are poorly, if at all, represented by global Chemistry Transport Models (CTM) forced by Numerical Weather Forecast (NWP) Models, enable to reproduce the impact of such events at global scale.

7 Conclusions

The three-year CALIOP aerosols profiles in the tropics reveal the existence of a fast cleansing process of the lower stratosphere up to 20 km. It is shown to take place in the Southern Hemisphere during the convective season, where overshooting systems seen by the TRMM precipitation radar are occurring in clean regions of the troposphere over African, South American rain forests and the Indonesian islands, as shown by the minimum MODIS Aerosols Optical Depth over these areas.

Such cleansing would be consistent with repeated injections of relatively clean air by overshooting updrafts in the air masses circumnavigating within less than two months in the tropical lower stratosphere, resulting in the progressive dilution of aerosols present there. The tropospheric air flux by those events might be 5–20 times larger than that corresponding to the previously assumed slow ascent by radiative heating. The fast extension of the cleansing to the northern tropics and the southern sub tropical lower stratosphere during the same season is consistent with the advection of the clean air injected at this level in the ITCZ region by the upper branch the Hadley cell. In contrast to the cleansing during the Southern Hemisphere summer, the aerosol increase observed in the Northern Hemisphere during the convective season, consistent with the large MODIS AOD within 10°–40° N over South Asia, suggesting instead, an injection of desert mineral dust and soot from biomass burning, but limited to around 16–17 km probably because of their fast sedimentation at higher altitude. The CALIPSO aerosols observations suggest that convective overshooting might be a major contributor of troposphere-to-stratosphere transport, with the many implications on the TTL top height, hydration, photochemistry and thermal structure of the lower stratosphere, a mechanism well captured by non-hydrostatic meso-scale models but missed by global meteorological and climatic models.

Appendix A

Overshoot updrafts mass flux

As shown by the conceptual overshoot model in Fig. 8, the mass flux conservation in the TTL can be written:

\[ D_c - E_c + D_{\text{ext}} - E_{\text{ext}} = 0 \]  

(A1)

where,

- \( D_c \) is the convective flow (\( D \) for Detrainment) (unit kg m\(^{-1}\) s\(^{-1}\))
- \( E_c \) is that extracted (\( E \) for Entrainment)
- \( D_{\text{ext}} \) is the input from horizontal transport and
- \( E_{\text{ext}} \) the horizontal outflow.

Neglecting the weak transport from mid-latitudes to the tropics allows simplifying Eq. (A1) in:

\[ D_c - E = 0 \quad (\text{with} \ E = E_c - E_{\text{ext}}) \]  

(A2)

The change of aerosol mixing ratio, \( \chi \), due to the mixing with clean air can be expressed:

\[ \rho \cdot A \cdot \frac{\partial \chi}{\partial t} = D_c \cdot \chi_c - E \cdot \chi \]  

(A3)

where \( \rho \) is the air density, \( A \) the surface of the layer, and \( \chi_c \) the aerosol mixing ratio in the convective cloud. The solution of Eq. (A3) is:

\[ \chi(t) = (\chi_0 - \chi_c) e^{\frac{D_c}{\rho A} t} + \chi_c \]  

(A4)

where \( \chi_0 \) is the initial aerosol mixing ratio in the layer and \( \chi(t) \) its change following an e-folding law, with \( \tau = \rho A / D_c \) assuming that the aerosol-mixing ratio in the convective cloud is negligible (clean air, \( \chi_c = 0 \)).

Since, \( \tau \) is the time required for reducing the aerosol mixing ratio by 63 %, the inflow \( F_c \) within each 1 km thick layer between 15–20 km can be expressed:

\[ F_c = \frac{D_c \cdot \Delta Z}{\tau} = \frac{\Delta Z \rho A}{\tau} \]

Assuming that the CALIPSO backscatter ratio (SR-1 = \( \frac{\beta_{\text{Ray}}}{\beta_{\text{mie}}} \)) is proportional to the aerosol mixing ratio. The tropospheric air flux required for cleansing the TTL can be derived from the decrease of CALIOP mean SR from November 2007 to February 2008 between 20° S–0° S shown in Fig. 5 using Eq. (A4). The resulting fluxes are displayed in Table 1.

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