Atmos. Chem. Phys., 14, 11687–11696, 2014 www.atmos-chem-phys.net/14/11687/2014/ doi:10.5194/acp-14-11687-2014 © Author(s) 2014. CC Attribution 3.0 License.





# Lidar-observed enhancement of aerosols in the upper troposphere and lower stratosphere over the Tibetan Plateau induced by the Nabro volcano eruption

Q. S. He<sup>1</sup>, C. C. Li<sup>2</sup>, J. Z. Ma<sup>3</sup>, H. Q. Wang<sup>4</sup>, X. L. Yan<sup>3</sup>, J. Lu<sup>5</sup>, Z. R. Liang<sup>1</sup>, and G. M. Qi<sup>6</sup>

<sup>1</sup>Shanghai Meteorological Service, Shanghai, China

<sup>2</sup>Department of Atmospheric and Oceanic Sciences, School of Physics, Peking University, Beijing, China

<sup>3</sup>Chinese Academy of Meteorological Sciences, Beijing, China

<sup>4</sup>College of Environmental Science and Engineering, Donghua University, Shanghai, China

<sup>5</sup>School of Computer and Software, Nanjing University of Information Science and Technology, Nanjing, China <sup>6</sup>Germu Meteorological Bureau, Qinghai, China

Correspondence to: C. C. Li (ccli@pku.edu.cn)

Received: 6 November 2013 – Published in Atmos. Chem. Phys. Discuss.: 31 January 2014 Revised: 24 August 2014 – Accepted: 24 September 2014 – Published: 6 November 2014

Abstract. Vertical profiles of aerosol extinction coefficients were measured by a micro-pulse lidar at Naqu (31.5° N, 92.1° E; 4508 m a.m.s.l.), a meteorological station located on the central part of the Tibetan Plateau during summer 2011. Observations show a persistent maximum in aerosol extinction coefficients in the upper troposphere-lower stratosphere (UTLS). These aerosol layers were generally located at an altitude of 18–19 km a.m.s.l., 1–2 km higher than the tropopause, with broad layer depth ranging approximately 3-4 km and scattering ratio of 4-9. Daily averaged aerosol optical depths (AODs) of the enhanced aerosol layers in UTLS over the Tibetan Plateau varied from 0.007 to 0.030, in agreement with globally averaged levels of  $0.018 \pm 0.009$  at 532 nm from previous observations, but the percentage contributions of the enhanced aerosol layers to the total AOD over the Tibetan Plateau are higher than those observed elsewhere. The aerosol layers in UTLS wore off gradually with the reducing intensity of the Asian monsoon over the Tibetan Plateau at the end of August. The eruption of Nabro volcano on 13 June 2011 is considered an important factor to explain the enhancement of tropopause aerosols observed this summer over the Tibetan Plateau.

## 1 Introduction

Aerosols in the upper troposphere–lower stratosphere (UTLS) play an important role in the global/regional climate system and the geochemical cycle (Hanson et al., 1994; Borrmann et al., 1997; Solomon et al., 1997). They also influence atmospheric ozone budgets through providing surface areas for efficient heterogeneous reactions (Keim et al., 1996; Solomon, 1999).

A volcanic eruption, though as an occasional event, can inject amounts of ash and sulfur dioxide (SO<sub>2</sub>) into the stratosphere, and the injected SO<sub>2</sub> is oxidized to sulfuric acid particles through homogeneous nucleation (Wu et al., 1994). The Nabro stratovolcano in Eritrea, northeastern Africa, erupted on 13 June 2011, injecting approximately 1.3 Tg of SO<sub>2</sub> to altitudes of 9 to 14 km in the upper troposphere, which resulted in a large aerosol enhancement in the stratosphere (Bourassa et al., 2012). This event has been observed by lidar networks such as EARLINET, MPLNET and the Network for the Detection of Atmospheric Composition Change (NDACC) with independent lidar groups and satellite Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) to track the evolution of the stratospheric aerosol layer in various parts of the globe (Uchino et al., 2012; Sawamura et al., 2013), and other instruments, such as the Infrared Atmospheric Sounding Interferometer (Clarisse et al., 2014) and the ground-based spectrometry of twilight sky brightness (Mateshvili et al., 2013). Bourassa et al. (2012) found that the aerosol enhancement developed while remaining confined for several weeks to the region between central Asia and the Middle East after eruption of Nabro volcano using the limb scanning Optical Spectrograph and Infra-Red Imaging System (OSIRIS) satellite instrument.

It should also be noted that aerosol enhancements in UTLS over the Tibetan Plateau have already been observed by many researchers before eruption of Nabro volcano. Using the Stratospheric Aerosol and Gas Experiment II (SAGE II) data, Li and Yu (2001) found that aerosol concentrations near 100 hPa are higher over the Tibetan Plateau than over China's central and northern regions in summer. Recent observations by balloon-borne optical particle counter (Tobo et al., 2007) and aircraft-borne measurements (Keim et al., 1996; Solomon, 1997) showed that soot-containing liquid aerosols with the major components of fine particles may also affect the aerosol layer near the tropopause. Appearance of cold tropopause in the upper troposphere (possibly in the lower stratosphere also) has been considered an important factor to explain the enhancement of tropopause aerosols observed in summer over the plateau (Kim et al., 2003). This observational fact is important from the point view of heterogeneous reactions on aerosol surfaces since gas-to-particle conversion processes are generally more active in low temperatures. During summer, the elevated surface heating and rising air associated with persistent deep convection over the Tibetan Plateau leads to anticyclonic circulation and divergence in the UTLS (Yanai et al., 1992; Hoskins and Rodwell, 1995; Highwood and Hoskins, 1998), where persistently enhanced pollutants such as aerosols, CO, methane and nitrogen oxides, as well as water vapor, can be linked to the rapid vertical transport of surface air from Asia, India and Indonesia in deep convection and confinement by strong anticyclonic circulation (Rosenlof et al., 1997; Jackson et al., 1998; Dethof et al., 1999; Park et al., 2004; Filipiak et al., 2005; Li et al., 2005a; Fu et al., 2006).

The aerosols from Nabro eruption might overlap with the background tropopause aerosols by deep convection in summer as mentioned in previous studies, changing their properties and evolvement in UTLS over the plateau. A clarification of the states that aerosols transport into and disperse out of the UTLS over the plateau is an important step toward understanding volcanic emission influences on hydration and chemical composition in the global stratosphere. Knowing the height dependence of the aerosol changes is important for understanding the volcano responsible for the transport of aerosols from the troposphere to the stratosphere over the Tibetan Plateau; however, a variety of aerosol vertical distributions and optical properties over the Tibetan Plateau have not been assessed in a satisfactory manner due to lack of continuous direct observations.

The vertical distributions of aerosol extinction coefficients were measured over the Tibetan Plateau in the summer of 2011, as part of the project Tibetan Ozone, Aerosol and Radiation (TOAR). In this study, the lidar and radiosonde measurement results are presented and compared with satellite data. We find a persistent maximum in aerosol extinction coefficients in the UTLS within the anticyclone, and show that such aerosol accumulation can be linked to the eruption of Nabro volcano. These results indicate that volcanic aerosol dispersed with the weakening of Tibetan anticyclonic circulation could primarily affect aerosol and hence radiation properties near the tropopause over the Tibetan Plateau.

#### 2 Measurements and data

#### 2.1 Micro-pulse lidar

An eye-safe, compact, solid-state micro-pulse lidar (MPL-4B, Sigma Space Corp., USA) was operated at the Naqu Meteorological Bureau  $(31.5^{\circ} \text{ N}, 92.1^{\circ} \text{ E}; 4508 \text{ m a.m.s.l.})$  on the central part of the Tibetan Plateau. The MPL is a backscatter lidar which uses an Nd:YLF laser with an output power of 12 µJ at 532 nm and 2500 Hz repetition rate. The diameter of the receiving telescope is 20 cm, and the field of view is 0.1 mrad. The vertical resolution of the lidar observation is 30 m, and the integration time is 30 s. Data obtained on the cloud-free days during nighttime were selected in order to avoid the disturbance of cloud and/or rain to columnaveraged lidar ratio and solar noise.

In general, the inversion of the lidar profile is based on the solution of the single scattering lidar equation:

$$P(r) = O_{\rm c}(r)CE\frac{\beta(r)}{r^2}\exp[-2\int_0^r \sigma(z)dz],$$
(1)

where r is the range, C is the lidar constant, which incorporates the transmission and the detection efficiency, and Eis the laser pulse energy.  $\beta(r)$  represents the total backscattering coefficient  $\beta(r) = \beta_{\rm m}(r) + \beta_{\rm a}(r), \sigma(r)$  is the total extinction coefficient  $\sigma(r) = \sigma_{\rm m}(r) + \sigma_{\rm a}(r)$ ,  $\beta_{\rm a}(r)$  and  $\sigma_{\rm a}(r)$ are aerosol backscattering and extinction coefficients, respectively.  $\beta_{\rm m}(r)$  and  $\sigma_{\rm m}(r)$  are molecular contributions to the backscattering and the extinction coefficients, respectively. They can be evaluated by the Rayleigh-scattering theory from the Standard Atmosphere 1976 (NASA, 1976). But here the molecular extinction coefficients are evaluated using temperature and pressure from the radiosondes released at the lidar field site twice a day.  $O_{\rm c}(r)$  is the overlap correction as a function of the range caused by field-of-view conflicts in the transceiver system. Systematic errors of  $P(\mathbf{r})$  were mainly observed in the lowest altitudes where an incomplete overlap between the emitted laser beam and the telescope field of view can lead to an underestimation of aerosol backscatter and extinction coefficients. Since the majority of aerosols are contained in the first several kilometers of the atmosphere, the overlap problem must be solved. Overlap is typically solved experimentally, using techniques outlined by Campbell et al. (2002). The starting point is an averaged data sample where the system is pointed horizontally with no obscuration. By choosing a time when the atmosphere is well mixed, such as late afternoon, when the aerosol loading is low, backscattering through the layer is roughly assumed to be constant with range (i.e., the target layer is assumed to be homogeneous). The similar overlap calibration was carried out at the beginning of this field experiment.

The vertical profile of aerosol extinction coefficient  $\sigma_a$  is determined by a near-end approach in solving the lidar equation as proposed by Fernald (1984). Considering the period of TOAR campaign was only 2 months after eruption of Nabro volcano, volcanic aerosols were still freshly nucleated particles with small size. The lidar ratios should therefore be rather high (Müller et al., 2007). Sawamura et al. (2013) employed the mean lidar ratio value of 50 sr at 532 nm for most groups of global lidar networks to trace the evolution of the stratospheric aerosol layer from Nabro volcano eruption. Therefore, a column-averaged lidar ratio of 50 sr is assumed for all measurements in this study.

We identify the boundaries of aerosol layer in the UTLS from the lidar extinction coefficient profiles. The lowest bin with  $\sigma_a = 0.002 \text{ km}^{-1}$  above 18 km is identified as the top of aerosol layer  $H_t$  and the bin with minimum value of  $\sigma_a$  between 10 and 16 km as the layer base  $H_b$ . The visible optical depth of the aerosol layer is derived by integrating the values of  $\sigma_a$  between  $H_b$  and  $H_t$ .

### 2.2 Radiosonde observations

During the field campaigns, 76 L band (GTS1) electronic radiosondes (Nanjing Bridge Machinery Co., Ltd., China) were launched to provide vertical profiles of pressure, temperature, and humidity up to 25 to 30 km high. The radiosondes were released at the lidar field site in Naqu twice a day at 00:00 and 12:00 UTC.

Eleven weather balloons with Vaisala RS92 radiosondes (Vömel et al., 2007) have been launched to provide profiles of air temperature, relative humidity (RH), wind speed and wind direction usually up to the mid-stratosphere. The RH can be measured between 0 and 100% with a resolution of 1% and an accuracy of 5% at -50 °C (Miloshevich et al., 2006; Währn et al., 2004), while Miloshevich et al. (2009) found that the RH measured by RS92 has a moist bias in the lower stratosphere (LS) and a dry bias in the upper troposphere (UT). To quantify the accuracy of RH measurement by RS92 over the Tibetan Plateau, we compared RS92 RH measurements with simultaneous water vapor measurements from a cryogenic frostpoint hygrometer (CFH) on 13 August 2011. The CFH is a lightweight (400 g) microprocessor-controlled instrument and operates on the chilled-mirror principle using a cryogenic liquid as a cooling agent. It includes several improvements over the similar NOAA/CMD instrument. It is currently designed to be combined with ozone sondes to provide simultaneous profiles of water vapor and ozone (Vömel et al., 2007). CFH has been taken in many intercomparison experiments as an absolute reference for water vapor measurements, including the validation of Aura Microwave Limb Sounder (MLS) water vapor products. After applying the time lag and solar radiation bias corrections, corrected RS92 RH measurements show agreement with CFH in the troposphere. The mean difference between corrected RS92 RH measurements and CFH is a dry bias of 2.9 % in the ground layer, while the mean differences in 5–10, 10–15 km and tropopause transition layer region are 1, 0.6 and 1.4 % moist bias, respectively. Therefore, the accuracy of corrected RS92 RH measurements is comparable to the accuracy of CFH in the UTLS (Yan, 2012).

### 2.3 Satellite observations

The Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) onboard CALIPSO (Winker et al., 2003) is used to characterize aerosol extinction profiles in the UTLS, which is a three-channel (532 nm parallel, 532 nm perpendicular, 1064 nm) elastic lidar receiving light at the same wavelength as the emitted laser frequency. CALIOP sends short and intense pulses (1064 and 532 nm) of linearly polarized laser light downward towards the Earth. The atmospheric backscatter profile is retrieved at 60 m vertical resolution from 8 to 20 km with a horizontal resolution of 1 km. The Level 2 aerosol extinctions at 532 nm of CALIOP (version 3.0) (available at: http://www-calipso.larc.nasa.gov/tools/data\_avail/) were used to compare with the ground-based MPL on the plateau. CALIOP data are selected over a  $300 \text{ km} \times 300 \text{ km}$  grid with an MPL location in its center.

We used the water vapor profile observations (version 3.3) from MLS on the NASA Aura satellite (Waters et al., 2006). Aura MLS measurements include water vapor, ozone and carbon monoxide, which are useful tracers of tropospheric and stratospheric air; these data have been used to document enhanced levels of carbon monoxide in the upper troposphere over the Asian monsoon (Li et al., 2005a; Filipiak et al., 2005) and also over the North American summer monsoon (Li et al., 2005b).

#### **3** Results

Figure 1a shows one case of aerosol enhancement observations in UTLS during the whole day of 7 August 2011. The case is the typical situation observed frequently over the Tibetan Plateau during the TOAR campaign. At nighttime (21:00–07:00 LST), aerosol enhancements were detected at relative constant altitudes from 17.0 to 18.5 km (a.m.s.l.) due to the higher signal to noise ratios (SNRs) of lidar compared with the daytime. A Vaisala RS92 radiosonde was launched at 06:56 UTC on 8 August and reached the tropopause at 17 km about 1 h later. Figure 1b presents the temperature and



Figure 1. (a) One case of aerosol enhancement observations during the TOAR campaign. Range-corrected 532 nm signals are shown with 30 s and 30 m resolution. (b) Temperature and RH profiles measured by the RS92 radiosonde.  $H_t$  and  $H_b$  denote the mean top and base heights of aerosol layer, respectively.

RH measured by this sounding along with lidar. These rather thick aerosol layers had optical depths around 0.01 and occurred in the temperature range between -70 and -80 °C. The radiosonde data indicate that the air in the aerosol layers was relatively dry with RH of about 5 % above an abrupt decrease of RH around the tropopause, which is very similar to the other cases measured this month with maximum RH of 10 %.

Figure 2 shows the vertical profiles of aerosol scattering ratios (SRs) measured at Naqu during 6–26 August 2011, along with the daily mean profiles of temperature. The measurements display relatively high aerosol extinction coefficients in the UTLS, which are 4–9 factors higher than those at altitudes below and close to (even higher than; such as on 6 and 12 August) molecular scattering coefficients at the same altitude. Compared with SR profiles of Nabro volcanic aerosols from MPLNET, EARLINET, NDACC and Hefei stations during June and July (Sawamura et al., 2013), the maximum SRs of aerosol layers in UTLS over the Tibetan Plateau are similar to those over Universitat Politècnica de Catalunya, Barcelona, Spain (41.39° N, 2.11° E), as one of EARLINET stations but larger than the other observations. The highest aerosol extinction coefficients in the UTLS over the Tibetan Plateau were generally located at 18–19 km altitudes, which are 1–2 km higher than the tropopause. Tropopause temperatures ranged from -70 to -80 °C, and the height of the tropopause varied from 80 to 100 hPa (from 17 to 18 km) during the observational period. Moreover, such relatively high aerosol extinction coefficients could extend over broad layers, ranging approximately 3–4 km.

The CALIOP aerosol extinction coefficients over UTLS are available for 12, 13, 18 and 20 August. Figure 3 compares the average extinction coefficient profile of MPL with that of CALIOP and shows a good agreement between the two instruments in both aerosol layers' altitude and the value of extinction coefficient. In particular, the MPL profiles show fewer standard errors at each vertical resolution altitude, possibly due to the good SNR of MPL observed at a high altitude



Figure 2. The nighttime mean aerosol scattering ratio profiles (dashed line) from MPL. The daily mean profiles of temperature (solid line  $^{\circ}$ C) from the two radiosondes each day are overlaid to indicate the altitude of the tropopause (~18 km a.m.s.l.).

of the lidar station and clear atmospheric environment over the Tibetan Plateau.

Figure 4 shows time series of total aerosol optical depth (AOD) and its daily averaged results, which varied from 0.075 to 0.142 with a maximum of 0.214 at 11:27 LST on 25 August. The percentage of AOD of the enhanced aerosol layers in total AOD each day also overlaps in this figure as the bar. The total AODs were derived from a Microtops II sun photometer collocated by lidar. The percentage of AOD varied between 5% (25 August) and 40% (12 August) with about 15 % for most days. AOD of the enhanced aerosol layers is only available at nighttime due to high lidar background noise at daytime, while data acquisition of total AODs from sun photometer is undergone at daytime, when more anthropogenic aerosols enter the atmosphere. Total AOD goes down at night, so the percentages of AOD would possibly be even greater than the above experimental values. However, the percentages of the UTLS AOD in the total AOD over the Tibetan Plateau are slightly higher than those observations, with the latter varying from 2 to 23 % at 532 nm. This might be on account of a more clean atmospheric environment over the Tibetan Plateau.

According to the period of occurrence of aerosol layers in UTLS, the continuous lidar observation can be split into



**Figure 3.** The average extinction coefficient profile of MPL (blue solid line) and the average extinction coefficient at each layer from CALIOP (red stars) during the whole observation period. The standard errors are marked as the error bar.

two stages: 6 to 12 (S1) and 22 to 26 (S2) August 2011 for the continuous maintenance stages of aerosol layer. Between the two stages, the existence of low clouds decayed the lidar signal to the extent that the available aerosol layer could not be observed in UTLS. Additionally, cirrus clouds in the upper troposphere would increase the retrieval error of extinction coefficients above the aerosol layer, and these cases were also removed from the data set. Figure 5 shows the daily variation in plateau monsoon index (PMI) and the 7-day-averaged PMI time series from 1 July to 31 August 2011, with an overlap of cirrus occurrence (He et al., 2013) and AOD in UTLS. PMI is an indicator of the daily mean intensity of the Tibetan Plateau monsoon. A larger PMI value indicates stronger monsoon in summer, which can be determined as follows (Tang et al., 1984):

$$PMI = H_1 + H_2 + H_3 + H_4 - 4H_0, (2)$$

where *H* is the daily deviation from the monthly mean geopotential height at 600 hPa. The subscript numbers 0 to 4 indicate the location of the center (90° E, 32.5° N), west (80° E, 32.5° N), south (90° E, 25° N), east (100° E, 32.5° N) and north (90° E, 40° N) of the plateau, respectively.

Many researchers have adopted the PMI to analyze the Tibetan Plateau monsoon variation. It is concluded that the index can reasonably describe the main characteristics of the Tibetan Plateau monsoon (e.g., Bai et al., 2001, 2005; Xun et al., 2011). These two stages might be caused by the different circulation systems due to an apparent time interval of about 10 days with PMI undergoing a substantial oscillation. During the first stage (from 6 to 12 August 2011), when the AOD decreased from 6 to 7 August and increased from 8 to 12 August, the values of PMI experienced an increasing trend from -20 on 6 August to 63 on 12 August. The values



**Figure 4.** Time series of total AOD (o) and its daily averaged results (+) derived from Microtops II sun photometer. The percentage of daily averaged AOD of the enhanced aerosol layers in the total AOD is defined as the bar.

sharply decreased to below -40 in the second stage with the low and continuous decreasing AOD over UTLS from 22 to 26 August 2011. Two obvious features can be found in the temporal variation of AOD: (i) AOD showed a decreasing trend accompanied by decreasing PMI during the campaign period, indicating that the aerosol layers wore off gradually with the reducing intensity of the Asian monsoon over the Tibetan Plateau at the end of August. Bourassa et al. (2012) found that the strong Asian monsoon anticyclone, which existed from June through September over Asia and the Middle East, where the Nabro volcanic aerosol was observed with OSIRIS, and the enhanced aerosol dispersed and quickly circulated throughout the Northern Hemisphere at the end of August, when the Asian monsoon anticyclone began to decay. And (ii) when the intensity of the Tibetan Plateau monsoon circulation subsided to PMI less than 0, the AOD in UTLS kept a persistent decline regardless of the variation trend of PMI, indicating that confinement of the air in the lower stratosphere induced by Asian monsoon anticyclone was destroyed to benefit the enhanced aerosol dispersing to the whole Northern Hemisphere.

According to the previous studies, deep convective activities are also considered to play an important role for transporting aqueous solution droplets of troposphere into stratosphere. It has been verified that deep convection over the Tibetan Plateau is likely to be a primary pathway for water vapor from the maritime boundary layer (e.g., Indian Ocean, South China Sea). Dessler and Sherwood (2004) have also suggested that convective transport plays a key role for the accumulation of water vapor near the tropopause, resulting in an increase of H<sub>2</sub>O mixing ratio by more than 5 ppmv near the tropopause (Gettelman et al., 2002; Park et al., 2004; Fu et al., 2006). But Tobo et al. (2007) used a growth model to calculate the possible growth under given atmospheric



**Figure 5.** Daily variation of PMI and the 7-day-averaged PMI time series from 1 July to 31 August 2011 and AOD in UTLS retrieved from MPL over the Tibetan Plateau. The days with cirrus occurrence are shaded (He et al., 2013). S1 and S2 represent two continuous maintenance stages of aerosol layer by lidar observation from 6 to 12 and from 22 to 26 August 2011, respectively.

conditions assuming the existence of liquid solutions at equilibrium with respect to  $H_2O$ ,  $H_2SO_4$  and  $HNO_3$ , and found that aerosol growth is sensitive to  $H_2O$  mixing ratios. According to the calculated growth curves of liquid solutions as a function of temperature and water vapor, the high  $H_2O$ mixing ratios (more than 5 ppmv) are indispensable conditions for producing high concentrations of fine particles near the tropopause. In fact, the  $H_2O$  mixing ratios near the tropopause from Vaisala RS92 radiosondes released in 6, 8, 11 and 23 August 2011 are not more than 2 ppmv, obviously less than the previous observations, as shown in Fig. 6. In consequence, the effects of gas-to-particle conversion from liquid solutions would likely be secondary to the enhancement of high tropopause aerosol extinction in these cases.

The continuous variation of water vapor distribution observed by satellite, despite lower vertical resolution, can also be used to investigate the contribution of liquid solution conversion to the enhancement of high tropopause aerosol extinction. Figure 7 shows the time series of water vapor profile derived from MLS, tropopause level from sounder temperature profiles, and the altitude of daily mean maximum aerosol extinction coefficients in this region. It can be clearly seen that almost all the abundant water vapor transported by deep convective systems is concentrated below 120 hPa altitude (about 15 km). Meanwhile, the temporal correlation of extinction coefficients in aerosol layer with water vapor from day to day is weak with a correlation coefficient of 0.36, suggesting that it is impossible that the enhanced tropopause aerosol is due to the condensation of water vapor.

In order to verify further that the enhanced tropopause aerosol is dominantly induced by the eruption of Nabro volcano, the aerosol loadings in UTLS over the Tibetan Plateau are compared with those over East China, where the influ-



**Figure 6.** Vertical profiles of water vapor from Vaisala RS92 radiosondes released in 6, 8, 11 and 23 August 2011, respectively. The black line along *y* axis represents the 5 ppmv of water vapor mixing ratio.

ence of the Asian monsoon anticyclonic circulation and deep convection transportation were significantly weak. Table 1 lists some statistical parameters of aerosol layer over Tibet and Shanghai (31.23° N, 121.53° E) for the same period. The aerosol parameters over Shanghai were also derived from an MPL with the same mode used in the Tibetan Plateau. The larger averaged extinction coefficient and higher AOD of the aerosol layer in UTLS over Shanghai demonstrate that the enhanced tropopause aerosol was dominated by the Nabro volcanic emissions with quickly circulation throughout the Northern Hemisphere at the end of August when the Asian monsoon anticyclone began to decay.

#### 4 Conclusions

In this study, we observed significantly increased aerosol extinction coefficients in UTLS over the Tibetan Plateau by continuous measurements with MPL during summer 2011. The retrieval of MPL showed a good agreement with CALIOP. The maximum SRs of aerosol layers, up to 4–9, in the UTLS were generally located in 18–19 km a.m.s.l., 1–2 km higher than in the tropopause, with broad layer depth ranging approximately 3–4 km. Daily averaged AODs of the enhanced aerosol layers in UTLS over the Tibetan Plateau varied from 0.007 to 0.030, which are at the same levels as the previous observations. The percentages of AOD for the enhanced aerosol layers in the total AOD are slightly higher than those observations by Sawamura et al. (2013).

The eruption of Nabro volcano is considered an important factor to explain the enhancement of tropopause aerosols observed in summer over the Tibetan Plateau. The aerosol layers wore off gradually with the reducing intensity of the

**Table 1.** Statistical parameters of aerosol layer over Tibet and Shanghai. Maximum extinction coefficient ( $EC_{max}$ ), averaged extinction coefficient ( $EC_{ave}$ ), aerosol layer depth (ALD), aerosol layer height (ALH) over sea level and aerosol optical depth (AOD) of the aerosol layer from 20:00 to 06:00 local standard time (LST). The numbers in parentheses correspond to the standard deviations.

	$EC_{max} (km^{-1})$	$EC_{ave} (km^{-1})$	ALD (km)	ALH (km)	AOD
Tibet	0.007	0.002 (0.002)	3.604 (1.626)	18.492 (0.248)	0.016 (0.002)
Shanghai	0.010	0.006 (0.003)	4.380 (0.764)	16.860 (0.839)	0.027 (0.006)



**Figure 7.** Altitude–time distributions of MLS water vapor (ppmv, color bar in natural logarithm) from 6 to 26 August 2011. Stars indicate the layer with nighttime mean maximum extinction coefficient and pluses stand for the tropopause level of each day, respectively.

Asian monsoon over the Tibetan Plateau at the end of August, and confinement of the air in the lower stratosphere induced by Asian monsoon anticyclone was destroyed to benefit the enhanced aerosol dispersion to the whole Northern Hemisphere. Deficiency in water vapor in UTLS indicates that the effects of gas-to-particle conversion from liquid solutions induced by deep convective activities would likely be secondary to the enhancement of high tropopause aerosol extinction in these cases.

It is must be noted that our interpretations are based on a short time observation. It is difficult to conclude that either one of the two processes is dominant due to lack of observations for trace gases. When further observations with more frequent soundings of water vapor and trace gases can be performed to investigate a correlation of high aerosol extinction with ambient temperatures, water vapor, trace gases, liquid solutions and transport processes, the result will be helpful in validating origination and mechanism of the enhanced aerosol extinction in UTLS.

Acknowledgements. This study was supported by Special Funds for Meteorological Research in the Public Interest (grant numbers: GYHY201106023, GYHY201006047 and GYHY201406001), the National Natural Science Foundation of China (NSFC, grant numbers: 40705013, 40975012 and 41175020), and the Shanghai Science and Technology Committee Research Special Funds (grant number: 10JC1401600). We thank all TOAR team members and the staff from the Tibet Meteorological Service for assisting our experiment work. We would like to thank the three anonymous reviewers and the editors of the manuscript, whose useful comments have greatly improved the paper.

Edited by: F. Yu

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