## Spatiotemporal variation of aerosol and potential long-range

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# transport impact over Tibetan Plateau, China

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## 25 Abstract:

26 The long-term temporal-spatial variations in the aerosol optical properties over the Tibetan 27 Plateau (TP) and the potential long-range transport from surrounding areas to TP were analysed in 28 this work, by using multiple years of supphotometer measurements (CE318) at five stations in the 29 TP, satellite aerosol products from the Moderate Resolution Imaging Spectroradiometer (MODIS) 30 and Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP), back-trajectory analysis from 31 the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) and model simulations 32 from the Goddard Earth Observing System (GEOS)-Chem chemistry transport model. The results 33 from the ground-based observations show that the annual aerosol optical depth (AOD) at 440 nm at 34 most TP sites increased in recent decades with trends of 0.001±0.003/year at Lhasa, 35  $0.013\pm0.003$ /year at Mt\_WLG,  $0.002\pm0.002$ /year at NAM\_CO, and  $0.000\pm0.002$ /year at QOMS\_CAS. The increasing trend was also found for the aerosol Extinction Ångstrom exponent 36 (EAE) at most sites with the exception of the Mt\_WLG site. Spatially, the AOD at 550 nm observed 37 from MODIS showed negative trends at the northwest edge close to the Taklimakan Desert and to 38 39 the east of the Qaidam Basin and slightly positive trends in most of the other areas of the TP. 40 Different aerosol types and sources contributed to a polluted day (with CE318 AOD at 440 nm > 41 0.4) at the five sites on the TP: dust was dominant aerosol type in Lhasa, Mt\_WLG and Muztagh 42 with sources in the Taklimakan Desert but fine aerosol pollution was dominant at NAM CO and

- 1 QOMS\_CAS with transport from South Asia. A case of aerosol pollution at Lhasa, NAM\_CO and
- 2 QOMS\_CAS during 28 April 3 May 2016 revealed that the smoke aerosols from South Asia were
- 3 lifted up to 10 km and transported to the TP, while the dust from the Taklimakan Desert could climb
- 4 the north slope of the TP and then be transported to the central TP. The long-range transport of
- 5 aerosol thereby seriously impacted the aerosol loading over the TP.
- 6 **Keywords:** Aerosol optical depth, Tibetan Plateau, aerosol pollution, long-range transport
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#### 1 **1. Introduction**

2 The heavy haze that has occurred in recent years in China has been largely attributed to the 3 atmospheric aerosols (Zhang et al., 2015). In addition, atmospheric aerosols can affect the climate through the interactions between aerosol-radiation and between aerosol-cloud (Takemura et al., 4 5 2005; Li et al., 2017), while the clouds and its precipitation are also connected to large scale 6 atmospheric circulations (Yang et al., 2010; Yang et al., 2017a). However, there is still a high level 7 of uncertainty about the impact of aerosols on the climate, which is mostly due to the high 8 spatiotemporal variability of aerosols. Therefore, studying the physical and chemical properties of 9 aerosols over different regions is essential.

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11 The Tibetan Plateau (TP), is the largest elevated plateau in East Asia and considered as one of 12 the most pristine terrestrial regions, along with the Arctic and Antarctic. However, in the past two 13 decades, the TP has been surrounded by an unprecedented growth of emissions of Asian air 14 pollutants from various sources. Consequently, some studies have demonstrated that the aerosols 15 transported from its around areas (South Asia and Taklimakan Desert) have polluted the TP (Huang et al., 2007; Xia et al., 2011; Kopacz et al., 2011; Lu et al., 2012; Liu et al., 2015). The increase in 16 17 aerosols over the TP may have an important impact on the regional or global climate. Lau et al. 18 (2006) has suggested that increased absorbing aerosols (dust and black carbon) over the TP may 19 create a positive tropospheric temperature anomaly over the TP and adjacent regions to the south, 20 causing the advance and enhancement of the Indian summer monsoon. Attempts have been made to 21 reveal the linkages between climate change (such as changes to glaciers and monsoons) and the air 22 pollutants around the TP (mainly absorbing carbonaceous materials) (Qian et al., 2011; Wang et al., 23 2016; Lee et al., 2013). However, the quantitative effect of the TP aerosol on climate variability 24 remains largely unknown, and it is very essential to fully understand the aerosol characteristics over 25 the TP.

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27 A large amount of attention has been paid to aerosol characteristics over the TP (Wan et al., 28 2015; Tobo et al., 2007; Zhao et al., 2013; Liu et al., 2008; Du et al., 2015). Although the seasonal 29 variations in aerosol properties over the TP have been analysed based on ground-based observations 30 or satellite products (Shen et al., 2015; Xia et al., 2008), analysis is needed of the long-term trends in the variation of aerosols over the TP to provide predictions and guidelines for environment 31 32 policies. In past studies, spring or summer have often been studied due to the important impacts of 33 dust and carbonaceous aerosols (Huang et al., 2007; Cong et al., 2007; Lee et al., 2013). However, 34 most studies of the aerosol properties based on ground-based measurements have been conducted 35 at a single site over the TP, such as NAM CO (Cong et al., 2009), Mt Yulong (Zhang et al., 2012), 36 and Mt\_WLG (Che et al., 2011). Past studies have mostly focused on single stations or short-term 37 variations due to the difficulty of taking a sufficient number of ground-based observations in 38 challenging weather conditions over the remote plateau.

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Ground-based measurements can offer more accurate data on aerosol properties, while largescale spatial observations of aerosol optical and physical properties require satellite remote-sensing methods (Li et al., 2015; Li et al., 2018; Xing et al., 2017). Thus, the long-term detection of aerosols from both ground and satellite platforms is absolutely necessary for improving our understanding of the climate effects of aerosol over the TP region. Consequently, based on multiple years of observations from five ground-based sunphotometers at the TP and the MODIS aerosol optical depth product over the TP region, our work here is focused on the long-term spatiotemporal variations in the aerosol optical properties over the TP and the aerosol properties and sources during the high aerosol loading over the TP. In addition, we also combined the observation and models to study the serosol transport process over the TP, thereby helping to reduce the uncertainties in estimating aerosol radiative forcing and aerosol sources.

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8 In this paper, section 2 describes the observation sites, data and methods. The results of the 9 analysis of the spatiotemporal variations in aerosol properties over the TP are shown in section 3. 10 The analysis of aerosol high loading and an aerosol transport case are presented in section 4 and 5, 11 respectively. The conclusions are presented in section 6.

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## 13 **2. Site, data and methodology**

## 14 **2.1 Sites**

15 In this study, five sites in the TP equipped with sun and sky scanning radiometers (CE318) were used (Figure 1). Table 1 shows the station locations and descriptions. Lhas a station is the only 16 17 urban site that suffers from local anthropogenic emissions. For the other four sites, local 18 anthropogenic emissions are extremely rare due to the low number of human inhabitants. However, 19 Mt\_WLG is in the northeast of the TP, where it is situated on the dust transport path from the largest 20 desert in China (the Taklimakan Desert). The Muztagh\_Ata site is located in the northwest corner 21 of the TP and next to the Central Asian Deserty and the Taklimakan Desert. NAM\_CO is located on 22 the central Tibetan Plateau, 220 km away from Lhasa. QOMS\_CAS is located at the northern slope 23 of Mt. Qomolangma on the border of Tibet and Nepal. Therefore, these five sites are representative 24 of the spatial features of the TP.

## 26 **2.2 Data**

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#### 27 2.2.1 CE318 aerosol optical properties

28 The column-integrated aerosol properties over the five TP sites are derived from CE318 29 measurements. Table 1 shows the observation period. The CE318 instrument measures direct solar 30 spectral radiation and the angular distribution of sky radiance. These spectral radiances can be used 31 to retrieve aerosol optical parameters (such as aerosol optical depth (AOD)) based on Beer's Law 32 and aerosol microphysical properties (such as volume size distribution) and the radiative forcing 33 through radiation transfer theory (Dubovik and King, 2000; Dubovik et al., 2006). The instruments 34 were periodically calibrated using the Langley method at AERONET global calibration sites (the 35 Izaña, Spain or the Mauna Loa, USA) or using the inter-comparison calibration method at the 36 Beijing-CAMS site (Che et al., 2015). The cloud-screened and quality-controlled data of AOD, 37 Extinction Ångstrom exponent (EAE), and aerosol volume size distribution (dV(r)/dlnr) are used in 38 this work (Giles et al., 2019). Eck et al. (1999) showed that the uncertainty of the AOD was approximately 0.01 to 0.02. The EAE was calculated from the AOD at 440 and 870 nm. The errors 39 40 of retrieval for dV(r)/dlnr were less than 10% in the maxima of the dV(r)/dlnr and may increase up 41 to 35% for the minimum values of dV(r)/dlnr within the radius range between 0.1 µm and 7 µm; for 42 the edges of the retrieval size, the errors increased apparently, but did not significantly affect the 43 derivation of the main feature of dV(r)/dlnr (Dubovik et al., 2002).

## 1 2.2.2 The MODIS AOD product

2 The Moderate Resolution Imaging Spectroradiometer (MODIS) instrument is a multi-spectral 3 sensor with a wide spectral range from 0.4 to 14.4 µm in 36 wavelength bands, onboard the Terra (morning descending direction) and Aqua (afternoon ascending direction) satellites in polar orbit. 4 5 It's broad swath of 2330 km permits retrieval aerosol products to cover the global word within 1-2 6 days. In this study, both Terra and Aqua MODIS Collection 6 Deep-Blue (DB) and Dark-Target (DT) 7 combined AOD at 550 nm product with 10km spatial resolution (Levy et al., 2013) from 2006 to 8 2017 were used. The MODIS AOD at 550 nm (MODIS\_AOD) combined the DT and DB algorithms merges the products from the two algorithms based on the normalized difference vegetation index 9 10 (NDVI) statistics as follows: 1) the DT AOD data are used for NDVI > 0.3; 2) the DB AOD data 11 are used for NDVI < 0.2; and 3) the mean of both the algorithms or AOD data with high quality flag 12 are used for  $0.2 \le \text{NDVI} \le 0.3$ . The MODIS\_AOD has been validated in the global or regional areas 13 (Bilal et al., 2018; Ma et al., 2016; Sayer et al., 2014). The root-mean-square error of MODIS\_AOD 14 was about 0.13, and the percentage of MODIS AOD data within the expected error was more than 71% at the Kunming site, which is near the TP (Zhu et al., 2016). 15

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## 17 2.2.3 The CALIOP profile data

18 The Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP), the primary instrument on 19 board of CALIPSO satellite, is a near-nadir viewing two wavelength (532 nm and 1064 nm) 20 polarization-sensitive lidar that performs global vertical profile measurements of aerosols and 21 clouds (Winker et al., 2010). It provides three primary calibrated and geolocated profile products: total attenuated backscatter at 532 nm and 1064 nm and the perpendicular polarization component 22 23 at 532 nm. The CALIOP (version 4.10) products used in this study include the attenuated 24 backscattering coefficient profiles from Level 1B and the vertical feature mask data products of 25 aerosol subtype from level 2 products under 15 km altitude, which were downloaded from the Langley Atmospheric Science Data Center (ASDC). Kumar et al. (2018) have showed that the AOD 26 27 from CALIOP version 4.10 agreed with the ground-based CE318 observation at a site in the central Himalayas with a correlation > 0.9 and  $\sim 87$  % matchup data were within the expected error. 28

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## 30 2.3 Methodology

The ground-based CE318 observations and MODIS AOD products were analysed to show the spatiotemporal variations in aerosol properties in TP.

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The CE318 observed AOD at 440 nm with values larger than 0.4 at each site was specially 34 35 analysed to study the aerosol properties of the high aerosol loading over the TP. The value of 0.4 36 was selected because the mean annual values of AOD observed by CE318 instruments at the TP 37 sites were less than ~0.1 in the past studies (Xia et al., 2016; Cong et al., 2009), and this value is 38 normally regarded as the high aerosol loading (Eck et al., 2010; Giles et al., 2012). Back trajectories were used for the aerosol source analysis in the TP. The back trajectories on the high aerosol loading 39 40 days were calculated by using the Hybrid Single-Particle Lagrangian Integrated Trajectory 41 (HYSPLIT) model which is driven by the one degree horizontal resolution archived meteorological 42 fields with (Draxler and Hess, 1998). 72-hour back trajectories ending at the five sites at 10 m above 43 ground level at 12 UTC on the days with high aerosol loading (AOD at 440 nm >0.4) were used to identify the air mass sources. 44

A case of long-range aerosol transport to the TP was selected based on the ground CE318 observations over Lhasa, NAM\_CO and QOMS\_CAS. The HYSPLIT back trajectories and the MODIS and CALIOP products were used to show the potential aerosol sources, spatial aerosol loading and the vertical features of the aerosol over the TP during the case period. In addition, the Goddard Earth Observing System (GEOS)-Chem chemistry transport model was used to simulate the AOD and its components (dust and carbon aerosol) during the case period, which may reflect the change in aerosol type during the case period.

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10 The GEOS-Chem chemical transport model (version 11-01) coupled with the online radiative 11 transfer calculations (RRTMG) at  $0.5^{\circ} \times 0.667^{\circ}$  horizontal resolution over East Asia domain (Bey 12 et al., 2001; Wang et al., 2004) was used. The model was driving by the Global Modeling and 13 Assimilation Office (GMAO) MERRA-2 meteorology with the temporal resolution of 3 hours for 14 meteorological parameters and 1 hour for surface fields. The simulation type of full chemistry in the troposphere was selected. The implementation of RRTMG in GEOS-Chem was described in 15 Heald et al. (2014). The AOD was calculated according to Martin et al. (2003). The default global 16 17 anthropogenic emissions were overwritten over East Asia by the MIX inventory from Li et al. (2014). 18 The Global Fire Emission Database (GFED) (van der Werf et al., 2010) has been used to specify 19 emissions from fire. More details on the model and the other emissions data used and the evaluation 20 of AOD in the east and south of the TP were shown in Zhu et al. (2017).

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In this study, the AOD from the CE318, MODIS, and GEOS-Chem model were used. For convenience, CE318\_AOD, MODIS\_AOD, and Model\_AOD stand for the AOD observed by CE318, MODIS, and the AOD simulated by the GEOS-Chem model, respectively. For CE318\_AOD, the 440 nm wavelength is often studied, while MODIS\_AOD and Model\_AOD generally use the data at 550 nm wavelength. Thus, unless otherwise specified, CE318\_AOD, MODIS\_AOD, and Model\_AOD hereinafter represent the ones at 440 nm, 550 nm, and 550 nm, respectively.

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## 30 **3. Temporal-spatial variations in aerosol properties**

31 **3.1** Aerosol properties observed by the CE318 instruments

The monthly, seasonal, and annual variations in aerosol properties observed from the CE318 instruments at the five TP sites were analyzed.

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35 The monthly variations in CE318 AOD and EAE at the five sites over the TP are shown in Figure 2. The monthly mean CE318\_AOD was highest in April at the Lhasa (0.19), NAM\_CO (0.09) 36 37 and QOMS\_CAS (0.10) sites, while the value at Mt\_WLG was highest in June (0.20). The monthly 38 mean CE318\_AOD rapidly increases from January to April and then slightly decreases until 39 December at the Lhasa, NAM\_CO and QOMS\_CAS sites. However, the monthly mean 40 CE318\_AOD at Mt\_WLG is nearly symmetrical form from January to December. The monthly 41 variation in EAE is different from the AOD. The highest monthly EAE values occur in September 42 at Lhasa (1.15), October at Mt\_WLG (1.15) and in January at the NAM\_CO (0.93) and 43 QOMS\_CAS (0.17) sites. The EAE at QOMS\_CAS also shows a high value of 0.17 in April, which 44 may be caused by the smoke aerosol transported from South Asia during this period. The monthly

1 mean EAE first decreases from January to March and then increases until September at Lhasa. The 2 monthly mean EAE values at NAM CO also decreases from January to March, but does not increase 3 apparently in the following months. The EAE at Mt\_WLG decreases from January to May and then increases obviously from May to October. The Lhasa, NAM CO, and QOMS CAS sites are near 4 5 and located in the south of the TP. Thus, the variations in the aerosol properties at these three sites 6 are similar. The Mt\_WLG site is located in the northeast of the TP, which is different from the 7 southern sites. The Muztagh Alt is in the northwest of the TP and is the closest site to the 8 Taklimakan Desert, which causes the high AOD there (a few observed data may be another reason). 9 Looking at the monthly CE318 AOD and EAE values together, high CE318 AOD is often 10 accompanied by low EAE at Lhasa, Mt\_WLG and NAM\_CO, indicating that these sites suffered 11 from coarse aerosols such as dust (Huang et al., 2007; Liu et al., 2015; Zhang et al., 2001). However, 12 the QOMS\_CAS site show high CE318\_AOD and high EAE in April, which may be related to 13 smoke aerosols transported from South Asia.

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Table 2 shows the seasonal statistics of CE318\_AOD and EAE. A distinct seasonal variation 15 in CE318 AOD and EAE can be found over the TP sites. The CE318 AOD mean values in fall 16 17 (SON) and winter (DJF) are lower at all sites except Muztagh. Muztagh\_Ata shows high 18 CE318 AOD in both observed seasons. Except for that in Muztagh, the maximum seasonal 19 CE318\_AOD is observed in spring (MAM) (Lhasa, NAM\_CO, and QOMS\_CAS) or in summer 20 (JJA) (Mt\_WLG). The minimum seasonal EAE occurred in spring (Lhasa, NAM\_CO and Mt\_WLG) 21 or summer (QOMS\_CAS), while the maximum EAE values are mostly observed in fall (Lhasa and 22 Mt\_WLG) and winter (NAM\_CO and QOMS\_CAS). These indicate frequent dust events over the 23 TP in the spring period at Lhasa, NAM CO and Mt WLG. Mt WLG is situated on the dust 24 transport path from the Taklimakan Desert, which causes the high CE318\_AOD observed in spring 25 and summer at this site.

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27 The seasonal size distributions of the five sites in Figure 3 also demonstrate that coarse mode 28 aerosol is dominant at the five TP sites in almost all seasons, which is different from those in the 29 eastern pollution regions of China, such as Yangtze River Delta, where fine mode aerosol is 30 dominant (Zhuang et al., 2018). This size distribution explained the relatively low annual averages 31 of EAE at the five sites (all annual EAE in Figure 2 are less than 1.0), compared to the those at the 32 inland urban and suburban sites in China (Xin et al., 2007), such as Beijing (1.19) (Fan et al., 2006), 33 Nanjing (1.20) (Zhuang et al., 2018; Zhuang et al., 2017), Kunming (1.25) (Zhu et al., 2016), and 34 Chengdu (1.09) (Che et al., 2015). What's more, spring is the season with a high-volume 35 concentration of coarse mode aerosol. Among the five sites, the southernmost sites, QOMS CAS, 36 showed the highest mean EAE and the size distribution was distinctly bimodal, especially in spring. 37 This was also because of the frequent biomass burning activity in India and Nepal, which can 38 transport the fine aerosol to the QOMS\_CAS site.

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The annual averages of CE318\_AOD (shown in Figure 2) are 0.05-0.14 over TP sites. These
average values are lower than those in other regional background sites, such as Longfengshan (0.35)
in Northeast China (Wang et al., 2010), Xinglong (0.28) in North China Plain (Zhu et al., 2014),
Lin'an (0.89) in Eastern China (Pan et al., 2010) and Dinghushan (0.91) in Southern China (Chen
et al., 2014). The low aerosol loading over the five TP sites indicates excellent air quality over the

- 1 TP region.
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3 However, the aerosol loading at the TP sites presents interannual changes. The annual variations in CE318 AOD and EAE over TP at the four sites, i.e. Lhasa, Mt WLG, NAM CO, and 4 5 QOMS\_CAS are shown in Figure 4. The data for the CE318 observations at Muztagh\_Ata site are 6 only available for 2010; thus, the annual variation at this site is not shown here. The annual 7 CE318\_AOD shows increasing trends of  $0.001 \pm 0.003$ /year at Lhasa,  $0.013 \pm 0.003$ /year at 8 Mt\_WLG, and 0.002±0.002/year at NAM\_CO during the CE318 observation period. The Mt\_WLG site shows the most obvious increase in CE318 AOD during 2009-2013. These results indicate an 9 10 increase in aerosol loading at the three sites. The long-term annual variation of CE318\_AOD at 11 OOMS CAS is very small  $(0.000\pm0.002/\text{vear})$ , but there are still short-term annual variations (the 12 values decreased from 2010 to 2013 and increased from 2013 to 2016). The annual trends of EAEs 13 are more evident than the CE318\_AOD at these four site. Most sites show an increasing tendency 14 in the average annual EAE except for Mt WLG site, which shows a large decreasing trend of -0.318 15  $\pm 0.081$ /year. This shows that the size of aerosol at the Mt\_WLG site increased, while the size of the aerosol decreased in the other three sites. Looking at the CE318 AOD and EAE values together, 16 17 the positive trend of CE318\_AOD and the positive trend of EAE in the long term variation at most 18 sites over TP indicates the addition of fine mode aerosol which may be related to the anthropogenic 19 impact or long-distance transport of dust to the TP. However, in the short term, the increase in the 20 average annual CE318\_AOD is often associated with the decrease in EAE over the TP, which 21 suggests the addition of coarse mode aerosol during the CE318 observation period.

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#### 3.2 Aerosol properties from MODIS

24 Ground-based observations can offer accurate aerosol optical properties at point locations but 25 lack spatial coverage. The MODIS aerosol product can provide the spatial variation in AOD over 26 the TP. Thus, we evaluated the MODIS\_AOD using the ground-based observation CE318\_AOD at 27 550 nm over the TP sites. The CE318\_AOD at 550 nm was interpolated from 440 nm, 675 nm, 870 nm and 1020 nm by using an established fitting method from Ångström (1929). The matchup 28 29 method was that the CE318 data within 1 hour of the MODIS overpass were compared with the 30 MODIS data within a 25 km radius of the ground-based site. The minimum requirement for a 31 matchup was at least 3 pixels from MODIS.

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33 Figure 5 shows the results of MODIS\_AOD compared to the collocated ground CE318 34 observations over the TP. There are 996 instantaneous matchups of Terra and Aqua MODIS during 35 the CE318 instrument measurement period at the five TP sites. The MODIS AOD overestimates 36 the AOD at 550 nm with a positive mean bias of 0.02 and a root mean squared error (RMSE) of 37 0.11. The RMSE value is lower than that of the North China Plain sites (~0.25) (Bilal et al., 2019). 38 The slope and intercept of the best-fit equation between the MODIS\_AOD and CE318\_AOD at 550 nm are 0.46 and 0.06, respectively, with a correlation coefficient (R) of 0.54. There are 67.8% of 39 the compared AODs within the expected error envelope of 0.05+0.15AOD (%EE). The R value is 40 41 lower than that in the global assessment statistics, while the %EE is higher than that in the global 42 evaluation (Bilal and Qiu, 2018). Overall, the results suggest that the MODIS\_AOD product can be 43 used to study the aerosol spatial variation over the TP region.

1 The spatial distribution of the annual MODIS\_AOD is shown in Figure 6. The MODIS\_AOD 2 agrees with the CE318 AOD at 550 nm at the five TP sites. The northwest area around the 3 Taklimakan Desert and the northern part of the TP on the transport path of the Taklimakan Desert dust showed high MODIS AOD (>0.25) in recent decades. In addition, the southern edge performed 4 5 slightly high MODIS\_AOD (0.2-0.25) influenced by the aerosol transport from South Asia. There 6 is some small area with high MODIS\_AOD (~0.2) in the centre of the TP, and the southeast region 7 shows low MODIS AOD ( $\sim 0.1$ ), which may be attributed to the aerosol transport and surface 8 features such as vegetation cover, since there are few inhabitants. The seasonal departure of 9 MODIS AOD (Figure 7) shows that high positive MODIS AOD departure often appears in spring, 10 especially for the northwest edge, northern area and southern edge of TP, which was a result of 11 aerosol transport from the frequent dust events in the Taklimakan Desert and the fire activities in 12 South Asia in spring.

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14 A linear regression analysis of the trends in annual MODIS AOD over the TP from 2006 to 15 2017 was conducted using the least squares method. The spatial distribution of the annual trends in MODIS AOD during 2006-2017 is illustrated in Figure 8. There are no statistically significant 16 17 trends in most areas during 2006-2007. The MODIS\_AOD showed negative trends in the northwest 18 edge close to the Taklimakan Desert and to the east of the Qaidam Basin and slightly positive trends 19 in most of the other areas. The areas where MODIS AOD decreased are mainly located near the 20 desert or on the transport path of the desert dust. This descending trend may be related to the 21 significant reduction in dust emissions caused by the decline in wind speed in recent years (Yang et 22 al., 2017b). The positive trend in other most areas may be due to the rapid increase in human 23 activities, such as the expansion of tourism to the TP and biomass burning in South Asia. 24

- 25 The seasonal trends in MODIS\_AOD at 550 nm over the TP during 2006-2017 are presented in Figure 9. The spring showed the most obvious decline in MODIS\_AOD (~ 0.02/year) at the 26 27 northern edges and northeast part of the TP during 2006-2017, which also suggested that the 28 reduction in dust impact from the Taklimakan Desert like the trend in the annual MODIS AOD 29 (seen in Figure 8). In summer, the positive trend in MODIS\_AOD over the TP was relatively 30 apparent, and most higher sporadic positive values of  $\sim 0.01$  occurred in the central and southern 31 part of the TP. Summer is the tourist season in the TP and tourism has developed in past decades, 32 which may be one of the reasons for the higher positive trend in summer in the TP. The positive 33 trends in autumn and winter were relatively lower than those in summer, and the most positive trends were located at the northern TP. The reason for this phenomenon needs to be explored. 34
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#### 36 4. Aerosol properties and potential sources during high aerosol loading

37 The annual mean AOD in the TP is normally low due to the few human inhabitants and high 38 altitude. However, some high CE318\_AOD values larger than 0.4, which is normally regarded as high aerosol loading (Eck et al., 2010; Giles et al., 2012), were observed at the five sites in the TP 39 40 by CE318. Thus, the CE318\_AOD larger than 0.4 over TP can be considered as the aerosol pollution. 41 The frequencies of high aerosol loading (CE318\_AOD > 0.4) during the CE318 measurements were 42 1.57%, 1.79%, 0.21%, 0.42% and 0.11% at the Lhasa, Mt\_WLG, Muztagh\_Ata, NAM\_CO, and 43 QOMS\_CAS sites, respectively. The aerosol properties and sources during high aerosol loading in 44 the TP need to be studied.

2 Figure 10 shows the CE318 AOD with values larger than 0.4 versus EAE observed by CE318 3 at the five sites in the TP. Except for the Lhasa and Mt\_WLG sites, almost all values of CE318\_AOD are less than 1.0, which reflects the relatively clear environment over the TP. The EAE shows two 4 5 centres at ~0.1 and ~1.5. The low EAE (~0.1) centre is related to dust events, which can cause higher 6 concentrations of coarse particles in the atmosphere. Besides, most values of the low EAE (<0.5) 7 part are less than 0.2 (only a few EAEs between 0.2-0.5 are observed at Lhasa and Mt WLG), 8 indicating that the pure dust type is more common than the polluted dust type in the TP according 9 to Eck et al. (2010). The high EAE centre at ~1.5 indicates mainly small sub-micron radius particles, 10 which can be attributed to anthropologic emissions. The values of EAE >1.0 at the NAM\_CO and 11 QOMS\_CAS sites are generally higher than those at the Lhasa and Mt\_WLG sites. According to 12 past studies, the EAE of biomass burning aerosol is generally higher than the urban/industry aerosol 13 (Giles et al., 2012; Eck et al., 2010), which may cause the higher EAE at NAM\_CO and 14 QOMS CAS (more biomass burning aerosol) than at Lhasa and Mt WLG (more urban/industry 15 aerosol). On the other hand, the values in the middle range of 0.5-1.0 are rare, indicating the less mix of natural and human sources. The percentage of EAE bins to the number of CE318\_AOD>0.4 16 17 is distinct (Table 3). The percentage of EAE <0.5 is high than that of EAE>1.0 at Lhasa, Mt\_WLG 18 and Muztagh Ata, indicating more nature dust pollution than the anthropogenic pollution at these 19 three sites. However, a greater number of high EAE values (>1.0) are observed than EAE<0.5 at the 20 NAM\_CO and QOMS\_CAS sites, suggesting that anthropogenic pollution is more than natural dust 21 pollution at these two sites.

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Figure 11 shows the aerosol size distribution binned by CE318\_AOD at the five sites in the TP. The volume concentration of coarse mode particles increases more apparently than fine mode at Lhasa, Mt\_WLG and Muztagh sites when the values of CE318\_AOD increase. However, the size distribution at NAM\_CO and QOMS\_CAS shows the dominant increase of fine mode aerosol. These indicate of the different aerosol type pollution in these five sites: dust dominant in Lhasa, Mt\_WLG and Muztagh and fine mode aerosol (mainly biomass burning aerosol) pollution dominant at NAM\_CO and QOMS\_CAS.

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The dominant aerosol pollution type showed obvious distinctions among the five sites on the 31 32 TP, then where is the distinct aerosol pollution source at each site? We used the HYSPLIT back-33 trajectory model and the MODIS\_AOD on the day with aerosol pollution (CE318\_AOD >0.4) to 34 show the aerosol source on the pollution day at each site. Figure 12 shows the 72 hour back-35 trajectories ended at the five site (10 m above ground level) in the TP overlaid with the mean 36 MODIS\_AOD on the aerosol pollution day observed by the ground-based CE318 37 (CE318\_AOD >0.4). The CE318 instruments observed 78, 20, 2, 15, and 14 days with instantaneous 38 AOD at 440 nm > 0.4 at Lhasa, Mt\_WLG, Muztagh\_Ata, NAM\_CO and QOMS\_CAS, respectively. The aerosol pollution days at Lhasa, Mt\_WLG, and Muztagh\_Ata observed by CE318 are often 39 40 with low EAE (black trajectories). The airflows ended at the Lhasa site on the polluted days are 41 mainly from the northwest and southwest. The MODIS\_AOD around Lhasa in the area of the back-42 trajectories with CE318 EAE <0.5 passing does not show significantly high values, especially in the 43 Taklimakan Desert, which indicates that the dust pollution at Lhasa is mainly from local or surrounding dust events rather than transport from the Taklimakan Desert. The Mt WLG shows that 44

1 the air mass on the pollution days comes from the west and east and the path of back trajectories has high MODIS AOD. The high values of MODIS AOD shows two transport paths of dust aerosol 2 3 to Mt\_WLG: one is through the Qaidam Basin and the other is through the northeast edge of the TP. The two polluted days observed by CE318 at the Muztagh Ata shows the easterly airflows 4 5 originating from the Taklimakan Desert. The direction of the back-trajectories of EAE<0.5 that 6 ended at NAM\_CO is similar to Lhasa, while the southerly air flows with high EAE (red trajectories) 7 originate from Nepal, where frequent biomass burning happened and caused the high MODIS AOD 8 values. The trajectories ended at QOMS\_CAS and the high MODIS\_AOD of the path revealed the 9 transport of finer aerosol from South Asia to this site.

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## 11 5. Case study of long-range transport to the TP

12 The long-range transport of aerosol can cause the aerosol pollution and affect the long-term 13 variation in aerosol over the TP. In addition, the dominant aerosol type may change at the TP sites 14 during a case of aerosol transport. Thus, a specific case of aerosol pollution during 27 April - 3 May 15 2016 was analysed further. This case is selected based on the observations from the CE318 instrument. During 28 April -1 May, the CE318\_AOD at Lhasa, NAM\_CO, QOMS\_CAS sites 16 17 showed the values larger than 0.4, which reached up to more than 3 times the mean values of 18 CE318 AOD of each site (0.11 at Lhasa, 0.05 at NAM CO and QOMS CAS). This was indicative 19 of aerosol pollution at the three sites. Then, how about the aerosol properties of this period and 20 where did the polluted aerosol come from?

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22 Figure 13 shows the daily CE318\_AOD and EAE during 27 April – 03 May at the three sites. 23 The mean values of CE318 AOD were 0.45, 0.38, and 0.23 at Lhasa, NAM CO and QOMS CAS, 24 respectively. These even reached 4 times the annual mean CE318\_AOD at each site. The mean 25 EAEs were 0.98, 1.22, and 1.44 at Lhasa, NAM\_CO and QOMS\_CAS, respectively, which was 26 higher than the annual averages and suggested the fine aerosol entrance. There were CE318\_AOD 27 peaks at the three sites during 27 April - 03 May. Lhasa showed an increase of CE318\_AOD from 0.30 on 27 April to 0.51 on 28 April and maintained high CE318 AOD to a value of 0.54 on 1 May, 28 29 after which it decreased to 0.34 on 2 May. NAM\_CO also showed an increase of CE318\_AOD 30 during the first two days of the period, but decreased after 29 April. QOMS\_CAM showed a slight increase in CE318\_AOD from 27 April to 30 April, which was later than those of the other two sites. 31 32 Combining the EAE on these days, fine mode aerosol was brought to Lhasa and NAM\_CO during 33 27-29 April, and then coarse aerosol began to occur on 30 April, and even became the dominant 34 aerosol in the following several days. The fine aerosol at the QOMS\_CAM site were maintained for 35 an additional day after those at the two sites, and then the coarse aerosol increased.

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37 The GEOS-Chem model simulation also supported the above results. Figure 14 shows the comparison between the Model\_AOD ( $0.5^{\circ} \times 0.667^{\circ}$ ) and CE318\_AOD at 550 nm and the ratios of 38 39 the model simulated aerosol types (dust, both organic carbon (OC) and black carbon (BC) aerosol) 40 to the total Model\_AOD during this case period at the three sites. The evaluation results showed 41 that the model underestimated the daily AOD at the three sites during this period, with negative 42 mean biases from -0.28 to -0.08. However, the Model\_AOD was relatively high correlated with the 43 CE318\_AOD at 550 nm, with the R values of 0.61 at Lhasa, 0.89 at NAM\_CO and 0.86 at 44 QOMS CAS. These R values were higher than the model evaluation in South China and the Indo-

1 China Plain (~0.5) (Zhu et al., 2017). Thus, the variation trend from Model\_AOD agreed well with 2 that measured by the CE318 instruments during these days. During the first 4 days of the case period 3 (27 April to 30 April), the ratios of different aerosols to the total Model\_AOD showed that the sum of OC and BC aerosols were higher than those of dust aerosol at all three sites. Besides, the sums 4 5 of OC and BC at Lhasa and QOMS\_CAS were higher than that of NAM\_CO. These indicated that 6 the smoke aerosol affected the three sites more severely than dust during the first 4 days and Lhasa 7 and QOMS CAS sites were nearer to smoke sources than NAM CO. After 30 April, the sum of BC 8 and OC decreased while dust increased, and the increase of dust at the three sites was NAM\_CO > 9 Lhasa > QOMS CAS. Therefore, the major aerosol source was changed and the NAM CO site was 10 closer to dust source after 30 April. This phenomenon had continued to 2 May at NAM\_CO and 11 Lhasa, and 1 May at QOMS\_CAS. In the last one or two days, the dust decreased while the smoke 12 obviously increased, which could cause the mixture of these two aerosols.

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14 Then, how was the spatial aerosol loading around the TP and the vertical feature of aerosol 15 transported to the TP? Figure 15 shows the MODIS\_AOD and 72-hour back trajectories at Lhasa (the first row), the CALIOP-derived vertical profile of total attenuated backscatter at 532 nm (the 16 17 second row), and the vertical feature mask of aerosol (the third row) on 28 April, 1 May, and 3 May 18 during the case study period. The MODIS AOD showed high values in the south (South Asia) and 19 north (Taklimakan Desert) on the three days. The high values in South Asia were caused by 20 anthropogenic aerosols (such as biomass burning) or dust polluted by anthropogenic aerosols, while 21 the high MODIS\_AOD in the Taklimakan Desert resulted from dust. The values and areas of the 22 high MODIS\_AOD in South Asia and Taklimakan Desert on 1 May and 3 May were higher and 23 larger than those on April 28. The back-trajectories ended at Lhasa on the three days were different. 24 On 28 April, the air flows originated from the southwest (South Asia region). However, the air 25 masses on 1 and 3 May were from the northwest (Taklimakan Desert).

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27 The CALIPSO ground tracks across the TP and through South Asia and the Taklimakan Desert 28 were chosen to show the aerosol transport to the TP sites. On 28 April, the Level-1 attenuated 29 backscatter at 532 nm derived from CALIOP (the second row) showed apparent aerosol layers in 30 the southern area (Bhutan and northeast India) and this aerosol layer even extended to an altitude of 31 ~10km over the TP along the southern slope of the TP. On 1 May, the CALIOP attenuated 32 backscatter not only showed the deep aerosol layers in south of the TP but also showed stronger 33 aerosol layers north of the TP (Taklimakan Desert area). Besides, the north aerosol layers also 34 climbed into the air over the TP, but not as high as the southern aerosol layer. On 3 May, there were 35 also aerosol layers in the south and north of the TP and that were both transported to above the TP, 36 but the aerosol loading over the TP was lower than that on 28 April and 1 May (the values of 37 attenuated backscatter on 3 May was lower), which corresponds to the lower CE318\_AOD on this 38 day than those on 28 April and 1 May at the three TP sites (Figure 13).

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The vertical feature mask of the aerosol from CALIOP (the third row) showed the aerosol types on the three days. On 28 April, the aerosol layer in the north (~ 35°N) and above the TP was mainly the smoke aerosol and was even higher than 10 km. The back trajectories ended at Lhasa also showed that the southern airflow brought the smoke aerosol and polluted dust from South Asia to the centre of the TP. On 1 May, the aerosol layer on the southern slope of the TP was also smoke

1 aerosol and polluted dust, while the aerosol layers in the northern TP and above the TP were almost 2 all dust aerosol, which could be explained by the northwest airflows carrying the dust aerosol from 3 the Taklimakan Desert. These may be the result of the lower EAE values at Lhasa and NAM\_CO than that at QOMS CAM (Figure 13). On 3 May, the aerosol type above the central TP and the 4 5 southern TP was occupied by polluted dust aerosol, and the EAE at NAM\_CO and QOMS\_CAM 6 also showed a slight increase on 3 May. These results agree with the aerosol simulation from GEOS-7 Chem. Jia et al. (2015) has shown that the dust from India polluted by anthropogenic aerosols can 8 be transported to the TP, but the back trajectories on 1 and 3 May illustrated that the airflows that 9 ended at Lhasa were from the north or northwest rather than the south, indicating that the polluted 10 dust over the TP on 3 May was more likely the mixing result of dust and smoke aerosol. In addition, 11 the lengths of the back trajectories (especially the back trajectories at 10 m and 500 m above ground 12 level) on 1 May showed that the airflows moved slowly, which allowed the possibility of aerosol 13 mixture over the TP. The observations and model simulations illustrated the following scene: first, 14 the smoke aerosol in South Asia was lifted up to 10 km, contaminating the TP sites, and transported 15 to the centre of the TP; then, the dust from the Taklimakan Desert could climb the north slope of the TP and be transported to the TP; finally, the dust and smoke aerosol over the TP were mixed. This 16 17 case of aerosol pollution shows that the anthropogenic aerosols (mainly smoke) in South Asia and 18 dust in the Taklimakan Desert could be transported to the centre of the TP and they both even can 19 cause mixed aerosol pollution above the TP. The past case studies of aerosol transport to the TP are 20 almost individual dust or smoke aerosol, while this case of aerosol pollution over the TP showed 21 the mixing pollution during the last two days of the case period.

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#### 23 6. Conclusion

The long-term spatiotemporal variations in the aerosol optical properties and the impacts of the long-range aerosol transport over the TP were analysed by using a combination of ground-based and satellite remote sensing aerosol products as well as model simulations. The major conclusions are drawn as follows:

- (1) The annual CE318\_AOD at most TP sites showed increasing trends (0-0.013/year) during the
   past decade. Increasing tendencies in the annual averaged EAE were also found at most TP sites.
   Spatially, the MODIS\_AOD showed negative trends in the northwest edge close to the
   Taklimakan Desert and the east of Qaidam Basin and slightly positive trends in most of the
   other areas of the TP.
- (2) Different aerosol types and sources contributed to the high aerosol loading at the five sites: dust
   was dominant in Lhasa, Mt\_WLG and Muztagh with sources from the Taklimakan Desert, but
   fine aerosol pollution was dominant at NAM\_CO and QOMS\_CAS with the transport from
   South Asia.
- (3) A case of smoke followed by dust pollution at Lhasa, NAM\_CO and QOMS\_CAS during 28
  April 3 May 2016 showed that the smoke aerosol in South Asia was first uplifted to 10 km
  and transported to the centre of TP. Then, the dust from the Taklimakan Desert could climb the
  northern slope of the TP and be transported to the TP, allowing the dust and smoke aerosol over
  the TP to mix.
- 42

43 There are some limitations in this study. First, ground-based remote sensing and MODIS\_AOD 44 products may have had missing data due to clouds interference. Second, only half of a year of 1 observations at the Muztagh\_Ata station may not be sufficient to fully reveal pollution days in the 2 northwest TP region, which could have affected the statistics to some extent. More long-term in situ 3 observations are needed in the TP. However, due to the remoteness and challenging weather conditions over the plateau, maintaining long-term in situ observation stations over the TP is very 4 5 difficult. The numerical model simulation is more practically feasible to study the aerosol properties 6 over the TP, but the model accuracy is far from ideal over the TP. Thus, long-term numerical model 7 simulation coupled with satellite observations and intensive short-term field campaigns should be 8 used to analyse the aerosol properties over the TP in the future.

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10 Data availability:

11 The four sites (Mt\_WLG, Muztagh\_Ata, NAM\_CO and QOMS\_CAS) data are available from 12 AERONET website (https://aeronet.gsfc.nasa.gov/). The dataset of Lhasa used in the study can be 13 requested by contacting the corresponding author. The MODIS aerosol products are available from 14 http://ladsweb.nascom.nasa.gov. The HYSPLIT model and meteorological fields' data can be from 15 https://www.arl.noaa.gov/hysplit/. The CALIPSO data are from https://eosweb.larc.nasa.gov. 16 GEOS-Chem model code and share data can be obtained from http://wiki.seas.harvard.edu/geos-17 chem.

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19 Competing interests.

The authors declare that they have no conflict of interest.

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22 Author contribution:

All authors help to shape the ideas and review this manuscript. JZ, XX and HC designed, and wrote the manuscript; JZ, XX, HC, JW help to analyze the data; HC, XZ, SK and ZC carried out the sunphotometer observations; JW, ZC, SK, TZ, XY, and YZ provided constructive comments on this study.

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## 1 **References**

- 2 Ångström, A.: On the atmospheric transmission of Sun radiation and on dust in the air, Geografiska
- 3 Annaler, 11, 156-166, <u>https://doi.org/10.1080/20014422.1929.11880498</u>, 1929.

4 Bey, I., Jacob, D. J., Yantosca, R. M., Logan, J. A., Field, B. D., Fiore, A. M., Li, Q., Liu, H. Y.,

5 Mickley, L. J., and Schultz, M. G.: Global modeling of tropospheric chemistry with assimilated

- 6 meteorology: Model description and evaluation, Journal of Geophysical Research: Atmospheres,
- 7 106, 23073-23095, <u>https://doi.org/10.1029/2001jd000807</u>, 2001.
- 8 Bilal, M., Nazeer, M., Qiu, Z., Ding, X., and Wei, J.: Global Validation of MODIS C6 and C6.1
- 9 Merged Aerosol Products over Diverse Vegetated Surfaces, Remote Sens-Basel, 10, 475,
- 10 https://doi.org/10.3390/rs10030475, 2018.
- 11 Bilal, M., and Qiu, Z.: Evaluation of Modis C6 Combined Aerosol Product at Global Scale, IGARSS
- 12 2018 2018 IEEE International Geoscience and Remote Sensing Symposium, 2018, 9126-9129,
- 13 Bilal, M., Nazeer, M., Nichol, J., Qiu, Z., Wang, L., Bleiweiss, M. P., Shen, X., Campbell, J. R., and
- 14 Lolli, S.: Evaluation of Terra-MODIS C6 and C6.1 Aerosol Products against Beijing, XiangHe, and

15 Xinglong AERONET Sites in China during 2004-2014, Remote Sens-Basel, 11, 486,
16 <u>https://doi.org/doi:10.3390/rs11050486</u>, 2019.

- 17 Che, H., Zhang, X. Y., Xia, X., Goloub, P., Holben, B., Zhao, H., Wang, Y., Zhang, X. C., Wang, H.,
- 18 Blarel, L., Damiri, B., Zhang, R., Deng, X., Ma, Y., Wang, T., Geng, F., Qi, B., Zhu, J., Yu, J., Chen,
- 19 Q., and Shi, G.: Ground-based aerosol climatology of China: aerosol optical depths from the China
- 20 Aerosol Remote Sensing Network (CARSNET) 2002–2013, Atmospheric Chemistry and Physics,
- 21 15, 7619-7652, <u>https://doi.org/10.5194/acp-15-7619-2015</u>, 2015.
- Che, H. Z., Wang, Y. Q., and Sun, J. Y.: Aerosol optical properties at Mt. Waliguan Observatory,
  China, Atmospheric Environment, 45, 6004-6009, <u>https://doi.org/10.1016/j.atmosenv.2011.07.050</u>,
  2011.
- Chen, J., Xin, J., An, J., Wang, Y., Liu, Z., Chao, N., and Meng, Z.: Observation of aerosol optical
   properties and particulate pollution at background station in the Pearl River Delta region,
- 27 Atmospheric Research, 143, 216-227, https://doi.org/10.1016/j.atmosres.2014.02.011, 2014.
- Cong, Z., Kang, S., Liu, X., and Wang, G.: Elemental composition of aerosol in the Nam Co region,
  Tibetan Plateau, during summer monsoon season, Atmospheric Environment, 41, 1180-1187,
  <a href="https://doi.org/10.1016/j.atmosenv.2006.09.046">https://doi.org/10.1016/j.atmosenv.2006.09.046</a>, 2007.
- Cong, Z., Kang, S., Smirnov, A., and Holben, B.: Aerosol optical properties at Nam Co, a remote
  site in central Tibetan Plateau, Atmospheric Research, 92, 42-48,
  https://doi.org/10.1016/j.atmosres.2008.08.005, 2009.
- Draxler, R. R., and Hess, G. D.: An overview of the HYSPLIT-4 Modelling system for trajectories,
   dispersion and deposition, Australian Meteorological Magazine, 47, 295-308, 1998.
- 36 Du, W., Sun, Y. L., Xu, Y. S., Jiang, Q., Wang, Q. Q., Yang, W., Wang, F., Bai, Z. P., Zhao, X. D.,

- 1 and Yang, Y. C.: Chemical characterization of submicron aerosol and particle growth events at a
- 2 national background site (3295 m a.s.l.) on the Tibetan Plateau, Atmospheric Chemistry and Physics,
- 3 15, 10811-10824, <u>https://doi.org/10.5194/acp-15-10811-2015</u>, 2015.

# 4 Dubovik, O., and King, M. D.: A flexible inversion algorithm for retrieval of aerosol optical

- 5 properties from Sun and sky radiance measurements, Journal of Geophysical Research:
- 6 Atmospheres, 105, 20673-20696, <u>https://doi.org/10.1029/2000jd900282</u>, 2000.
- 7 Dubovik, O., Holben, B., Eck, T. F., Smirnov, A., Kaufman, Y. J., King, M. D., Tanré, D., and
- 8 Slutsker, I.: Variability of Absorption and Optical Properties of Key Aerosol Types Observed in
- 9 Worldwide Locations, Journal of the Atmospheric Sciences, 59, 590-608,
- 10 https://doi.org/10.1175/1520-0469(2002)059<0590:voaaop>2.0.co;2, 2002.
- 11 Dubovik, O., Sinyuk, A., Lapyonok, T., Holben, B. N., Mishchenko, M., Yang, P., Eck, T. F., Volten,

H., Munoz, O., Veihelmann, B., van der Zande, W. J., Leon, J. F., Sorokin, M., and Slutsker, I.:
Application of spheroid models to account for aerosol particle nonsphericity in remote sensing of
desert dust, J Geophys Res-Atmos, 111, D11208.11201-D11208.11234,

- 15 <u>https://doi.org/10.1029/2005jd006619</u>, 2006.
- Eck, T. F., Holben, B. N., Reid, J. S., Dubovik, O., Smirnov, A., O'Neill, N. T., Slutsker, I., and
  Kinne, S.: wavelength dependence of the optical depth of biomass burning, urban, and desert dust
  aerosols, Journal of Geophysical Research, 104, 31,333-331,349,
  https://doi.org/10.1029/1999JD900923, 1999.
- 20 Eck, T. F., Holben, B. N., Sinyuk, A., Pinker, R. T., Goloub, P., Chen, H., Chatenet, B., Li, Z., Singh,
- 21 R. P., Tripathi, S. N., Reid, J. S., Giles, D. M., Dubovik, O., O'Neill, N. T., Smirnov, A., Wang, P.,
- 22 and Xia, X.: Climatological aspects of the optical properties of fine/coarse mode aerosol mixtures,
- 23 J Geophys Res-Atmos, 115, <u>https://doi.org/10.1029/2010jd014002</u>, 2010.
- Fan, X., Chen, H., Goloub, P., Xia, X., Zhang, W., and Chatenet, B.: Analysis of column-integrated
   aerosol optical thickness in Beijing from AERONET observations, China Particulogy, 4, 330-335,
- 26 https://doi.org/10.1016/S1672-2515(07)60285-1, 2006.
- 27 Giles, D. M., Holben, B. N., Eck, T. F., Sinyuk, A., Smirnov, A., Slutsker, I., Dickerson, R. R.,

Thompson, A. M., and Schafer, J. S.: An analysis of AERONET aerosol absorption properties and
classifications representative of aerosol source regions, J Geophys Res-Atmos, 117, 127-135,
https://doi.org/10.1029/2012JD018127, 2012.

- Giles, D. M., Sinyuk, A., Sorokin, M. G., Schafer, J. S., Smirnov, A., Slutsker, I., Eck, T. F., Holben,
  B. N., Lewis, J. R., Campbell, J. R., Welton, E. J., Korkin, S. V., and Lyapustin, A. I.: Advancements
  in the Aerosol Robotic Network (AERONET) Version 3 database automated near-real-time
  quality control algorithm with improved cloud screening for Sun photometer aerosol optical
  depth (AOD) measurements, Atmos Meas Tech, 12, 169-209, <a href="https://doi.org/10.5194/amt-12-169-2019">https://doi.org/10.5194/amt-12-169-2019</a>, 2019.
- 37 Heald, C. L., Ridley, D. A., Kroll, J. H., Barrett, S. R. H., Cady-Pereira, K. E., Alvarado, M. J., and

- 1 Holmes, C. D.: Contrasting the direct radiative effect and direct radiative forcing of aerosols,
- 2 Atmospheric Chemistry and Physics, 14, 5513-5527, https://doi.org/10.5194/acp-14-5513-2014,
- 3 2014.
- 4 Huang, J., Minnis, P., Yi, Y., Tang, Q., Wang, X., Hu, Y., Liu, Z., Ayers, K., Trepte, C., and Winker,
- 5 D.: Summer dust aerosols detected from CALIPSO over the Tibetan Plateau, Geophys Res Lett, 34,
  6 529-538, https://doi.org/10.1029/2007gl029938, 2007.
- Jia, R., Liu, Y., Chen, B., Zhang, Z., and Huang, J.: Source and transportation of summer dust over
  the Tibetan Plateau, Atmospheric Environment, 123, 210-219,
  https://doi.org/10.1016/j.atmosenv.2015.10.038, 2015.
- 10 Kopacz, M., Mauzerall, D., Wang, J., Leibensperger, E., Henze, D., and Singh, K.: Origin and
- 11 radiative forcing of black carbon transported to the Himalayas and Tibetan Plateau, Atmospheric
- 12 Chemistry and Physics, 11, 2837-2852, <u>https://doi.org/10.5194/acp-11-2837-2011</u>, 2011.
- 13 Kumar, A., Singh, N., Anshumali, and Solanki, R.: Evaluation and utilization of MODIS and
- 14 CALIPSO aerosol retrievals over a complex terrain in Himalaya, Remote Sensing of Environment,
- 15 206, 139-155, <u>https://doi.org/10.1016/j.rse.2017.12.019</u>, 2018.
- Lau, K. M., Kim, M. K., and Kim, K. M.: Asian summer monsoon anomalies induced by aerosol
  direct forcing: the role of the Tibetan Plateau, Clim Dynam, 26, 855-864,
  https://doi.org/10.1007/s00382-006-0114-z, 2006.
- 19 Lee, W.-S., Bhawar, R. L., Kim, M.-K., and Sang, J.: Study of aerosol effect on accelerated snow
- melting over the Tibetan Plateau during boreal spring, Atmospheric Environment, 75, 113-122,
   https://doi.org/10.1016/j.atmosenv.2013.04.004, 2013.
- 22 Levy, R. C., Mattoo, S., Munchak, L. A., Remer, L. A., Sayer, A. M., Patadia, F., and Hsu, N. C.:
- The Collection 6 MODIS aerosol products over land and ocean, Atmos Meas Tech, 6, 2989-3034,
  https://doi.org/10.5194/amt-6-2989-2013, 2013.
- Li, J., Huang, J., Stamnes, K., Wang, T., Lv, Q., and Jin, H.: A global survey of cloud overlap based
  on CALIPSO and CloudSat measurements, Atmospheric Chemistry and Physics, 15, 519-536,
  https://doi.org/10.5194/acp-15-519-2015, 2015.
- Li, J., Lv, Q., Zhang, M., Wang, T., Kawamoto, K., Chen, S., and Zhang, B.: Effects of atmospheric
  dynamics and aerosols on the fraction of supercooled water clouds, Atmospheric Chemistry and
  Physics, 17, 1847-1863, https://doi.org/10.5194/acp-17-1847-2017, 2017.
- Li, J., Jian, B., Huang, J., Hu, Y., Zhao, C., Kawamoto, K., Liao, S., and Wu, M.: Long-term variation of cloud droplet number concentrations from space-based Lidar, Remote Sensing of Environment, 213, 144-161, https://doi.org/10.1016/j.rse.2018.05.011, 2018.
- 34 Li, M., Zhang, Q., Streets, D. G., He, K. B., Cheng, Y. F., Emmons, L. K., Huo, H., Kang, S. C., Lu,
- 35 Z., Shao, M., Su, H., Yu, X., and Zhang, Y.: Mapping Asian anthropogenic emissions of non-
- 36 methane volatile organic compounds to multiple chemical mechanisms, Atmospheric Chemistry and

- 1 Physics, 14, 5617-5638, <u>https://doi.org/10.5194/acp-14-5617-2014</u>, 2014.
- 2 Liu, Y., Sato, Y., Jia, R., Xie, Y., Huang, J., and Nakajima, T.: Modeling study on the transport of
- 3 summer dust and anthropogenic aerosols over the Tibetan Plateau, Atmospheric Chemistry and
- 4 Physics, 15, 12581-12594, https://doi.org/10.5194/acp-15-12581-2015, 2015.
- Liu, Z., Liu, D., Huang, J., Vaughan, M., Uno, I., Sugimoto, N., Kittaka, C., Trepte, C., Wang, Z.,
  Hostetler, C., and Winker, D.: Airborne dust distributions over the Tibetan Plateau and surrounding
  areas derived from the first year of CALIPSO lidar observations, Atmospheric Chemistry and
  Physics, 8, 5045-5060, https://doi.org/10.5194/acp-8-5045-2008, 2008.
- 9 Lu, Z., Streets, D. G., Zhang, Q., and Wang, S.: A novel back-trajectory analysis of the origin of
  10 black carbon transported to the Himalayas and Tibetan Plateau during 1996-2010, Geophys Res Lett,
  11 39, n/a-n/a, https://doi.org/10.1029/2011gl049903, 2012.
- 12 Ma, Y., Li, Z., Li, Z., Xie, Y., Fu, Q., Li, D., Zhang, Y., Xu, H., and Li, K.: Validation of MODIS
- 13 Aerosol Optical Depth Retrieval over Mountains in Central China Based on a Sun-Sky Radiometer
- 14 Site of SONET, Remote Sens-Basel, 8, 111, <u>https://doi.org/10.3390/rs8020111</u>, 2016.
- Martin, R. V., Jacob, D. J., Yantosca, R. M., Chin, M., and Ginoux, P.: Global and regional decreases
   in tropospheric oxidants from photochemical effects of aerosols, J Geophys Res-Atmos, 108,
   <u>https://doi.org/10.1029/2002jd002622</u>, 2003.
- 18 Pan, L., Che, H., Geng, F., Xia, X., Wang, Y., Zhu, C., Chen, M., Gao, W., and Guo, J.: Aerosol
- 19 optical properties based on ground measurements over the Chinese Yangtze Delta Region,
- 20 Atmospheric Environment, 44, 2587-2596, <u>https://doi.org/10.1016/j.atmosenv.2010.04.013</u>, 2010.
- 21 Qian, Y., Flanner, M. G., Leung, L. R., and Wang, W.: Sensitivity studies on the impacts of Tibetan
- 22 Plateau snowpack pollution on the Asian hydrological cycle and monsoon climate, Atmospheric
- 23 Chemistry and Physics, 11, 1929-1948, <u>https://doi.org/10.5194/acp-11-1929-2011</u>, 2011.
- 24 Sayer, A. M., Munchak, L. A., Hsu, N. C., Levy, R. C., Bettenhausen, C., and Jeong, M. J.: MODIS
- 25 Collection 6 aerosol products: Comparison between Aqua's e-Deep Blue, Dark Target, and "merged"
- 26 data sets, and usage recommendations, Journal of Geophysical Research: Atmospheres, 119,
- 27 13,965-913,989, <u>https://doi.org/10.1002/2014jd022453</u>, 2014.
- Shen, R. Q., Ding, X., He, Q. F., Cong, Z. Y., Yu, Q. Q., and Wang, X. M.: Seasonal variation of
  secondary organic aerosol tracers in Central Tibetan Plateau, Atmospheric Chemistry and Physics,
  15, 8781-8793, https://doi.org/10.5194/acp-15-8781-2015, 2015.
- Takemura, T., Nozawa, T., Emori, S., Nakajima, T. Y., and Nakajima, T.: Simulation of climate response to aerosol direct and indirect effects with aerosol transport-radiation model, Journal of
- 33 Geophysical Research, 110, -, <u>https://doi.org/10.1029/2004jd005029</u>, 2005.
- Tobo, Y., Iwasaka, Y., Shi, G.-Y., Kim, Y.-S., Ohashi, T., Tamura, K., and Zhang, D.: Balloon-borne
  observations of high aerosol concentrations near the summertime tropopause over the Tibetan
  Plateau, Atmospheric Research, 84, 233-241, https://doi.org/10.1016/j.atmosres.2006.08.003, 2007.

- 1 van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Mu, M., Kasibhatla, P. S., Morton,
- 2 D. C., DeFries, R. S., Jin, Y., and van Leeuwen, T. T.: Global fire emissions and the contribution of
- 3 deforestation, savanna, forest, agricultural, and peat fires (1997–2009), Atmospheric Chemistry and
- 4 Physics, 10, 11707-11735, <u>https://doi.org/10.5194/acp-10-11707-2010</u>, 2010.
- 5 Wan, X., Kang, S., Wang, Y., Xin, J., Liu, B., Guo, Y., Wen, T., Zhang, G., and Cong, Z.: Size
- 6 distribution of carbonaceous aerosols at a high-altitude site on the central Tibetan Plateau (Nam Co
- 7 Station, 4730ma.s.l.), Atmospheric Research, 153, 155-164,
- 8 <u>https://doi.org/10.1016/j.atmosres.2014.08.008</u>, 2015.
- 9 Wang, P., Che, H. Z., Zhang, X. C., Song, Q. L., Wang, Y. Q., Zhang, Z. H., Dai, X., and Yu, D. J.:
- 10 Aerosol optical properties of regional background atmosphere in Northeast China, Atmospheric
- 11 Environment, 44, 4404-4412, <u>https://doi.org/10.1016/j.atmosenv.2010.07.043</u>, 2010.
- Wang, X., Ren, J., Gong, P., Wang, C., Xue, Y., Yao, T., and Lohmann, R.: Spatial distribution of
  the persistent organic pollutants across the Tibetan Plateau and its linkage with the climate systems:
  a 5-year air monitoring study, Atmospheric Chemistry and Physics, 16, 6901-6911,
  https://doi.org/10.5194/acp-16-6901-2016, 2016.
- 16 Wang, Y. X., McElroy, M. B., Jacob, D. J., and Yantosca, R. M.: A nested grid formulation for
- chemical transport over Asia: Applications to CO, Journal of Geophysical Research: Atmospheres,
- 18 109, n/a-n/a, <u>https://doi.org/10.1029/2004jd005237</u>, 2004.
- 19 Winker, D. M., Pelon, J., Coakley, J. A., Ackerman, S. A., Charlson, R. J., Colarco, P. R., Flamant,
- 20 P., Fu, Q., Hoff, R. M., Kittaka, C., Kubar, T. L., Le Treut, H., McCormick, M. P., Megie, G., Poole,
- 21 L., Powell, K., Trepte, C., Vaughan, M. A., and Wielicki, B. A.: THE CALIPSO MISSION A Global
- 3D View of Aerosols and Clouds, B Am Meteorol Soc, 91, 1211-1229,
   https://doi.org/10.1175/2010bams3009.1, 2010.
- Xia, X., Che, H., Zhu, J., Chen, H., Cong, Z., Deng, X., Fan, X., Fu, Y., Goloub, P., Jiang, H., Liu,
  Q., Mai, B., Wang, P., Wu, Y., Zhang, J., Zhang, R., and Zhang, X.: Ground-based remote sensing
  of aerosol climatology in China: Aerosol optical properties, direct radiative effect and its
  parameterization, Atmospheric Environment, 124, 243-251,
  https://doi.org/10.1016/j.atmosenv.2015.05.071, 2016.
- Xia, X. G., Wang, P. C., Wang, Y. S., Li, Z. Q., Xin, J. Y., Liu, J., and Chen, H. B.: Aerosol optical
  depth over the Tibetan Plateau and its relation to aerosols over the Taklimakan Desert, Geophys Res
  Lett, 35, 96-106, https://doi.org/10.1029/2008gl034981, 2008.
- Xia, X. G., Zong, X. M., Cong, Z. Y., Chen, H. B., Kang, S. C., and Wang, P. C.: Baseline continental
   aerosol over the central Tibetan plateau and a case study of aerosol transport from South Asia,
- 34 Atmospheric Environment, 45, 7370-7378, <u>https://doi.org/10.1016/j.atmosenv.2011.07.067</u>, 2011.
- 35 Xin, J. Y., Wang, Y. S., Li, Z. Q., Wang, P. C., Hao, W. M., Nordgren, B. L., Wang, S. G., Liu, G. R.,
- 36 Wang, L. L., Wen, T. X., Sun, Y., and Hu, B.: Aerosol optical depth (AOD) and Angstrom exponent
- of aerosols observed by the Chinese Sun Hazemeter Network from August 2004 to September 2005,

- 1 J Geophys Res-Atmos, 112, 1703-1711, <u>https://doi.org/10.1029/2006jd007075</u>, 2007.
- 2 Xing, C., Liu, C., Wang, S., Chan, K. L., Gao, Y., Huang, X., Su, W., Zhang, C., Dong, Y., Fan, G.,
- 3 Zhang, T., Chen, Z., Hu, Q., Su, H., Xie, Z., and Liu, J.: Observations of the vertical distributions
- 4 of summertime atmospheric pollutants and the corresponding ozone production in Shanghai, China,
- 5 Atmospheric Chemistry and Physics, 17, 14275-14289, <u>https://doi.org/10.5194/acp-17-14275-2017</u>,
- 6 2017.
- 7 Yang, R., Wang, J., Zhang, T., and He, S.: Change in the relationship between the Australian summer
- 8 monsoon circulation and boreal summer precipitation over Central China in the late 1990s,
- 9 Meteorology and Atmospheric Physics, 131, 105-113, https://doi.org/10.1007/s00703-017-0556-3,
- 10 2017a.
- Yang, R. W., Tao, Y., and Cao, J.: A Mechanism for the Interannual Variation of the Early Summer
   East Asia-Pacific Teleconnection Wave Train, Acta Meteorol Sin, 24, 452-458, 2010.
- Yang, Y., Russell, L. M., Lou, S., Liao, H., Guo, J., Liu, Y., Singh, B., and Ghan, S. J.: Dust-wind
  interactions can intensify aerosol pollution over eastern China, Nature communications, 8, 15333,
  https://doi.org/10.1038/ncomms15333, 2017b.
- 16 Zhang, B., Wang, Y., and Hao, J.: Simulating aerosol–radiation–cloud feedbacks on meteorology
- 17 and air quality over eastern China under severe haze conditions in winter, Atmospheric Chemistry
- 18 and Physics, 15, 2387-2404, <u>https://doi.org/10.5194/acp-15-2387-2015</u>, 2015.
- Zhang, N., Cao, J., Ho, K., and He, Y.: Chemical characterization of aerosol collected at Mt. Yulong
  in wintertime on the southeastern Tibetan Plateau, Atmospheric Research, 107, 76-85,
  https://doi.org/10.1016/j.atmosres.2011.12.012, 2012.
- Zhang, X. Y., Arimoto, R., Cao, J. J., An, Z. S., and Wang, D.: Atmospheric dust aerosol over the
  Tibetan Plateau, Journal of Geophysical Research: Atmospheres, 106, 18471-18476,
  https://doi.org/10.1029/2000jd900672, 2001.
- Zhao, Z., Cao, J., Shen, Z., Xu, B., Zhu, C., Chen, L. W. A., Su, X., Liu, S., Han, Y., Wang, G., and
  Ho, K.: Aerosol particles at a high-altitude site on the Southeast Tibetan Plateau, China: Implications
  for pollution transport from South Asia, Journal of Geophysical Research: Atmospheres, 118,
  11,360-311,375, https://doi.org/10.1002/jgrd.50599, 2013.
- Zhu, J., Che, H., Xia, X., Chen, H., Goloub, P., and Zhang, W.: Column-integrated aerosol optical
  and physical properties at a regional background atmosphere in North China Plain, Atmospheric
- 31 Environment, 84, 54-64, <u>https://doi.org/10.1016/j.atmosenv.2013.11.019</u>, 2014.
- Zhu, J., Xia, X., Che, H., Wang, J., Zhang, J., and Duan, Y.: Study of aerosol optical properties at
   Kunming in southwest China and long-range transport of biomass burning aerosols from North
- 34 Burma, Atmospheric Research, 169, 237-247, <u>https://doi.org/10.1016/j.atmosres.2015.10.012</u>, 2016.
- Zhu, J., Xia, X., Wang, J., Zhang, J., Wiedinmyer, C., Fisher, J. A., and Keller, C. A.: Impact of
   Southeast Asian smoke on aerosol properties in Southwest China: First comparison of model

- 1 simulations with satellite and ground observations, Journal of Geophysical Research: Atmospheres,
- 2 122, 3904-3919, <u>https://doi.org/10.1002/2016jd025793</u>, 2017.

Zhuang, B. L., Wang, T. J., Liu, J., Li, S., Xie, M., Han, Y., Chen, P. L., Hu, Q. D., Yang, X. Q., Fu,
C. B., and Zhu, J. L.: The surface aerosol optical properties in the urban area of Nanjing, west
Yangtze River Delta, China, Atmospheric Chemistry and Physics, 17, 1143-1160,
https://doi.org/10.5194/acp-17-1143-2017, 2017.

7 Zhuang, B. L., Wang, T. J., Liu, J. N. E., Che, H. Z., Han, Y., Fu, Y., Li, S., Xie, M., Li, M. M., Chen,

8 P. L., Chen, H. M., Yang, X. Q., and Sun, J. N.: The optical properties, physical properties and direct

9 radiative forcing of urban columnar aerosols in the Yangtze River Delta, China, Atmospheric

10 Chemistry and Physics, 18, 1419-1436, <u>https://doi.org/10.5194/acp-18-1419-2018</u>, 2018.

11

#### **1** Figure captions

- 2 Figure 1. Topography of the Tibetan Plateau (TP) and the five CE318 stations located in the TP
- 3 (Lhasa, Mt\_WLG, Mutztagh\_Ata, NAM\_CO, and QOMS\_CAS).
- 4 Figure 2. Box plots of the monthly AOD and EAE at the five sites located on the Tibetan Plateau,
- 5 i.e., Lhasa, Mt\_WLG, Muztagh\_Alt, NAM\_CO, and QOMS\_CAS. In each box, the red-line in the
- 6 centre is the median and the lower and upper limits are the first and the third quartiles, respectively.
- 7 The lines extending vertically from the box indicate the spread of the distribution with the length
- 8 being 1.5 times the difference between the first and the third quartiles. The asterisk symbols indicate
- 9 the geometric means in each month. The annual mean values and standard errors are also shown in
- 10 each subgraph.
- 11 Figure 3. Seasonal variation in aerosol size distribution at the five sites located in Tibetan Plateau.
- Figure 4. Annual averages of and trends in aerosol optical depth (AOD) and Extinction Ångstrom
   exponent (EAE) at four sites located in Tibetan Plateau.
- 14 Figure 5. Comparisons of the 550 nm AOD measured by the CE318 instrument (CE318\_AOD) over
- 15 Tibetan Plateau stations with the MODIS retrieval Deep-Blue/Dark-Target combined AOD of 10
- 16 km spatial resolutions (MODIS AOD). The statistical parameters in this figure include the number
- 17 of matchup data (N), the slope and intercept at the y-axis of linear regression (read line), the mean
- 18 bias (MB), root mean squared error (RMSE), correlation coefficient (R), and the percentage of data
- 19 within the expected error 0.05+0.15AOD (%EE) which is used as the MODIS AOD expected
- 20 uncertainty over land (green lines).
- 21 Figure 6. Spatial distribution of MODIS C6 AOD at 550 nm over the Tibetan Plateau (only the
- altitude > 3000 m) during 2006-2017. The color-filled circles are the CE318 observation AOD
- 23 averages at TP sites.
- Figure 7. The seasonal departure of MODIS AOD over the TP (altitude > 3000 m).
- Figure 8. Trend in the MODIS AOD at 550 nm during 2006-2017.
- Figure 9. Trends in the MODIS AOD at 550 nm during 2006-2017 in each season.
- Figure 10. AOD vs EAE (only CE318 AOD at 440 nm > 0.4 was considered) observed by CE318
- at the five sites on the Tibetan Plateau.
- Figure 11. Aerosol size distribution binned by CE318 AOD at the five sites on the Tibetan Plateau.
- 30 Figure 12. The back-trajectories ended at the five sites (10 m above ground level) on the TP overlaid
- 31 with the mean MODIS C6 AOD at 550 nm on the aerosol pollution day observed by ground-based
- 32 CE318 (CE318 AOD >0.4). Red stands for EAE >1.0, black for EAE < 0.5, and green for EAE
- 33 within 0.5-1.0.
- Figure 13. CE318 observed daily AOD at 440 nm and EAE during 27April, 2016 3May, 2016 at
- 35 Lhasa, NAM\_CO and QOMS\_CAS.
- 36 Figure 14. The GEOS-Chem model simulated the daily average AOD vs CE318 observed daily
- 37 AOD at 550 nm, and the ratios of dust or organic carbon (OC) and black carbon (BC) aerosol to the

- 1 total AOD during 27April, 2016 3 May, 2016 at Lhasa, NAM\_CO and QOMS\_CAS. The statistical
- 2 parameters used in the modal evaluation are the same as those in Figure 5.
- 3 Figure 15. The MODIS C6 AOD at 550 nm and 72-hour back trajectories ended at Lhasa at three
- 4 heights above the ground level (10 m in black, 500 m in red and 1000 m in blue lines) (the first
- 5 row); the CALIOP-derived vertical profile of total attenuated backscatter at 532 nm (the second
- 6 row); and the vertical feature mask of aerosol on 28 April, 1 May, and 3 May, 2016 over the
- 7 ground track shown in the first row (green lines) (the third row).
- 8



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- 4 95% significance.
- 5





Figure 5. Comparisons of the 550 nm AOD measured by the CE318 instrument (CE318\_AOD)
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Site name	Lat(° N)	Lon(° E)	° E) Site description, observation days and period			
Lhasa	29.50	91.13	Urban station on the Tibetan Plateau, 3648 m a.s.l.			
			1554 days, 2007.05~2017.12			
Mt_WLG	36.28	100.90	Mountain, 3816 m a.s.l., 314 days, 2009.09~2013.07			
Muztagh_Ata	38.41	75.04	Mountain, 3674 m a.s.l., 84 days, 2011.06~2011.10			
NAM_CO	30.77	90.96	Mountain, 4740 m a.s.l., 1061 days, 2006.08~2016.08			
QOMS_CAS	28.36	86.95	Mountain, 4276 m a.s.l., 1623 days, 2009.10~2017.11			

Table 1. Site location and description. 

 $1 \qquad \text{Table 2. Seasonal aerosol optical depth (AOD_{440nm}) and extinction Angstrom exponent (EAE_{440-870nm}) at}$ 

Site	AOD				EAE			
	MAM	JJA	SON	DJF	MAM	JJA	SON	DJF
Lhasa	0.16+0.	0.12+0.	0.10+0.	0.09+0.	0.72+0.	0.97+0.	1.11+0.	0.91+0.
	10	08	18	08	37	40	38	52
Mt_WLG	0.13+0.	0.14+0.	0.08+0.	0.08+0.	0.37+0.	0.65+0.	1.04+0.	0.58+0.
	16	07	11	07	38	40	80	69
Muztagh_	NaN	0.14+0.	0.14+0.	NaN	NaN	0.73+0.	0.64+0.	NaN
Ata		06	05			30	27	
NAM_CO	0.07+0.	0.06+0.	0.03+0.	0.03+0.	0.63+0.	0.62+0.	0.65+0.	0.78+0.
	07	04	05	01	44	45	32	43
QOMS_C	0.08+0.	0.06+0.	0.03+0.	0.03+0.	1.04+0.	0.76+0.	0.85+0.	1.10+0.
AS	06	04	01	02	38	43	51	67

2 the five sites in the TP.

3

% EAE>1.0/N Site N of AOD>0.4 % EAE<0.5/N % 0.5<EAE<1.0/N 36.0 Lhasa 655 60.6 3.4 Mt\_WLG 290 73.4 0 26.6 Muztagh\_Ata 5 100 0 0 27.9 NAM\_CO 140 2.8 69.3 QOMS\_CAS 59 23.7 0 76.3

1 Table 3. The percentages of EAE <0.5, 0.5-1.0, and >1.0 for high AOD observations at the five sites.