# Observations of the vertical distributions of summertime atmospheric pollutants in Nam Co: OH production and source analysis

Chengzhi Xing<sup>1</sup>, Cheng Liu<sup>1,2,3,4,\*</sup>, Chunxiang Ye<sup>5,\*</sup>, Jingkai Xue<sup>6</sup>, Hongyu Wu<sup>6</sup>, Xiangguang Ji<sup>7</sup>, Jinping Ou<sup>8</sup>, and Qihou Hu<sup>1</sup>

<sup>1</sup> Key Lab of Environmental Optics & Technology, Anhui Institute of Optics and Fine Mechanics, Hefei Institutes of Physical Science, Chinese Academy of Sciences, Hefei, 230031, China

<sup>2</sup>Department of Precision Machinery and Precision Instrumentation, University of Science and Technology of China, Hefei, 230026, China

<sup>3</sup> Center for Excellence in Regional Atmospheric Environment, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen, 361021, China

<sup>4</sup> Key Laboratory of Precision Scientific Instrumentation of Anhui Higher Education Institutes, University of Science and Technology of China, Hefei, 230026, China

<sup>5</sup> College of Environmental Sciences and Engineering, Peking University, 100871 Beijing

<sup>6</sup> School of Environmental Science and Optoelectronic Technology, University of Science and Technology of China, Hefei, 230026, China

<sup>7</sup> Institute of Physical Science and Information Technology, Anhui University, Hefei, 230601, China

<sup>8</sup> The Department of Health Promotion and Behavioral Sciences, School of Public Health, Anhui Medical University, Hefei, 230032, China

\*Corresponding author. E-mail: chliu81@ustc.edu.cn; c.ye@pku.edu.cn

### 1 Abstract

2 The Tibetan Plateau (TP) plays a key role in regional environment and global climate change, however, 3 the lack of vertical observation of atmospheric species, such as HONO and O<sub>3</sub>, hinders a deeper 4 understanding of the atmospheric chemistry and atmospheric oxidation capacity (AOC) on the TP. In 5 this study, we conducted multi-axis differential optical absorption spectroscopy (MAX-DOAS) 6 measurements at Nam Co, the central TP, to observe the vertical profiles of aerosol, water vapor (H<sub>2</sub>O), 7 NO<sub>2</sub>, HONO and O<sub>3</sub> from May to July 2019. In addition to NO<sub>2</sub> mainly exhibiting a Gaussian shape 8 with the maximum value appearing at 300-400 m, other four species all showed an exponential shape 9 and decreased with the increase of height. The maximum values of monthly averaged aerosol (0.17 km<sup>-1</sup>) and O<sub>3</sub> (66.71 ppb) occurred on May, H<sub>2</sub>O ( $3.68 \times 10^{17}$  molec cm<sup>-3</sup>) and HONO (0.13 ppb) 10 appeared on July, while NO<sub>2</sub> (0.39 ppb) occurred on June at 200-400 m layer. H<sub>2</sub>O, HONO and O<sub>3</sub> all 11 exhibited a multi-peak pattern, and aerosol appeared a bi-peak pattern for their averaged diurnal 12 variations. The averaged vertical profiles of OH production rates from O<sub>3</sub> and HONO all exhibited an 13 14 exponential shape decreasing with the increase of height with maximum values of 2.61 ppb/h and 0.49 ppb/h at the bottom layer, respectively. The total OH production rate contributed by HONO and O<sub>3</sub> on 15 the TP was obviously larger than that in low-altitude areas. In addition, source analysis for HONO and 16 17 O<sub>3</sub> at different height layers were conducted. The heterogeneous reaction of NO<sub>2</sub> on wet surfaces was a 18 significant source of HONO. The maximum values of HONO/NO<sub>2</sub> appeared around H<sub>2</sub>O being  $1.0 \times$ 10<sup>17</sup> molec cm<sup>-3</sup> and aerosol being lager 0.15 km<sup>-1</sup> under 1.0 km, and the maximum values usually 19 accompanied with H<sub>2</sub>O being  $1.0-2.0 \times 10^{17}$  molec cm<sup>-3</sup> and aerosol being lager 0.02 km<sup>-1</sup> at 1.0-2.0 km. 20 O<sub>3</sub> was potentially sourced from south Asian subcontinent and Himalayas through long-range transport. 21 22 Our results enrich the new understanding of vertical distribution of atmospheric components and

23 explained the strong AOC on the TP.

24

### 25 **1 Introduction**

The TP spans 2.5 million square kilometers with an average altitude of over 4000 m. Therefore, the TP 26 27 is called the "Third Pole" of the earth (Ma et al., 2020; Kang et al., 2022). It is the home to tens of thousands of glaciers and nourishes more than 10 of Asia's rivers, thus it also acts the role of "Water 28 29 Tower of Asia" (Qu et al., 2019; Ma et al., 2022). Due to its special topography, the TP is the heat source of atmosphere due the strong solar radiation, which as the driven force to profoundly affect the 30 31 regional atmospheric circulation, global weather conditions and climate change (Yanai et al., 1992; 32 Boos et al., 2010; Chen et al., 2015; Liu et al., 2022; Zhou et al., 2022). Monsoon rainfall in Asia, flood 33 over the Yangtze River valley, and El Niño in the Pacific Ocean are strongly associated with the TP (Hsu et al., 2003; Li et al., 2016; Lei et al., 2019). In addition, the cyclone circulations caused by the 34 35 TP heat source also can inhibit the diffusion of atmospheric pollutants in the areas around the TP, such as the Sichuan Basin, causing regional pollution (Zhang et al., 2019). Therefore, observations of the 36 37 atmospheric species on the TP are essential to enhance the in-depth understanding of its atmospheric 38 physicochemical processes.

However, deciphering the atmospheric environment of the TP is highly challenging and dangerous, due 39 40 to its complex topography and harsh environment (Barnett et al., 2005; Bolch et al., 2012; Cong et al., 41 2015; Kang et al., 2016). In order to unveil the feature of atmospheric composition over the TP and their corresponding climate feedback, a large number of field observation stations have been 42 43 established, and a series of field campaigns have continued to be carried out recently, especially after 44 the performance of "the Second Tibetan Plateau Scientific Expedition and Research Program" (Che and Zhao 2021; Wang et al., 2021; Ran et al., 2022). The China National Environmental Monitoring Center 45 46 (CNEMC) has established an in-situ monitoring network with more than 12 stations over the TP, such 47 as Lhasa, Shigatse, Shannan, Nyingchi, Nagqu, Ngari, Qamdo, Diqing, Aba, Guoluo, Xining, and Haixi, to continuously monitor the surface concentrations of six atmospheric components (i.e. PM<sub>10</sub>, 48 49 PM<sub>2.5</sub>, NO<sub>2</sub>, SO<sub>2</sub>, O<sub>3</sub> and CO) since 2013 (Gao et al., 2020; Li et al., 2020; Sun et al., 2021). The 50 Institute of Tibetan Plateau Research, Chinese Academy of Sciences, has also established six long-term 51 field observation stations to measure meteorological parameters and small amounts of atmospheric 52 composition (i.e. black carbon, aerosol optical density (AOD)) (Ma et al., 2020). In addition, scientists 53 are relying on advancements in satellite remote sensing technology, such as the tropospheric 54 monitoring instrument (TROPOMI), the ozone monitoring instrument (OMI), the moderate-resolution imaging spectroradiometer (MODIS) and the cloud-aerosol lidar and infrared pathfinder satellite 55 observation (CALIPSO), to monitor the spatial and temporal evolutions of atmospheric composition on 56 57 the TP (Zhu et al., 2019; Li et al., 2020; Rawat and Naja 2022). Their advantage is to obtain the column 58 densities of pollutants in a large-scale space of the TP. Although CALIPSO could detect aerosol 59 vertical profiles, the spatiotemporal resolution (i.e. ~5.0 km horizontal resolution, 0.06 km vertical resolution and ~16 d temporal resolution) is limited and the data uncertainty in the planetary boundary 60 layer (PBL) is large due to the low signal-to-noise ratio (Huang et al., 2007). However, several studies 61 62 also revealed that the formation, aging and transport processes of atmospheric composition on the TP 63 occurs not only near the ground surface but also at high altitudes (Xu et al., 2020; Xu et al., 2022). The high PBL on the TP caused by its strong solar radiation and undulating terrain promotes the 64 atmospheric exchange between the bottom troposphere and stratosphere (Yang et al., 2003; Seidel et al., 65 66 2010). Therefore, the lack of vertical profiles of hinders the understanding of the evolution of trace 67 gases and their environmental and climate effects over the TP. In recent years, balloon and lidar vertical measurements on the TP are occasionally carried out (Fang et al., 2019; Zhang et al., 2020; 68 69 Dong et al., 2022), but their limited detection species (i.e. aerosol and  $O_3$ ) and high cost are obstacles 70 that limit long-term continuous observation and the conduction of more in-depth scientific research. 71 MAX-DOAS has the technical advantage of low-cost continuous observation of multiple atmospheric 72 components (i.e. aerosol, O<sub>3</sub> and their precursors) (Wang et al., 2018; Ma et al., 2020; Cheng et al., 73 2021; Xing et al., 2021; Li et al., 2022; Cheng et al., 2023a, 2023b). Combining these data with better 74 scientific models can reduce the modeling bias and promote to better understand the physical, chemical 75 and dynamical processes.

The strong convergent airflow formed under the combined action of monsoon, subtropical anticyclone and the airflow of subtropical westerlies could promote the accumulation of  $O_3$  on the TP in summer

(Ye and Gao 1997). Therefore, several studies have revealed the high O<sub>3</sub> concentration on the TP (Li et 78 79 al., 2022; Yang et al., 2022; Yu et al., 2022). The strong solar radiation, high O<sub>3</sub> concentration and 80 relatively high humidity on the TP provide great potential for high OH production. Lin et al. (2008) and 81 Ye (2019) also confirmed that the high OH over the TP is mainly related to the reaction between  $O(^{1}D)$ and H<sub>2</sub>O. The  $O(^{1}D)$  is produced from the photolysis of O<sub>3</sub> by UV radiation. Therefore, a hypothesis of 82 83 "strong AOC over the TP" was put forward. Previous studies pointed out that HONO also play an 84 important role in AOC at low-altitude areas, and its contribution to OH can reach 40-60%, and even 85 more than 80% in the early morning (Michoud et al., 2012; Ryan et al., 2018; Xue et al., 2020). 86 However, few HONO studies on the TP have been reported. Our previous study operated at the 87 Qomolangma Atmospheric and Environmental Observation and Research Station, Chinese Academy of Sciences (OOMS-CAS) revealed that the HONO mainly distributed in the lower PBL and peaked in 88 89 summer with 1.11 ppb, which is comparable to the average level of HONO in other low-altitude areas 90 (Luo et al., 2010; Xing et al., 2021a, 2021b; Yang et al., 2021). It indicates that it is also necessary to 91 study the contribution of HONO to AOC on the TP. Furthermore, understanding the vertical 92 distribution of OH is of great significance for learning about the atmospheric chemical processes and 93 the evolution of atmospheric components on the TP (Zhou et al., 2015). Identifying the sources of O<sub>3</sub> 94 and HONO is the basis for studying the AOC on the TP. The limited researches concluded that the 95 atmospheric HONO on the TP is mainly sourced from the emissions of vehicles, biomass burning and 96 soil, except for the NO<sub>2</sub> heterogeneous reaction on aerosol surfaces (Xing et al., 2021). The lower 97 tropospheric O<sub>3</sub> on the TP is mainly dominated by local photochemical reactions, regional horizontal 98 transport, vertical mixing and the intrusion from stratosphere (Yin et al., 2017; Xu et al. 2018).

In this study, we firstly analyzed the temporal and vertical characteristics of several atmospheric components (i.e. aerosol,  $H_2O$ ,  $NO_2$ , HONO and  $O_3$ ) based on MAX-DOAS observations in Nam Co. Afterwards, the contributions of  $O_3$  and HONO to OH in the vertical space were discussed through the tropospheric ultraviolet and visible (TUV) radiative transfer model and MAX-DOAS measurements. Finally, the potential sources of  $O_3$  and HONO at different altitudes were analyzed based on the MAX-DOAS retrievals.

# 105 2 Method and methodology

### 106 **2.1 Site**

107 The Nam Co Monitoring and Research Station for Multisphere Interactions, CAS (NAMORS) 108 (30.774°N, 90.988°E; 4730 m a.s.l.) is located at the southeast banks of Nam Co lake and the foothills 109 of the northern Mt. Nyaingêntanglha (Fig. 1). The station land is covered by alpine meadows with soil 110 type of sandy silt loam. The southwest monsoon can carry abundant moisture from Indian Ocean to this 111 station in summer to increase humidity and precipitation there. Moreover, due to the summertime huge 112 evaporation from Nam Co lake, the atmospheric H<sub>2</sub>O around CAS (NAMORS) is more abundant than 113 in other areas of the TP, resulting in lush grass vegetation and making the area around this station an 114 important summertime pasture. In addition, there are not large industries and cities within 100 km of 115 the CAS (NAMORS). The closest town to CAS (NAMORS) is Dangxiong county which is about 60 km away from this station and lower about 500 m than this station. Only a small number of vehicles 116 117 pass through this area during summer tourism season. Therefore, no obvious anthropogenic sources of 118 air pollutants exist near this station. Averaged spatial distributions of AOD, O<sub>3</sub>, NO<sub>2</sub> and HCHO 119 monitored by satellite from May to July 2019 are shown in Figure S1. Elevated AOD, NO<sub>2</sub>, and O<sub>3</sub> are 120 mainly distributed in South Asian subcontinent (e.g. India and Nepal), the southern foothills of the 121 Himalayas, which is located in the upwind direction of the southwest monsoon potentially affecting the 122 atmospheric composition over CAS (NAMORS).

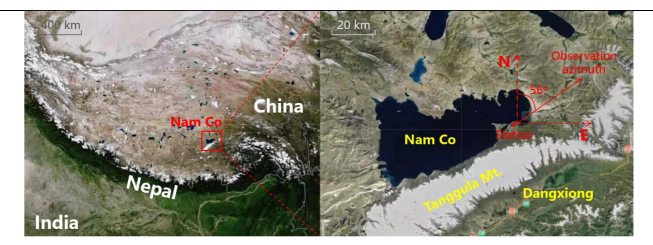




Figure 1. Geographical location of CAS (NAMORS) on the Tibet plateau.

### 125 **2.2 Measurements**

### 126 2.2.1 Instrument setup and spectral analysis

The MAX-DOAS instrument installed at CAS (NAMORS) was operated from 01 May to 09 July 2019. It consists of three major parts: telescope unit, spectrometer unit and control unit. The detailed description of this instrument can be found in Xing et al. (2021). In this study, the elevation angle sequence was set to 1, 2, 3, 4, 5, 6, 8, 10,15, 30, and 90° with an exposure time of 60 s to each individual spectrum. The azimuth angle was set to 56° pointing to Nagqu direction. Moreover, only spectra collected under solar zenith angle (SZA) less than 75° was used for spectral analysis to avoid the strong stratospheric absorption.

134 The differential slant column densities (DSCDs) of O<sub>4</sub>, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub> were retrieved using 135 QDOAS software (http://uvvis.aeronomie.be/software/QDOAS/) developed by Royal Belgian Institute for Space Aeronomy (BIRA-IASB). The zenith spectrum measured at every sequence were selected as 136 137 scan Frauenhofer reference spectrum. The retrieval configurations of O<sub>4</sub>, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub> 138 followed Xing et al. (2017), Lin et al. (2020), Xing et al. (2021), Wang et al. (2020) and Wang et al. 139 (2018), respectively. The detailed DOAS fit settings of above five species were listed in Table 1. 140 Corrected I<sub>0</sub> (Aliwell et al., 2002) was used in this study. Fig. 2 shows a typical DOAS retrieval example for above five species. DOAS fit results with root mean square (RMS) values larger than  $5 \times$ 141  $10^{-4}$ ,  $5 \times 10^{-4}$ ,  $5 \times 10^{-4}$ ,  $1 \times 10^{-3}$ , and  $6 \times 10^{-4}$  for O<sub>4</sub>, H<sub>2</sub>O, NO<sub>2</sub>, HONO, and O<sub>3</sub>, respectively, were 142 filtered out. In addition, we calculated color index (CI) to remove cloud effect (Wagner et al., 2016). 143 144 The data filter criteria according to CI followed by Ryan et al. (2018) and Xing et al. (2020). 145 Afterwards, the quantified DSCDs of O<sub>4</sub>, H<sub>2</sub>O, NO<sub>2</sub>, HONO, and O<sub>3</sub> remained 91.33%, 91.97%, 146 92.16%, 86.42% and 81.09%, respectively.

147 2.2.2 Vertical profile retrieval

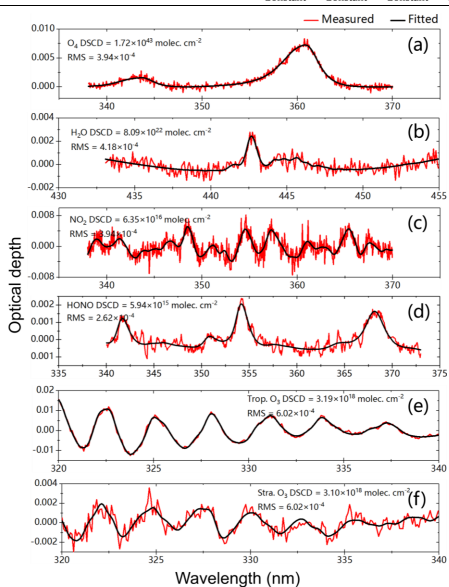
148 The vertical profiles of aerosol and trace gases (i.e. H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub>) were retrieved using 149 algorithm based on optimal estimation method (OEM). A linearized pseudo-spherical vector discrete 150 ordinate radiative transfer model VLIDORT was used as forward model and a Gauss-Newton (GN) scheme was used as the inversion strategy (Wedderburn et al., 1974). The detailed description of this 151 152 algorithm can be found in Liu et al. (2021), Xing et al. (2021) and Wang et al. (2018). The detailed 153 retrieval processes were depicted in Sect. S1 of the supplement. In this study, the initial a priori profile 154 shape of above five species was set to exponential decreasing shape, and the AOD and vertical column 155 densities (VCDs) simulated by weather research and forecasting model coupled chemistry (WRF-Chem) 156 were also used as initial input a priori information to constrain the retrieval process. For the  $O_3$  profile retrieval, the stratospheric O<sub>3</sub> profile was deducted using TROPOMI O<sub>3</sub> profile (Zhao et al., 2021). We 157 158 set 20 vertical layers from 0.0 to 4.0 km with a vertical resolution of 0.2 km. The correlation height was 159 set to 1.0 km. Moreover, the surface albedo, single scattering albedo and asymmetry parameter were set 160 to fixed constant of 0.08, 0.85 and 0.65, respectively (Irie et al., 2008). The retrieved vertical profiles

161 were removed under the condition of degree of freedom (DOF) and relative error less than 1.0 and

162 100%, respectively.

Parameter	Data source	Fitting intervals (nm)					
		O <sub>4</sub>	$H_2O$	$NO_2$	HONO	O <sub>3</sub>	
Wavelength range		338-370	433-455	338-370	340-373	320-340	
NO <sub>2</sub>	298K, I <sub>0</sub> -corrected, Vandaele et al. (1998)	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	
NO <sub>2</sub>	220K, Io-corrected, Vandaele et al. (1998)	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	×	
O <sub>3</sub>	223K, Io-corrected, Serdyuchenko et al. (2014)	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	
O <sub>3</sub>	243K, Io-corrected, Serdyuchenko et al. (2014)	$\checkmark$	×	$\checkmark$	$\checkmark$	×	
O <sub>3</sub>	293K, Io-corrected, Serdyuchenko et al. (2014)	×	×	×	×	$\checkmark$	
O <sub>4</sub>	293K, Thalman and Volkamer (2013)	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	
НСНО	298K, Meller and Moortgat (2000)	$\checkmark$	×	$\checkmark$	$\checkmark$	$\checkmark$	
Glyoxal	298K, Volkamer (2005)	×	$\checkmark$	×	×	×	
H <sub>2</sub> O	HITEMP (Rothman et al. 2010)	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	×	
BrO	223K, Fleischmann et al. (2004)	$\checkmark$	×	$\checkmark$	$\checkmark$	×	
HONO	296K, Stutz et al. (2000)	×	×	$\times$	$\checkmark$	×	
Ring	Calculated with QDOAS	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	
Polynomial degree		Order 3	Order 3	Order 3	Order 5	Order 3	
Intensity offset		Constant	Constant	Constant	Constant	No	

163 Table 1. Detailed DOAS retrieval settings for O<sub>4</sub>, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub>.



164

- 165 Figure 2. DOAS fit examples of  $O_4$ ,  $H_2O$ ,  $NO_2$ , HCHO, tropospheric  $O_3$  and stratospheric  $O_3$ . The red
- 166 line and black line represent the measured and fitted results, respectively.
- 167 2.2.3 Error analysis

168 The error sources can be divided into four different types: smoothing error, noise error, forward model

error, and model parameter error (Rodgers, 2004). However, in terms of this classification, some errors 169 170 are difficult to be calculated or estimated. For example, the forward model error, which is caused by an 171 imperfect representation of the physics of the system, is hard to be quantified due to the difficulty of acquiring an improved forward model. Given calculation convenience and contributing ratios of 172 173 different errors in total error budget, we mainly took into account following error sources, which were 174 smoothing and noise errors, algorithm error, cross section error, and uncertainty related to the aerosol retrieval (only for trace gas). In this study, we estimated the contribution of different error sources to 175 176 the AOD and VCDs of trace gases, and near-surface (0–200 m) trace gases' concentrations and aerosol 177 extinction coefficients (AECs), respectively. The detailed demonstrations and estimation methods are 178 displayed below.

- a. Smoothing errors arise from the limited vertical resolution of profile retrieval. Noise errors denote the noise in the spectra (i.e., the error of DOAS fits). Considering the error of the retrieved state vector equaling the sum of these two independent errors, we calculated the sum of smoothing and noise errors on near-surface concentrations and column densities, which were 13 and 5 % for aerosols, 13 and 36 % for H<sub>2</sub>O, 12 and 14 % for NO<sub>2</sub>, 18 and 21 % for HONO, and 12 and 32 % for O<sub>3</sub>, respectively.
- 185 b. Algorithm error is denoted by the differences between the measured and simulated DSCDs. This error contains forward model error from an imperfect approximation of forward function, parameter 186 187 error of forward model, and other errors, such as detector noise (Rodgers, 2004). Algorithm error is 188 a function of the viewing angle, and it is difficult to assign this error to each altitude. Thus, this 189 error on the near-surface values and column densities is estimated through calculating the average 190 relative differences between the measured and simulated DSCDs at the minimum and maximum 191 elevation angle (except 90°), respectively (Wagner et al., 2004). In this study, we estimated these 192 errors on the near-surface values and the column densities at 4 and 8 % for aerosols, 3 and 11 % for 193 NO<sub>2</sub>, and 20 and 20 % for HONO referring to Wang et al. (2017, 2020), 1 and 8 % for H<sub>2</sub>O 194 referring to Lin et al. (2020), and 6 and 10 % for O<sub>3</sub> referring to Ji et al. (2023), respectively.
- c. Cross section error arises from the uncertainty in the cross section. According to Thalman and Volkamer, (2013), Lin et al. (2020), Vandaele et al. (1998), Stutz et al. (2000), and Serdyuchenko et al. (2014), we adopted 4, 3, 3, 5, and 2 % for O<sub>4</sub> (aerosols), H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub>, respectively.
- d. The profile retrieval error for trace gases is sourced from the uncertainty of aerosol extinction profile retrieval and propagated to trace gas profile. This error could be roughly estimated based on a linear propagation of the total error budgets of the aerosol retrievals. The errors of the learned four trace gases were roughly estimated at 14 % for VCDs and 10 % for near-surface concentrations, respectively.
- The total uncertainty was the sum of all above errors in the Gaussian error propagation, and the error results were listed in Table 2. We found that the smoothing and noise errors played a dominant role in the total uncertainties of aerosol and trace gases. Moreover, improving the accuracy and temperature gradient of the absorption cross section is another important means to reduce the uncertainty of the vertical profiles in the future, especially for  $O_3$ .
- Table 2. Error budget estimation (in %) of the retrieved near-surface (0–200 m) concentrations of trace gases and AECs, and AOD and VCDs.

			Error sources			
		Smoothing and	moothing and Algorithm error Cross section Related to the			
		noise errors	-	error	aerosol retrieval	
Near-surface	aerosol	13	4	4	/	14
	$H_2O$	13	1	3	14	19
	$NO_2$	12	3	3	14	18
	HONO	18	20	5	14	29

	O <sub>3</sub>	12	6	2	14	19
VCD or AOD	AOD	5	8	4	/	10
	$H_2O$	36	8	3	10	38
	$NO_2$	14	11	3	10	20
	HONO	21	20	5	10	31
	$O_3$	32	10	2	10	35

### 212 2.3 TUV model

213 The calculation of photolysis rates of HONO and O<sub>3</sub> used TUV radiation model 214 (https://www2.acom.ucar.edu/modeling/tropospheric-ultraviolet-and-visible-tuv-radiation-model)

215 based on a full FORTRAN code. In order to ensure the accuracy of model running, we only selected data in sunny and cloudless days. Moreover, we developed a cloud classification method based on the 216 217 diurnal variations of Color Index (CI= $I_{330}/I_{360}$ ) in Figure S2. The initial input parameters were as 218 follows: the AOD at 361 nm was derived from aerosol extinction profiles measured by MAX-DOAS; the daily total ozone column density was measured by TROPOMI with a value range of 260-280 DU; 219 the single scattering albedo (SSA) was calculated based on the regression analysis of multi-wavelength 220 (361 and 477 nm) O<sub>4</sub> absorptions measured by MAX-DOAS (Xing et al., 2019); fixed Ångström 221 222 exponents of 0.508, 0.581 and 0.713 were used in May, June and July, respectively, referring to Xia et 223 al. (2011).

### 224 **2.4 Backward trajectory, PSCF and CWT analysis**

The 48-h backward trajectories at five heights of 200, 600, 1000, 1400 and 1800 m were calculated using the Hybrid Single-particle Lagrangian Integrated Trajectory (HYSPLIT) model based on the Global Data Assimilation System (GDAS) to identify the major transport pathways of O<sub>3</sub> (Draxler and Hess, 1998). Moreover, the calculated backward trajectories were clustered into three groups using Ward's variance method and Angle Distance algorithm (Ward 1963; Wang et al., 2006).

230 In order to determine the potential source locations of  $O_3$  over CAS (NAMORS), the Potential Source 231 Contribution Function (PSCF) model and Concentration Weighted Trajectory (CWT) model were used 232 (Hong et al., 2019; Ou et al., 2021). The PSCF was calculated through the number of air trajectory 233 endpoints being divided by the number of air trajectory endpoints. Moreover, a weighting function was 234 introduced to reduce the increased uncertainties of PSCF with the increase of the distance between the 235 grid and sampling point. In this study, the set of this weighting function referred to Yin et al. (2017). CWT can be used to calculate the weight concentration through averaging the concentrations 236 associated with trajectories crossing the grid cell. Above weighting function was also introduced to 237 calculate the WCWT (Hsu, et al., 2003). The detailed description of these two models can be found in 238 239 Wang et al., 2006.

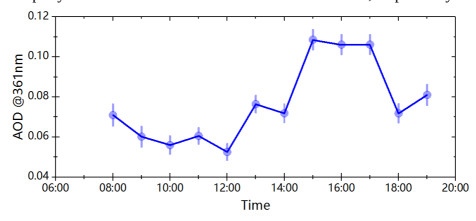
### 240 2.5 Ancillary data

241 The surface NO<sub>2</sub>, HONO and O<sub>3</sub> concentrations used to validate the corresponding MAX-DOAS 242 measurements were monitored by broadband cavity enhanced spectrometer (BBCES) (Fang et al., 243 2017), long path absorption photometer (LOPAP) (Kleffmann et al., 2008) and Thermo Electron 49i (Shi et al., 2009), respectively. The PBL height was simulated using WRF with spatiotemporal 244 resolutions of  $20 \times 20$  km<sup>2</sup> and 1.0 hour (detailed configurations in Sect. S3 of the supplement). 245 Moreover, the large-scaled spatial distributions of AOD, O<sub>3</sub> and NO<sub>2</sub> over CAS (NAMORS) were 246 247 monitored by Himawari-8 (Bessho et al., 2016), OMI (Veefkind et al., 2004) and TROPOMI (Griffin et al., 2018; Su et al., 2020), respectively. 248

### 249 **3 Results**

### **3.1 Overview of the measurements**

Figure 3 showed the averaged diurnal variation of AOD from 1<sup>st</sup> May to 9<sup>th</sup> July 2019, with an average value of 0.076 km<sup>-1</sup> during 08:00-19:00. The AOD was 0.071 km<sup>-1</sup> at 08:00, and then gradually decreased to a minimum value of 0.052 km<sup>-1</sup> at 12:00. Subsequently, the AOD increased significantly, reaching maximum values during 15:00-17:00 (average of 0.107km<sup>-1</sup>), which was about 1.408 times the diurnal average value. Considering the diurnal variation of wind speed (Figure S3), such an enhancement of AOD may be related to the long-range transport of aerosol from southern Asia (Yang et al., 2020; Bi et al., 2023). Moreover, 15:00-17:00 was the active time of tourists and local residents (i.e. cooking), and these kinds of anthropogenic sources contributed to the atmospheric AOD of NAMORS through short-distance transport (Yin et al., 2017; Zhang et al., 2017). After 17:00, the AODs decreased rapidly to 0.071 km<sup>-1</sup> at 18:00 and 0.081 km<sup>-1</sup> at 19:00, respectively.



261

Figure 3. Averaged diurnal variation of AOD at CAS (NAMORS). The error bars represent the mean retrieved errors of AOD.

As shown in Figure S4, the diurnal variation of PBL in Nam Co from May to July 2019 was lower in the early morning and late afternoon, but higher between 11:00 and 17:00, a relatively long period, with the maximum PBL larger than 2.0 km. Zhang et al. (2017) and Yang et al., (2017) also reported that the PBL in Nam Co was usually larger than 1.0 km during daytime in spring and summer. In order to investigate the height-dependent variations of aerosol, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub> within the PBL during the measurements, five height layers under the PBL (0.0-0.2 km, 0.4-0.6 km, 0.8-1.0 km, 1.2-1.4 km and 1.6-1.8 km) were thus selected.

Figure 4 showed the time series of the daily averaged aerosol, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub> at above five 271 272 layers from 1<sup>st</sup> May to 9<sup>th</sup> July 2019. Aerosol mainly distributed at 0.0-0.2 km with an average 273 extinction coefficient of 0.138 km<sup>-1</sup>, and the ratios of aerosol extinction at 0.4-0.6 km, 0.8-1.0 km, 274 1.2-1.4 km and 1.6-1.8 km to those at 0.0-0.2 km were 39.34%, 18.77%, 7.29% and 2.62%, 275 respectively. That indicated that the aerosol was usually local-emitted at the surface, and the occasionally appearance of strong aerosol extinction at 0.4-0.6 km, such as 13<sup>th</sup> and 30<sup>th</sup> June, was 276 associated with long-range transport from south Asia (Figure S5, Wan et al., 2015; Li et al., 2016). The 277 average concentration of H<sub>2</sub>O at 0.0-0.2 km was  $2.35 \times 10^{17}$  molec cm<sup>-3</sup>, and the ratios of H<sub>2</sub>O at 278 0.4-0.6 km, 0.8-1.0 km, 1.2-1.4 km and 1.6-1.8 km to those at 0.0-0.2 km were 83.40%, 68.08%, 279 280 50.64% and 35.74%, respectively, which should attribute to the transport of H<sub>2</sub>O from southern Asia 281 driven by the Indian ocean monsoon and the elevated evaporation from Nam Co lake to lead to its not obvious vertical gradient (Figure S6, Lei et al., 2014; Zhu et al., 2019). The average concentration of 282 283 NO<sub>2</sub> at 0.0-0.2 km was 0.193 ppb, and its high concentration mainly distributed at 0.4-0.6 km after 15<sup>th</sup> May. The ratios of NO<sub>2</sub> at 0.4-0.6 km, 0.8-1.0 km, 1.2-1.4 km and 1.6-1.8 km to those at the bottom 284 layer were 104.03%, 59.05%, 24.62% and 12.84%, respectively. The elevation of the distribution 285 height of high concentration  $NO_2$  should be attributed to the transport process from the  $NO_x$  produced 286 287 by ice and snow on the top of Mt. Tanggula under strong ultraviolet radiation (Boxe et al., 2005; Fisher 2005; Lin et al., 2021). As depicted in Figure S7, the WPSCF passing through Mt. Tanggula showed 288 high values at 300-400 m layer, especially at 400 m (> 0.3). It also indirectly indicated that the 289 290 important contribution to NO<sub>x</sub> from ice and snow on the top of mountains under strong ultraviolet 291 radiation on the TP. HONO mainly distributed at 0.0-0.2 km with an average value of 0.087 ppb, and 292 the ratios of HONO at 0.4-0.6 km, 0.8-1.0 km, 1.2-1.4 km and 1.6-1.8 km to those at 0.0-0.2 km were 293 58.49%, 44.64%, 31.30% and 21.67%, respectively. That indicated that the primary and secondary 294 sources of HONO were mainly at the surface (Section 4.2). The vertical gradient of daily averaged O<sub>3</sub> 295 concentration was also not obvious, which was associated with its vertical mixing and photochemical

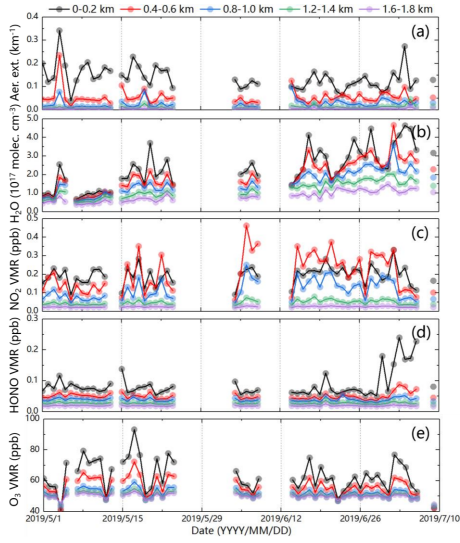
296 production (Yin et al., 2017). As shown in Figure S8, the corresponding TROPOMI O<sub>3</sub> profiles in Nam

297 Co and O<sub>3</sub> profiles measured by lidar and ozonesonde around Nam Co reported in several previous

298 studies also exhibited an exponential shape (Fang et al., 2019; Zhang et al., 2020; Yu et al., 2022). The

299  $O_3$  average concentration at 0.0-0.2 km was 63.030 ppb, and the ratios of  $O_3$  at 0.4-0.6 km, 0.8-1.0 km,

300 1.2-1.4 km and 1.6-1.8 km to those at surface were 89.25%, 82.44%, 80.16% and 79.13%, respectively.



301

Figure 4. Time series of daily averaged (a) aerosol extinction, (b) H<sub>2</sub>O, (c) NO<sub>2</sub>, (d) HONO, and (e) O<sub>3</sub> monitored by MAX-DOAS at 0-0.2, 0.4-0.6, 0.8-1.0, 1.2-1.4 and 1.6-1.8 km five height layers from 01 May to 09 July 2019.

# 305 **3.2 Vertical distributions of aerosol, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub>**

306 The first row in Figure 5 provided the averaged vertical profiles of aerosol, H<sub>2</sub>O, NO<sub>2</sub>, HONO and O<sub>3</sub> 307 from May to July 2019. We found that the vertical profiles of aerosol, H<sub>2</sub>O, HONO and O<sub>3</sub> all 308 exhibited an exponential shape with maximum values near the surface, while NO<sub>2</sub> exhibited a Gaussian shape with the maximum value of 0.321 ppb occurring at 0.3-0.4 km layer. In addition to the effect of 309 NO<sub>x</sub> transport, Xu et al. (2018) also revealed that the long-range high-altitude transport process from 310 311 the northern south Asian subcontinent can significantly enhance the Nam Co's peroxyacetyl nitrate 312 (PAN) level which is a reservoir of NO<sub>x</sub>. As shown in the second row of Figure 5, the monthly averaged aerosol vertical profiles from May to July 2019 all exhibited an exponential shape, and varied 313 314 in the order of May  $(0.17 \text{ km}^{-1}) > \text{July} (0.14 \text{ km}^{-1}) > \text{June} (0.11 \text{ km}^{-1})$ . Xu et al. (2018) and Neupane et al. (2019) also reported a similar monthly variations of black carbon (BC) from May to July over the 315 316 TP, and revealed that it was mainly associated with the anthropogenic emissions (i.e. biomass burning) 317 and its transport from south Asia. The monthly averaged vertical profile of  $H_2O$  in May and July 318 exhibited an exponential shape, while its maximum concentration layer slightly elevated to 0.1-0.2 km

319 in June which was related to the strongest monsoon transport (Figure S9). It varied in the order of July  $(3.68 \times 10^{17} \text{ molec cm}^{-3})$  > June  $(2.71 \times 10^{17} \text{ molec cm}^{-3})$  > May  $(2.26 \times 10^{17} \text{ molec cm}^{-3})$ , and its 320 maximum concentration occurring in July was strongly associated with the enhanced evaporation from 321 the Nam Co lake (Xu et al., 2011). The monthly averaged vertical profiles of NO<sub>2</sub> all exhibited a 322 Gaussian shape from May to July, and its maximum values mainly distributed at 0.2-0.4 km layer 323 324 varying in the order of June (0.39 ppb) > May (0.31 ppb) > July (0.28 ppb). It indicated that the 325 regional transport from the NO<sub>x</sub> produced from ice and snow under strong shortwave radiation (Figure 326 S7),  $NO_2$  emitted from vehicles due to the increased tourism, anthropogenic emissions from local 327 residents (i.e. biomass burning and religious activities) played an important role in the vertical distribution characteristic of NO<sub>2</sub> (Boxe et al., 2005; Chen et al., 2019). The monthly averaged vertical 328 329 profiles of HONO from May to July all exhibited an exponential shape, with maximum values near the 330 surface varying in the order of July (0.13 ppb) > May (0.07 ppb) > June (0.06 ppb). The local direct 331 emissions from biomass burning, vehicles and soil should be main sources of the surface HONO (Xing et al., 2021). Moreover, the heterogeneous reaction of NO<sub>2</sub> on wet surfaces should be another important 332 333 source of HONO at different height layers (Section 4.2). For example, the aerosol extinction coefficient, 334 and the concentrations of H<sub>2</sub>O and NO<sub>2</sub> were all relatively large at the bottom layer in July, correspondingly, we observed the highest concentration of HONO near the surface in this month. The 335 336 monthly averaged O<sub>3</sub> vertical profiles all showed an exponential shape from May to July, and its 337 surface concentration varied in the order of May (66.71 ppb) > July (61.45 ppb) > June (59.55 ppb). This kind of monthly variation trend of O<sub>3</sub> was also reported by several previous studies (Yin et al., 338 339 2017; Xu et al., 2018). The O<sub>3</sub> in Nam Co was mainly sourced from stratospheric intrusion, 340 photochemical reactions, long-range transport and local vertical mixing (Yin et al., 2017; Chen et al., 341 2019).

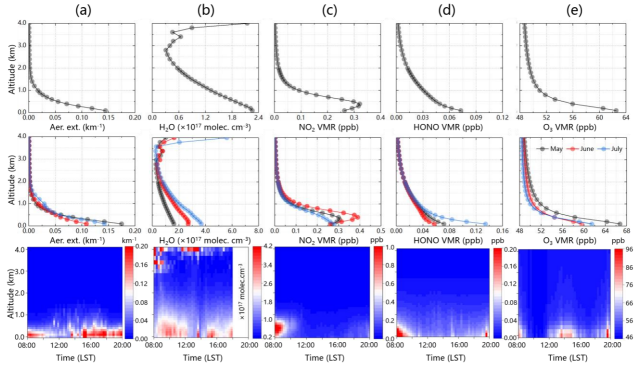




Figure 5. Vertical profiles of (a) aerosol extinction, (b) H<sub>2</sub>O, (c) NO<sub>2</sub>, (d) HONO, and (e) O<sub>3</sub>. The top row shows the averaged vertical profiles from 01 May to 09 July 2019. The middle row shows the monthly averaged vertical profiles. The bottom row shows the averaged diurnal vertical profiles from 01 May to 09 July 2019.

The third row in Figure 5 illustrated the averaged diurnal variations in vertical profiles of aerosol,  $H_2O$ , NO<sub>2</sub>, HONO and O<sub>3</sub> from May to July 2019. Aerosol mainly distributed under 1.0 km, especially 0.6 km, and its mixing height was gradually increased with the rise of the PBL height after 12:00. Moreover, the diurnal variation of aerosol showed a bi-peak pattern, which was in line with the investigation reported by Pokharel et al. (2019). The first peak occurred between 08:00-10:00, and

another appeared after 15:00. The first peak should be attributed to the local emission of aerosol and 352 the diurnal cycle of PBL (Zhang et al., 2017; Pokharel et al., 2019). The second peak was driven by 353 354 regional transport and the interaction between local sandy silt loam surface and local meteorology. The high wind speed (> 4.5 m/s) at surface appeared after 15:00, which coincided with the appearance of 355 356 the second aerosol peak (Figure S3). Moreover, the high extinction during the second peak was 357 extended to 1.0 km associated with the wind speed larger than 8 m/s (Figure S10), which created a 358 favorable condition for high-altitude aerosol transport. H<sub>2</sub>O mainly distributed under 1.0 km and above 359 3.0 km, and its diurnal variation exhibited a multi-peak pattern. The first peak appeared between 360 08:00-12:00, which was mainly affected by the monsoon drived long-range transport of H<sub>2</sub>O (Cong et al., 2009; Xu et al., 2020). The second and third peaks occurred at 15:00-16:00 and after 17:00, 361 respectively. In addition to long-range transport, the enhanced evaporation from the Nam Co lake also 362 363 significantly contributed to the appearance of these two peaks of H<sub>2</sub>O (Xu et al., 2011). NO<sub>2</sub> mainly distributed at 0.2-0.4 km, and peaked before 10:00 and after 18:00 which were dominated by the effects 364 of local emissions and regional transport from the NOx formed through ice and snow on the top of Mt. 365 366 Tanggula under strong ultraviolet radiation (Figure S7) (Boxe et al., 2005; Fisher 2005; Chen et al., 367 2019; Lin et al., 2021). Moreover, its diurnal mixing height was obviously correlated to the diurnal 368 evolution of PBL height. HONO mainly distributed under 1.0 km, especially 0.4 km. Its diurnal 369 variation showed a multi-peak pattern with three obvious peaks before 10:00, 15:00-16:00, and after 370 19:00. In addition to local emissions (i.e. vehicle emission, biomass burning and soil emission), the heterogeneous reaction of NO<sub>2</sub> on wet surfaces should be also an important HONO source (Xing et al., 371 372 2021). We found that there were larger aerosol extinction (>  $0.12 \text{ km}^{-1}$ ) and higher concentrations of NO<sub>2</sub> (> 0.20 ppb) and H<sub>2</sub>O (>  $2.27 \times 10^{17}$  molec cm<sup>-3</sup>) around three HONO peaks. O<sub>3</sub> mainly 373 distributed under 0.4 km, and its diurnal variation exhibited a multi-peak pattern with three peaks 374 375 appearing before 09:00, 13:00-15:00 and after 19:00. The appearance of O<sub>3</sub> peaks was mainly 376 associated with the influence of the complex topography of the TP, long-range transport, local vertical 377 mixing and stratospheric intrusion (Yin et al., 2017; Chen et al., 2019; Qian et al., 2022). The active 378 photochemical reaction should be another important source of O<sub>3</sub>, especially for its second peak at 379 13:00-15:00.

### 380 **3.3 Validation with independent data**

381 In order to validate the MAX-DOAS dataset, we extracted the concentrations of NO<sub>2</sub>, HONO and O<sub>3</sub> at 382 the bottom layer (0.0-0.1 km) from their corresponding vertical profiles to compare with in situ measurements. As shown in Figure 6(a-c), we found good agreements between MAX-DOAS and in 383 situ observations with Pearson correlation coefficients (R) of 0.91, 0.62 and 0.82 (regression slope of 384 0.89, 1.05 and 0.82) for NO<sub>2</sub>, HONO and O<sub>3</sub>, respectively. That indicated the good reliability of trace 385 386 gases from MAX-DOAS retrievals. Moreover, we also compared the MAX-DOAS PBL and WRF PBL, 387 and a similar variation trend was found. However, WRF PBL showed a significantly difference in 388 height values with MAX-DOAS PBL before 12:00. That should be due to the simulation uncertainties 389 for WRF model at Tibetan plateau with complex topography and meteorology (Yang et al., 2016; Xu et 390 al., 2019).

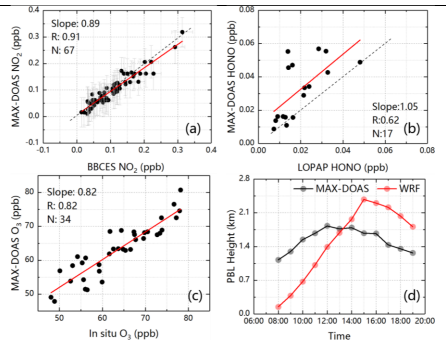


Figure 6. Validations of (a) MAX-DOAS NO<sub>2</sub> vs in situ NO<sub>2</sub> (error bars represent the retrieved errors
of NO<sub>2</sub> from MAX-DOAS and BBCES), (b) MAX-DOAS HONO vs LOPAP HONO, (c)
MAX-DOAS O<sub>3</sub> vs in situ O<sub>3</sub>, and (d) MAX-DOAS PBL vs WRF PBL.

# 395 **4 Discussion**

# 396 4.1 OH production

HONO and  $O_3$  are two important precursors of OH redical to enhance the AOC (Kleffmann et al., 2005; Ryan et al., 2018; Xing et al., 2021). In order to evaluate the AOC on the TP, we tried to analyze the OH production from HONO and  $O_3$  at different height layers through vertical observations and TUV calculations. The OH production rates from HONO and  $O_3$  were calculated using the following two equations:

402  $P(OH)_{HONO} = J(HONO) \times [HONO]$ 

403  $P(OH)_{O_2} = 2 \times f \times J(O(^1D)) \times [O_3]$ 

404 Where J(HONO) and  $J(O(^{1}D))$  were the photolysis rates of HONO and O(^{1}D) calculated using TUV 405 model. O(^{1}D) was the product from O<sub>3</sub> photolysis by UV radiation. *f* was the fraction of the process 406 O(^{1}D) + H<sub>2</sub>O  $\rightarrow$  2OH.

407 Figure 7(a-b) showed the averaged diurnal vertical distributions of the photolysis rates J(HONO) and 408  $J(O(^{1}D))$  from May to July 2019. We found that the maximum J(HONO) and  $J(O(^{1}D))$  were all appeared at the bottom layer between 12:30 and 15:30 with values of  $2.0 \times 10^{-3}$  and  $6.75 \times 10^{-5}$  s<sup>-1</sup>. 409 respectively. The maximum values were usually larger than that at low-altitude areas due to the 410 stronger solar UV radiation on the TP (Su et al., 2008; Xing et al., 2021; Yang et al., 2021; Liu et al., 411 412 2022), but being consistent with the values on the TP reported by Lin et al. (2008). Moreover, it should 413 be noted that the values of J(HONO) and  $J(O(^{1}D))$  all decreased with the increase of altitude, which was significantly different with previous studies in low altitudes (Ryan et al., 2018; Xing et al., 2021; 414 415 Xu et al., 2021).

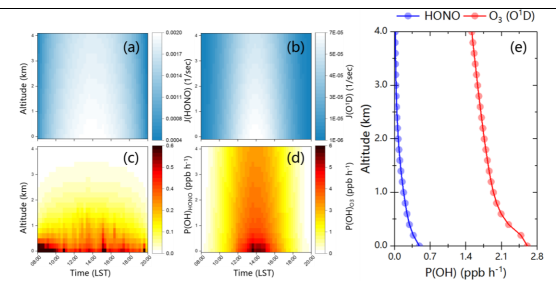


Figure 7. Averaged diurnal vertical profiles of the (a) photolysis rate J(HONO), (b) photolysis rate  $J(O(^{1}D))$ , (c) OH radical production rates from HONO photolysis, (d) OH radical production rates from O<sub>3</sub> photolysis. (e) shows the averaged vertical profiles of OH radical production rates from HONO and

420 O<sub>3</sub> photolysis from 01 May to 09 July 2019.

421 Figure 7(c-d) showed the averaged diurnal vertical profiles of OH production rates from HONO and O<sub>3</sub> 422 photolysis from May to July 2019. P(OH)<sub>HONO</sub> exhibited a multi-peak pattern which mainly appeared 423 before 10:00, 15:00-16:00, and after 19:00 at 0-0.4 km with a maximum value of 0.81 ppb/h. While 424 P(OH)O<sub>3</sub> showed a unimodal pattern occurring at 13:00-15:00 under 0.4 km with a maximum value of 425 6.20 ppb/h. The averaged vertical profiles of P(OH)<sub>HONO</sub> and P(OH)<sub>O3</sub> during the observation were depicted in Figure 7(e). We found that the maximum values of P(OH)<sub>HONO</sub> (0.49 ppb/h) and P(OH)<sub>O3</sub> 426 427 (2.61 ppb/h) all appeared at the bottom layer, and decreased with height. That indicated O<sub>3</sub> was the main contributor of OH production (> 80%) on the TP, which was about 5-6 times to HONO. 428 429 Moreover, the OH production rates from HONO and  $O_3$  in other cities of China were depicted in Table 3. The contribution percentage of O<sub>3</sub> to P(OH) in Nam Co was significantly higher than that in other 430 431 cities, which was due to the relatively high concentrations of O<sub>3</sub> and H<sub>2</sub>O, and the strong radiation in 432 Nam Co. In addition, P(OH)<sub>HONO</sub> in Nam Co was close to that in relatively dry areas (i.e. Beijing and 433 Xianghe), but slightly lower than that in areas with relatively high humidity which can enhance the 434 heterogeneous production of HONO (Ryan et al., 2018; Liu et al., 2019; Xing et al., 2021).

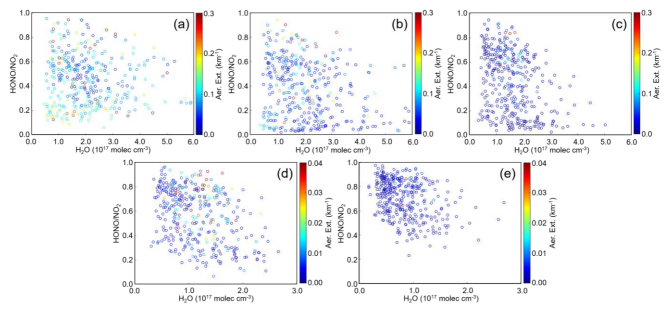
Location	Date	P(OH) <sub>HONO</sub> (ppb/h)	$P(OH)_{O3}$ (ppb/h)	References	
Xianghe (China)	Jul. 2008-Apr. 2009	4.1 2008 Apr 2000 ~0.80 in Spring		Hendrick et al. (2014)	
Alanghe (Chilla)	Jul. 2008-Apt. 2009	~0.70 in Summer	~0.45 in Summer	rienunck et al. (2014)	
Beijing (China)	Mar. 2010-Dec. 2012	~1.25 in Spring,	~0.10 in Spring,	Hendrick et al. (2014)	
Beljing (China)	Wiai. 2010-Dec. 2012	~0.70 in Summer	~0.55 in Summer	Hendrick et al. (2014)	
East China Sea (China)	Jun. 2017	~1.75	~1.20	Cui et al. (2019)	
Chengdu (China)	AugSep. 2019	~3.25	-	Yang et al. (2021)	
Qingdao (China)	JulAug. 2019	~1.30	~1.00	Yang et al. (2021)	
Nam Co (China)	May-Jul. 2019	0.81	6.20	This study	

435	Table 3. The maximum	n OH	production rates	contributed from	HONO	and $O_3$ at c	lifferent locations.	

### 436 **4.2 Possible daytime HONO sources**

437 Atmospheric HONO mainly sourced from direct emission, homogeneous reaction and heterogeneous 438 reaction (Fu et al., 2019; Ren et al., 2020; Chai et al., 2021; Crilley et al., 2021; Li et al., 2021). There 439 were less anthropogenic emissions for HONO around NAMORS, however, the open burning of crop 440 residues and soil emissions should be important HONO sources considering the pasture environment 441 and large amounts of animal manure (Cui et al., 2021a; 2021b). Moreover, the background of low-level 442 NO on the TP leaded to the homogeneous reaction not to be the main source of HONO at NAMORS 443 (Lin et al., 2019; Xing et al., 2021; Li et al., 2022). Heterogeneous reaction of NO<sub>2</sub> on wet surfaces became an important potential source of HONO around NAMORS, which affected by the humidity, 444

temperature, solar radiation, aerosol concentration and corresponding specific surface area. In order to 445 446 remove the effect of diurnal PBL evolution, we used HONO/NO<sub>2</sub> to indicate the extent of the heterogeneous reaction process. As shown in Figure 8, scatter plots between HONO/NO<sub>2</sub> and H<sub>2</sub>O 447 448 were illustrated. We found that the maximum value of HONO/NO<sub>2</sub> appeared around water vapor being around  $1.0 \times 10^{17}$  molec cm<sup>-3</sup> under 1.0 km, and being around  $0.5 \cdot 1.0 \times 10^{17}$  molec cm<sup>-3</sup> at 1.0-2.0 km 449 height layer. This phenomenon of HONO/NO<sub>2</sub> firstly increasing and then decreasing with the 450 increasing of H<sub>2</sub>O (or relative humidity) was usually found in low-altitude areas in previous studies 451 452 (Wang et al., 2013; Liu et al., 2019; Xing et al., 2021; Xu et al., 2021). When the  $H_2O$  was greater than above mentioned critical values at different heights, HONO/NO2 gradually decreased, which was 453 related to the efficient uptake of HONO and the decrease of NO<sub>2</sub> reactivity with the increase of H<sub>2</sub>O 454 (Liu et al., 2019; Xu et al., 2021). That indicated H<sub>2</sub>O has significant enhancement for the conversion 455 rate of NO<sub>2</sub> to HONO. Moreover, we found that the high value areas of HONO/NO<sub>2</sub> at above five 456 457 height layers were all accompanied by high aerosol extinction (> 0.15 km<sup>-1</sup> under 1.0 km, and > 0.02 458 km<sup>-1</sup> at 1.0-2.0 km). It indicated that aerosol surface has contribution to the heterogeneous reaction 459 process of NO<sub>2</sub>. The scatter plots between HONO and NO<sub>2</sub> at above five layers (Figure S11) also 460 confirmed the possibility of the NO<sub>2</sub> heterogeneous reaction to generate HONO on the TP, and the contribution of atmospheric H<sub>2</sub>O and aerosol extinction to this process. 461



#### 462

Figure 8. Scatter plots between HONO/NO<sub>2</sub> and H<sub>2</sub>O colored by aerosol extinction at (a) 0.0-0.2 km, (b) 0.4-0.6 km, (c) 0.8-1.0 km, (d) 1.2-1.4 km, (and e) 1.6-1.8 km from  $1^{st}$  May to  $9^{th}$  July 2019.

In Figure 9, the vertical profile of HONO/NO<sub>2</sub> from May to July 2019 was depicted. We found that HONO/NO<sub>2</sub> firstly decreased and then increased with the increasing of height, which was opposite to previous studies in low-altitude areas (Meng et al., 2020; Zhang et al., 2020; Xing et al., 2021; Xu et al., 2021). The minimum average HONO/NO<sub>2</sub> occurred at 0.3-0.4 km height layer with a value of 0.37. The relatively high values of HONO/NO<sub>2</sub> at the bottom layer should be related to the non-deducted HONO direct emissions.

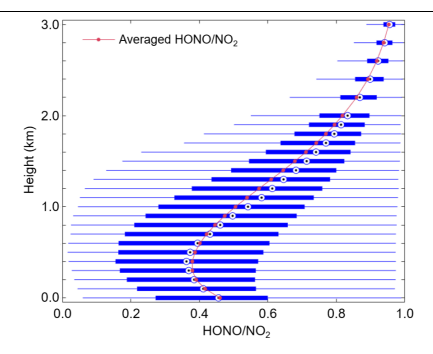


Figure 9. Statistics for the vertical profile of HONO/NO<sub>2</sub> from  $1^{st}$  May to  $9^{th}$  July 2019. The left and right of the blue box represent the  $25^{th}$  and  $75^{th}$  percentiles, respectively; the dot within the box represents the mean.

### 475 **4.3 Possible daytime O3 sources**

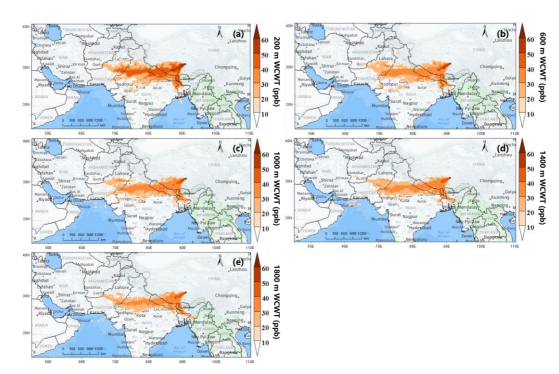
476 In addition to local photochemistry process, long-range transport was the main source of  $O_3$  on the TP (Yin et al., 2017; Xu et al., 2018). To further understand the transport pathway and potential source of 477 O<sub>3</sub>, cluster analysis, WPSCF and WCWT models were used to assess the regional representativity of 478 479 O<sub>3</sub> at five typical heights (200 m, 600 m, 1000 m, 1400 m and 1800 m). As shown in Figure S12 and Table 4, the backward trajectories arriving at NAMORS during the observation were classified into 480 three clusters at 200 m, 600 m, 1400 m, 1800 m, and four clusters at 1000 m. We found that cluster 3 481 482 was associated with the highest  $O_3$  concentration at 200 m (65.48±17.41 ppb) and 1800 m (49.69± 2.21 ppb), and cluster 1 were related to the highest  $O_3$  concentration at 600 m (54.67  $\pm$  6.94 ppb), 1000 483 484 m (51.61 $\pm$ 3.84 ppb) and 1400 m (50.51 $\pm$ 2.89 ppb). These two clusters were all originating from 485 northwestern of south Asian subcontinent passing through Himalayas, which was also reported by Yin et al. (2017) during springtime from 2011 to 2015. In Figure S13 and 10, WPSCF and WCWT analysis 486 487 told us that the high  $O_3$  concentration at above heights potentially sourced from northern India, central 488 Pakistan, Nepal, western Bhutan and northern Bangladesh through long-range transport. It should be 489 noted that the potential contribution to O<sub>3</sub> at NAMORS at 200 m from above potential source areas 490 were all over 40 ppb. These contributions from the mentioned potential source areas at other four 491 heights were also over 20-30 ppb. The massive fire emissions during springtime were an important 492 source of  $O_3$  in south Asia (Jena et al., 2015), and the obvious burning during the observation was 493 observed in Figure S14. Moreover, the abundant precursors and high photochemical activity were 494 another significant sources of O<sub>3</sub> in south Asia (Kumar et al., 2012; Sharma et al., 2017).

In addition, Figure 10 showed that the contribution of O<sub>3</sub> transported from Himalayas can even up to 50 ppb, especially under 600 m. Several previous studies have revealed that the stratospheric O<sub>3</sub> intrusion events were frequent in the Himalayas during spring and summer (Cristofanelli et al., 2010; Chen et al., 2011; Škerlak et al., 2014; Putero et al., 2016). Therefore, the O<sub>3</sub> from stratospheric intrusions in the Himalayas can affect the O<sub>3</sub> at NAMORS through long-range transport.

Table 4. Trajectory ratios and averaged O<sub>3</sub> concentration for all trajectory clusters arriving in Nam Co at 200 m, 600 m, 1000 m, 1400 m and 1800 m from May to July 2019.

Classie	т : <i>с</i> :	O <sub>3</sub> concentration (ppb)
Cluster	Traj_ratio	Mean±SD

	1	55.86%	$61.50 \pm 18.15$
200	2	11.85%	54.57±14.67
200 m	3	32.28%	65.48±17.41
	All	100.00%	61.14±17.74
	1	62.55%	54.67±6.94
600	2	14.32%	50.43±6.64
600 m	3	23.13%	53.27±7.63
	All	100.00%	53.39±7.26
	1	49.16%	$51.61 \pm 3.84$
	2	8.81%	49.60±3.99
1000 m	3	22.73%	$50.72 \pm 4.21$
	4	19.30%	51.39±4.49
	All	100.00%	$50.98 \pm 4.30$
	1	80.14%	$50.51 \pm 2.89$
1400 m	2	4.95%	49.12±2.73
1400 III	3	14.92%	$49.44 \pm 3.85$
	All	100.00%	$50.07 \pm 3.15$
	1	83.75%	49.68±2.55
1800 m	2	0.00%	$49.07 \pm 2.23$
1000 111	3	16.25%	$49.69 \pm 2.21$
	All	100.00%	$49.59 \pm 2.49$



503

Figure 10. Spatial distributions of WCWT values for  $O_3$  at (a) 200 m, (b) 600 m, (c) 1000 m, (d) 1400 m, and (e) 1800 m height layers from  $01^{st}$  May to  $09^{th}$  July 2019 over CAS (NAMORS).

# 506 **5 Summary and conclusions**

507 MAX-DOAS measurements were performed to clarify the vertical distributions of several atmospheric 508 components (aerosol,  $H_2O$ ,  $NO_2$ , HONO and  $O_3$ ), and to explore the AOC in vertical space in Nam Co 509 from May to July 2019. The MAX-DOAS NO<sub>2</sub>, HONO and O<sub>3</sub> agreed well with in situ measurements, 510 with correlation coefficients of 0.91, 0.62 and 0.82, respectively. We found that the averaged vertical

- 511 profiles of aerosol, H<sub>2</sub>O, HONO and O<sub>3</sub> all exhibited an exponential shape, while NO<sub>2</sub> showed a 512 Gaussian shape with a maximum value of 0.32 ppb appearing at 300-400 m. The maximum 513 concentrations of monthly averaged aerosol (0.17 km<sup>-1</sup>) and O<sub>3</sub> (66.71 ppb) appeared on May, H<sub>2</sub>O 514  $(3.68 \times 10^{17} \text{ molec cm}^{-3})$  and HONO (0.13 ppb) appeared on July, and NO<sub>2</sub> (0.39 ppb) occurred on 515 June. For the diurnal variation, above five species all mainly distributed under 1.0 km, and mostly 516 exhibited a multi-peak pattern considering the effect of regional transport and local chemical reaction.
- 517 O<sub>3</sub> and HONO were the main source of OH on the TP. The diurnal averaged OH production rate from 518 HONO during the observation exhibited a multi-peak pattern appearing before 10:00, 15:00-16:00 and 519 after 19:00 under 0.4 km with the maximum value of 0.81 ppb/h. The OH production rate from O<sub>3</sub> 520 shown a unimodal pattern occurring at 13:00-15:00 under 0.4 km with the maximum value of 6.20 521 ppb/h which was obviously higher than that at low-altitude areas. In addition to direct emission, the 522 heterogeneous reaction of NO<sub>2</sub> on wet surfaces was also an important source of HONO in Nam Co. We 523 found that HONO/NO<sub>2</sub> first increasing and then decreasing with the increasing of H<sub>2</sub>O. The maximum value of HONO/NO<sub>2</sub> appeared around H<sub>2</sub>O being around  $1.0 \times 10^{17}$  molec cm<sup>-3</sup> under 1.0 km, and 524 being around  $1.0-2.0 \times 10^{17}$  molec cm<sup>-3</sup> at 1.0-2.0 km height layer. Moreover, high values of 525 HONO/NO<sub>2</sub> usually accompanied by high aerosol extinction. O<sub>3</sub> under 2.0 km were potentially sourced 526 from Himalayas, northern India, central Pakistan, Nepal, western Bhutan and northern Bangladesh 527 528 through long-range transport. Our results draw a picture of further understanding the spatial and 529 temporal variations in oxidation chemistry under PBL and provided a new perspective for source analysis of major atmospheric components through vertical observation on the TP. 530

# 531 Acknowledgements

532 We firstly would like to thank @Tibet group for effectively organizing the Nam Co observation. We also would like to thank Peking University (Chunxiang Ye's group) and Anhui Institute of Optics and 533 534 Fine Mechanics (Weixiong Zhao's group) to provide the DOAS validation data of HONO, O<sub>3</sub> and NO<sub>2</sub>. 535 We thank the National Oceanic and Atmospheric Administration (NOAA) Air Resources Laboratory 536 (ARL) for providing the open HYSPLIT transport and dispersion model. This study was supported by 537 the National Natural Science Foundation of China (42225504 and U21A2027), the Anhui Provincial 538 Natural Science Foundation (2108085QD180), and the Presidential Foundation of the Hefei Institutes 539 of Physical Science, Chinese Academy Sciences (YZJJ2021QN06).

# 540 **Compliance with ethics guidelines**

- 541 All authors declare that they have no conflict of interest or financial conflicts to disclose.
- 542

# 543 **References**

- 544 [1] Kang, S., Chen, P., Li, C., Liu, B., and Cong, Z.: Atmospheric Aerosol Elements over the Inland Tibetan Plateau:
- 545 Concentration, Seasonality, and Transport, Aerosol Air Qual. Res., 16: 789–800, doi: 10.4209/aaqr.2015.05.0307, 546 2016.
- 547 [2] Xia, X., Zong, X., Cong, Z., Chen, H., Kang, S., and Wang, P.: Baseline continental aerosol over the central 548 Tibetan plateau and a case study of aerosol transport from South Asia, Atmos. Environ., 45, 7370-7378, doi:
- 549 10.1016/j.atmosenv.2011.07.067, 2011.
- [3] Xing, C., Liu, C., Wang, S., Hu, Q., Liu, H., Tan, W., Zhang, W., Li, B., and Liu, J.: A new method to determine
  the aerosol optical properties from multiple-wavelength O<sub>4</sub> absorption by MAX-DOAS observation, Atmos. Meas.
  Tech., 12, 3289-3302, doi.org/10.5194/amt-12-3289-2019, 2019.
- [4] Zhao, F., Liu, C., Cai, Z., Liu, X., Bak, J., Kim, J., Hu, Q., Xia, C., Zhang, C., Sun, Y., Wang, W., and Liu, J.:
  Ozone profile retrievals from TROPOMI: Implication for the variation of tropospheric ozone during the outbreak of
  COVID-19 in China, Sci. Total Environ., 764, 142886, doi.org/10.1016/j.scitotenv.2020.142886, 2021.
- 555 (5) Fang, B., Zhao, W., Xu, X., Zhou, J., Ma, X., Wang, S., Zhang, W., Venables, D.S., and Chen, W.: Portable
- broadband cavity-enhanced spectrometer utilizing Kalman filtering: application to real-time, in situ monitoring of
- 558 glyoxal and nitrogen dioxide, Opt. Express, 25(22), 26910-26922, doi.org/10.1364/OE.25.026910, 2017.
- 559 [6] Kleffmann, J., Wiesen, P.: Technical Note: Quantification of interferences of wet chemical HONO LOPAP 560 measurements under simulated polar conditions, Atmos. Chem. Phys., 8, 6813-6822, 561 doi.org/10.5194/acp-8-6813-2008, 2008.

- 562 [7] Bessho, K., Date, K., Hayashi, M., Ikeda, A., Imai, T., Inoue, H., Kumagai, Y., Miyakawa, T., Murata, H., Ohno,
- 563 T., Okuyama, A., Oyama, R., Sasaki, Y., Shimazu, Y., Shimoji, K., Sumida, Y., Suzuki, M., Taniguchi, H., 564 Tsuchiyama, H., Uesawa, D., Yokota, H., and Yoshida, R.: An Introduction to Himawari-8/9-Japan's New-Generation 565 Geostationary Meteorological Satellites, J. Meteorol. Soc. Jan., 94(2), 151-183, doi: 10.2151/jmsj.2016-009, 2016.
- 566 [8] Veefkind, J.P., de Haan, J.F., Brinksma, E.J., Kroon, M., and Levelt, P.F.: Total Ozone From the Ozone
- 567 Monitoring Instrument (OMI) Using the DOAS Technique, IEEE T. Geosci. Remote Sens., 44(5), 1239-1244, doi: 568 10.1109/TGRS.2006.871204, 2004.
- 569 [9] Griffin, D., Zhao, X., Mclinden, C.A., Boersma, F., Bourassa, A., Dammers, E., Degenstein, D., Eskes, H., Fehr,
- 570 L., Fioletov, V., Hayden, K., Kharol, S.K., Li, S., Makar, P., Martin, R.V., Mihele, C., Mittermeier, R.L., Krotkov, N.,
- 571 Sneep, M., Lamsal, L.N., ter Linden, M., van Geffen, J., Veefkind, P., and Wolde, M.: High-Resolution Mapping of
- 572 Nitrogen Dioxide With TROPOMI: First Results and Validation Over the Canadian Oil Sands, Geophys. Res, Lett., 573 46, 1049-1060, doi: 10.1029/2018GL081095, 2018.
- 574 [10] Su, W., Liu, C., Chan, K.L., Hu, Q., Liu, H., Ji, X., Zhu, Y., Liu, T., Zhang, C., Chen, Y., and Liu, J.: An 575 improved TROPOMI tropospheric HCHO retrieval over China, Atmos. Meas. Tech., 13, 6271-6292, 576 doi.org/10.5194/amt-13-6271-2020, 2020.
- 577 [11] Grell, G.A., Peckham, S.E., Schmitz, R., McKeen, S.A., Frost, G., Skamarock, W.C., and Eder, B.: Fully coupled 578 "online" chemistry WRF model, Atmos, 6957-6975. with the Environ., 39(37), 579 doi.org/10.1016/j.atmosenv.2005.04.027, 2005.
- 580 [12] Shi, G., Yang, L., Wang, Y., Kobayashi, K., Zhu, J., Tang, H., Pan, S., Chen, T., Liu, G., and Wang, Y.: Impact 581 of elevated ozone concentration on yield of four Chinese rice cultivars under fully open-air filed conditions, Agr. 582 Ecosys. Environ., 131(3-4), 178-184, doi.org/10.1016/j.agee.2009.01.009, 2009.
- 583 [13] Yin, X., Kang, S., de Foy, B., Cong, Z., Luo, J., Zhang, L., Ma, Y., Zhang, G., Rupakheti, D., and Zhang, Q.: 584 Surface ozone at Nam Co in the inland Tibetan Plateau: variation, synthesis comparison and regional 585 representativeness, Atmos. Chem. Phys., 17, 11293-11311, doi.org/10.5194/acp-17-11293-2017, 2017.
- 586 [14] Draxler, R.R., Hess, G.: An overview of the HYSPLIT\_4 modelling system for trajectories, Aust. Meteorol. Mag., 587 47, 295-308, 1998.
- 588 [15] Hong, Q., Liu, C., Hu, Q., Xing, C., Tan, W., Liu, H., Huang, Y., Zhu, Y., Zhang, J., Geng, T., and Liu, J.: 589 Evolution of the vertical structure of air pollutants during winter heavy pollution episodes: The role of regional 590 transport and potential sources, Atmos. Res., 228, 106-222, doi.org/10.1016/j.atmosres.2019.05.016, 2019.
- 591 [16] Ou, J., Hu, Q., Liu, H., Hong, Q., Xing, C., Tan, W., Lin, H., Wang, X., Xu, H., Zhu, P., and Liu, W.: Vertical 592 characterization and potential sources of aerosols in different seasons over the Yangtze River Delta using
- 593 ground-based MAX-DOAS, Environ. Pollut., 279, 116898, doi.org/10.1016/j.envpol.2021.116898, 2021.
- 594 [17] Hsu, Y.K., Holsen, T.M., Hopke, P.K.: Comparison of hybrid receptor models to locate PCB sources in Chicago, 595 Atmos. Environ., 37, 545-562, doi.org/10.1016/S1352-2310(02)00886-5, 2003.
- 596 [18] Wang, Y., Zhang, X., Draxler, R.R.: TrajStat: GIS-based software that uses various trajectory statistical analysis 597 methods to identify potential sources from long-term air pollution measurement data, Environ. Model Softw., 24, 598 938-939, doi.org/10.1016/j.envsoft.2009.01.004, 2009.
- 599 [19] Ye, C.: The first constraint of atmospheric oxidative capacity in Namco, a background Tibetan Plateau research 600 site, AGU Fall Meeting Abstracts, 2019:A51C-08, 2019.
- 601 [20] Lin, W., Zhu, T., Song, Y., Zou, H., Tang, M., Tang, X., and Hu, J.: Photolysis of surface O<sub>3</sub> and production 602 potential of OH radicals in the atmosphere over the Tibetan Plateau, J. Geophys. Res.-Atmos., 113, D02309,
- 603 doi:10.1029/2007JD008831, 2008.
- 604 [21] Michoud, V., Kukui, A., Camredon, M., Colomb, A., Borbon, A., Miet, K., Aumont, B., Beekmann, M.,
- 605 Durand-Jolibois, R., Perrier, S., Zapf, P., Siour, G., Ait-Helal, W., Locoge, N., Sauvage, S., Afif, C., Gros, C., Furger, 606 M., Ancellet, G., and Doussin, J.F.: Radical budget analysis in a suburban European site during the MEGAPOLI 607
- summer field campaign, Atmos. Chem. Phys., 12, 11951-11974, doi.org/10.5194/acp-12-11951-2012, 2012.
- 608 [22] Ryan, R.G., Rhodes, S., Tully, M., Wilson, S., Jones, N., Frieß, U., and Schofield, R.: Daytime HONO, NO<sub>2</sub> and 609 aerosol distributions from MAX-DOAS observations in Melbourne, Atmos. Chem. Phys., 18, 13969-13958, 610 doi.org/10.5194/acp-18-13969-2018, 2018.
- 611 [23] Xue, C., Zhang, C., Ye, C., Liu, P., Catoire, V., Krysztofiak, G., Chen, H., Ren, Y., Zhao, X., Wang, J., Zhang, F.,
- 612 Zhang, C., Zhang, J., An, J., Wang, T., Chen, J., Kleffmann, J., Mellouki, A., and Mu, Y.: HONO budget and its role 613 nitrate formation in rural North China Plain, Environ. Sci. Tech., 54, 18, 11048-11057, in 614 doi.org/10.1021/acs.est.0c01832, 2020.
- 615 [24] Xing, C., Liu, C., Wu, H., Lin, J., Wang, F., Wang, S., and Gao, M.: Ground-based vertical profile observations
- 616 of atmospheric composition on the Tibetan Plateau (2017-2019), Earth Syst. Sci. Data, 13, 4897-4912, 617 doi.org/10.5194/essd-13-4897-2021, 2021a.
- 618 [25] Xing, C., Liu, C., Hu, Q., Fu, Q., Wang, S., Lin, H., Zhu, Y., Wang, S., Wang, W., Javed, Z., Ji, X., Liu, J.: 619 Vertical distributions of wintertime atmospheric nitrogenous compounds and the corresponding OH radicals 620 production in Leshan, southwest China, J. Environ. Sci., 105, 44-55, doi.org/10.1016/j.jes.2020.11.019.
- 621 [26] Luo, S., Holland, F., Rohrer, F., Lu, K., Bohn, B., Brauers, T., Chang, C.C., Fuchs, H., Häseler, R., Kita, K., 622 Kondo, Y., Li, X., Shao, M., Zeng, L., Wahner, A., Zhang, Y., Wang, W., Hofzumahaus, A.: Atmospheric OH

- reactivities in the Pearl River Delta-China in summer 2006: measurement and model results, Atmos. Chem. Phys., 10,
  11243-11260, doi.org/10.5194/acp-10-11243-2010, 2010.
- 625 [27] Yang, Y., Wang, Y., Huang, W., Yao, D., Zhao, S., Wang, Y., Ji, D., Zhang, R., Wang, Y.: Parameterized 626 atmospheric oxidation capacity and speciated OH reactivity over a suburban site in the North China Plain: A 627 Total comparative study between summer and winter, Sci. Environ., 773, 145264, 628 doi.org/10.1016/j.scitotenv.2021.145264, 2021.
- 629 [28] Ma, Y., Zhong, L., Su, Z.: Energy and water cycles in the third pole, Water, 14(7), 1175, 630 doi.org/10.3390/w14071175, 2022.
- 631 [29] Kang, S., Zhang, Y., Chen, P., Guo, J., Zhang, Q., Cong, Z., Kaspari, S., Tripathee, L., Gao, T., Niu, H., Zhong,
- K., Chen, X., Hu, Z., Li, X., Li, Y., Neupane, B., Yan, F., Rupakheti, D., Gul, C., Zhang, W., Wu, G., Yang, L., Wang,
  Z., Li, C.: Black carbon and organic carbon dataset over the Third Pole, Earth Syst. Sci. Data, 14, 683–707,
- 634 doi.org/10.5194/essd-14-683-2022, 2022.
- [30] Ma, Y., Hu, Z., Xie, Z., Ma, W., Wang, B., Chen, X., Li, M., Zhong, L., Sun, F., Gu, L., Han, C., Zhang, L., Liu,
  X., Ding, Z., Sun, G., Wang, S., Wang, Y., and Wang, Z.: A long-term (2005–2016) dataset of integrated
  land-atmosphere interaction observations on the Tibetan Plateau, Earth Syst. Sci. Data, 12, 2937-2957,
  doi:10.5194/essd-12-2937-2020, 2020.
- [31] Qu, B., Zhang, Y., Kang, S., Sillanpää, M.: Water quality in the Tibetan Plateau: Major ions and trace elements
  in rivers of the "Water Tower of Asia", Sci. Total Environ., 649, 571-581, doi.org/10.1016/j.scitotenv.2018.08.316,
  2019.
- [32] Zhou, S., Sun, F., Wang, M., Zhou, S., and Qing, Y.: Effects of atmospheric heat source on the Tibetan Plateau
- vortex in different stages: A case study in June 2016, Atmosphere, 13(5), 689, doi.org/10.3390/atmos13050689, 2022.
- 644 [33] Liu, J., Guan, X., Gao, Z., Huang, X., Ma, J., He, Y., and Xie, T.: Inter-decadal variability of the heat source over 645 the Tibetan Plateau, Clim. Dynam., 58, 729-739, doi.org/10.1007/s00382-021-05929-z, 2022.
- [34] Chen, P., Kang, S., Bai, J., Sillanpää, M., Li, C.: Yak dung combustion aerosols in the Tibetan Plateau: Chemical
  characteristics and influence on the local atmospheric environment, Atmos. Res., 156, 58-66,
  doi.org/10.1016/j.atmosres.2015.01.001, 2015.
- 649 [35] Boos, W. R. and Kuang, Z.: Dominant control of the South Asian monsoon by orographic insulation versus 650 plateau heating, Nature, 463, 218–222, doi:10.1038/nature08707, 2010.
- [36] Yanai, M., Li, C., and Song, Z.: Seasonal Heating of the Tibetan Plateau and Its Effects on the Evolution of the
   Asian Summer Monsoon, J. Meteorol. Soc. Jpn. Ser. II, 70, 319–351, doi:10.2151/jmsj1965.70.1B\_319, 1992.
- 653 [37] Li, C., Zou, Q., Xu, X., and Gao, S.: Water vapor transport around the Tibetan Plateau and its effect on summer 654 rainfall over the Yangtze River valley, J. Meteorol. Res., 30, 472-482, doi: 10.1007/s13351-016-5123-1, 2016.
- [38] Lei, Y., Zhu, Y., Wang, B., Yao, T., Yang, K., Zhang, X., Zhai, J., and Ma, N.: Extreme lake level changes in the
  Tibetan Plateau associated with the 2015/2016 El Niño, Geophys. Res. Lett., 46, 11, 5889-5898,
  doi.org/10.1029/2019GL081946, 2019.
- [39] Hsu, H-H., and Liu, X.: Relationship between the Tibetan Plateau heating and East Asian summer monsoon
   rainfall, Geophys. Res. Lett., 30, 20, doi.org/10.1029/2003GL017909, 2003.
- [40] Zhang, L., Guo, X., Zhao, T., Gong, S., Xu, X., Li, Y., Luo, L., Gui, K., Wang, H., Zheng, Y., and Yin, X.: A
  modelling study of the terrain effects on the haze pollution in Sichuan Basin, Atmos. Environ., 196, 77-85,
  doi.org/10.1016/j.atmosenv.2018.10.007, 2019.
- 663 [41] Barnett, T. P., Adam, J. C., and Lettenmaier, D. P.: Potential impacts of a warming climate on water availability 664 in snow-dominated regions, Nature, 438, 303–309, doi:10.1038/nature04141, 2005.
- 665 [42] Bolch, T., Kulkarni, A., Kaab, A., Huggel, C., Paul, F., Cogley, J. G., Frey, H., Kargel, J. S., Fujita, K., Scheel,
- 666 M., Bajracharya, S., and Stoffel, M.: The State and Fate of Himalayan Glaciers, Science, 336, 310–314, doi:10.1126/science.1215828, 2012.
- [43] Cong, Z., Kang, S., Kawamura, K., Liu, B., Wan, X., Wang, Z., Gao, S., and Fu, P.: Carbonaceous aerosols on
   the south edge of the Tibetan Plateau: concentrations, seasonality and sources, Atmos. Chem. Phys., 15, 1573–1584,
- 670 https://doi.org/10.5194/acp-15-1573-2015, 2015.
- 671 [44] Kang, S. C., Huang, J., Wang, F. Y., Zhang, Q. G., Zhang, Y. L., Li, C. L., Wang, L., Chen, P. F., Sharma, C. M.,
- Li, Q., Sillanpaa, M., Hou, J. Z., Xu, B. Q., and Guo, J. M.: Atmospheric Mercury Depositional Chronology
- Reconstructed from Lake Sediments and Ice Core in the Himalayas and Tibetan Plateau, Environ. Sci. Technol., 50,
  2859–2869, doi:10.1021/acs.est.5b04172, 2016.
- [45] Ran, L., Deng, Z., Wu, Y., Li, J., Bai, Z., Lu, Y., Zhuoga, D., and Bian J.: Measurement report: Vertical profiling of particle size distributions over Lhasa, Tibet tethered balloon-based in situ measurements and source
- 677 apportionment, Atmos. Chem. Phys., 22, 6217–6229, doi.org/10.5194/acp-22-6217-2022, 2022.
- [46] Wang, K., Hattori, S., Lin, M., Ishino, S., Alexander, B., Kamezaki, K., Yoshida, N., and Kang, S.: Isotopic
   constraints on atmospheric sulfate formation pathways in the Mt. Everest region, southern Tibetan Plateau, Atmos.
- 680 Chem. Phys., 21, 8357–8376, https://doi.org/10.5194/acp-21-8357-2021, 2021.
- [47] Che, J. and Zhao, P.: Characteristics of the summer atmospheric boundary layer height over the Tibetan Plateau
- 682 and influential factors, Atmos. Chem. Phys., 21, 5253–5268, https://doi.org/10.5194/acp-21-5253-2021, 2021.

- 683 [48] Sun, Y., Yin, H., Cheng, Y., Zhang, Q., Zheng, B., Notholt, J., Lu, X., Liu, C., Tian, Y., Liu, J.: Quantifying 684 variability, source, and transport of CO in the urban areas over the Himalayas and Tibetan Plateau, Atmos. Chem.
- 685 Phys., 21, 9201–9222, https://doi.org/10.5194/acp-21-9201-2021, 2021.
- 686 [49] Li, R., Zhao, Y., Zhou, W., Meng, Y., Zhang, Z., and Fu, H.: Developing a novel hybrid model for the estimation 687 of surface 8-h ozone (O<sub>3</sub>) across the remote Tibetan Plateau during 2005-2018, Atmos. Chem. Phys., 20, 6159–6175,
- 688 https://doi.org/10.5194/acp-20-6159-2020.
- 689 [50] Gao, M., Gao, J., Zhu, B., Kumar, R., Lu, X., Song, S., Zhang, Y., Jia, B., Wang, P., Beig, G., Hu, J., Ying, Q.,
- 690 Zhang, H., Sherman, P., and McElroy, M. B.: Ozone pollution over China and India: seasonality and sources, Atmos. 691 Chem. Phys., 20, 4399–4414, https://doi.org/10.5194/acp-20-4399-2020, 2020.
- 692 [51] Rawat, P., and Naja, M.: Remote sensing study of ozone, NO<sub>2</sub>, and CO: some contrary effects of SARS-CoV-2 693 lockdown over India, Environ. Sci. Pollut. Res., 29, 22515-22530, doi:10.1007/s11356-021-17441-2, 2022.
- 694 [52] Huang, J., Minnis, P., Yi, Y., Tang, Q., Wang, X., Hu, Y., Liu, Z., Ayers, K., Trepte, C., and Winker, D.: 695 Summer dust aerosols detected from CALIPSO over the Tibetan Plateau, Geophys. Res. Lett., 34, L18805,
- 696 https://doi.org/10.1029/2007GL029938, 2007.
- 697 [53] Li, R., Zhao, Y., Zhou, W., Meng, Y., Zhang, Z., and Fu, H.: Developing a novel hybrid model for the estimation 698 of surface 8 h ozone (O3) across the remote Tibetan Plateau during 2005–2018, Atmos. Chem. Phys., 20, 6159–6175, 699 https://doi.org/10.5194/acp-20-6159-2020, 2020.
- 700 [54] Zhu, J., Xia, X., Che, H., Wang, J., Cong, Z., Zhao, T., Kang, S., Zhang, X., Yu, X., and Zhang, Y.: 701 Spatiotemporal variation of aerosol and potential long-range transport impact over the Tibetan Plateau, China, Atmos. 702 Chem. Phys., 19, 14637–14656, https://doi.org/10.5194/acp-19-14637-2019, 2019.
- 703 [55] Xu, X., Sun, C., Chen, D., Zhao, T., Xu, J., Zhang, S., Li, J., Chen, B., Zhao, Y., Xu, H., Dong, L., Sun, X., and 704 Zhu, Y.: A vertical transport window of water vapor in the troposphere over the Tibetan Plateau with implications for 705 global climate change, Atmos. Chem. Phys., 22, 1149–1157, https://doi.org/10.5194/acp-22-1149-2022, 2022.
- 706 [56] Xu, X., Wu, H., Yang, X., and Xie, L.: Distribution and transport characteristics of dust aerosol over Tibetan 707 Plateau and Taklimakan Desert in China using MERRA-2 and CALIPSO data, Atmos. Environ., 237, 117670, 708 https://doi.org/10.1016/j.atmosenv.2020.117670, 2020.
- 709 [57] Yang, K., Koike, T., and Yang, D.: Surface flux parameterization in the Tibetan Plateau, Bound.-Lay. Meteorol., 710 106, 245-262, doi:10.1023/A:1021152407334, 2003.
- 711 [58] Seidel, D. J., Ao, C. O., and Li, K.: Estimating climatological planetary boundary layer heights from radiosonde 712 observations: Comparison of methods and uncertainty analysis, J. Geophys. Res., 115, D16113, 713 https://doi.org/10.1029/2009JD013680, 2010.
- 714 [59] Dong, Q., Huang, Z., Li, W., Li, Z., Song, X., Liu, W., Wang, T., Bi, J., and Shi, J.: Polarization lidar 715 measurements of dust optical properties at the junction of the Taklimakan Desert-Tibetan Plateau, Remote Sens., 716 14(3), 558, https://doi.org/10.3390/rs14030558, 2022.
- 717 [60] Zhang, J., Xia, X., and Wu, X.: First in situ UV profile across the UTLS accompanied by ozone measurement 718 over the Tibetan Plateau, J. Environ., Sci., 98, 71–76, doi:10.1016/j.jes.2020.05.020, 2020.
- 719 [61] Fang, X., Li, T., Ban, C., Wu, Z., Li, J., Li, F., Cen, Y., and Tian, B.: A mobile differential absorption lidar for 720 simultaneous observations of tropospheric and stratospheric ozone over Tibet, Opt. Express, 27, 4126–4139, 721 doi:10.1364.OE.27.004126, 2019.
- 722 [62] Wang, Y., Pukite, J., Wagner, T., Donner, S.; Beirle, S., Hilboll, A., Vrekoussis, M., Richter, A., Apituley, A.,
- 723 Piters, A., Allaart, M., Eskes, H., Frumau, A., van Roozendael, M., Lampel, J., Platt, U., Schmitt, S., Swart, D., and
- 724 Vonk, J.: Vertical profles of tropospheric ozone from MAX-DOAS measurement during the CINDI-2 campaign: part 725 1-Development of a new retrieval algorithm. J. Geophys. Res. Atmos. 123 (18), 10-637.
- 726 https://doi.org/10.1029/2018JD028647, 2018.
- 727 [63] Xing, C., Liu, C., Wang, S., Chan, K.L., Gao, Y., Huang, X., Su, W., Zhang, C., Dong, Y., Fan, G., Zhang, T.,
- 728 Chen, Z., Hu, Q., Su, H., Xie, Z., and Liu, J.: Observations of the vertical distributions of summertime atmospheric 729 pollutants and the corresponding ozone production in Shanghai, China. Atmos. Chem. Phys. 17, 14275–14289. 730 https://doi.org/10.5194/acp-17-14275-2017, 2017.
- 731 [64] Xing, C., Liu, C., Wang, S., Hu, Q., Liu, H., Tan, W., Zhang, W., Li, B., and Liu, J.; A new method to determine 732 the aerosol optical properties from multiple-wavelength O4 absorptions by MAX-DOAS observation. Atmos. Meas. 733 Tech. 12, 3289–3302. https://doi.org/10.5194/amt-12-3289-2019, 2019.
- 734 [65] Xing, C., Liu, C., Hu, Q., Fu, Q., Lin, H., Wang, S., Su, W., Wang, W., Javed, Z., and Liu, J.: Identifying the 735
- wintertime sources of volatile organic compounds (VOCs) from MAX-DOAS measured formaldehyde and glyoxal in
- 736 Chongqing, Southwest China. Sci. Total Environ. 715, 136258 https://doi.org/10.1016/j.scitotenv.2019.136258, 2020.
- 737 [66] Ye, D. Z., and Gao, Y. X.: The Meteorology of the Tibetan Plateau (in Chinese), 278pp., Science Press, Beijing, 738 pp. 39–48, 1979.
- 739 [67] Liu, Y., and Li, W.: Deepening of the ozone valley over Tibetan Plateau and its possible influences (Chinese with 740 English abstract), Acta Meteorologica Sinica, 59(1), 97–106, 2001.
- 741 [68] Yang, J., Kang, S., Hu, Y., Chen, X., Rai, M.: Influence of South Asian biomass burning on ozone and aerosol
- 742 concentrations over the Tibetan Plateau, Adv. Atmos. Sci., 39, 1184-1197, doi:10.1007/s00376-022-1197-0, 2022.

- [69] Yu, J., Meng, L., Chen, Y., Zhang, H., and Liu, J.: Ozone profiles, precursors, and vertical distribution in urban
  Lhasa, Tibetan Plateau, Remote Sens., 14(11), 2533, https://doi.org/10.3390/rs14112533, 2022.
- [70] Li, M., Mao, J., Chen, S., Bian, J., Bai, Z., Wang, X., Chen, W., and Yu, P.: Significant contribution of lightning
  NO<sub>x</sub> to summertime surface O<sub>3</sub> on the Tibetan Plateau, Sci, Total, Environ., 829, 154639,
  doi.org/10.1016/j.scitotenv.2022.154639, 2022.
- [71] Zhou, L., Zhang, X., Zhang, J.: Temporal and spatial distributions of atmospheric hydroxyl radicals based on the
   observation with the aura microwave limb sounder. Science & Technology Review, 33(17): 69-77, 2015.
- 750 [72] Yin, X., Kang, S., de Foy, B., Cong, Z., Luo, J., Zhang, L., Ma, Y., Zhang, G., Rupakheti, D., and Zhang, Q.:
- Surface ozone at Nam Co in the inland Tibetan Plateau: variation, synthesis comparison and regional
   representativeness, Atmos. Chem. Phys., 17, 11293–11311, https://doi.org/10.5194/acp-17-11293-2017, 2017.
- [73] Xu, X., Zhang, H., Lin, W., Wang, Y., Xu, W., and Jia, S.: First simultaneous measurements of peroxyacetyl nitrate (PAN) and ozone at Nam Co in the central Tibetan Plateau: impacts from the PBL evolution and transport processes, Atmos. Chem. Phys., 18, 5199–5217, https://doi.org/10.5194/acp-18-5199-2018, 2018.
- [74] Bi, H., Chen, S., Zhao, D., Lu, F., Chen, Y., and Guan, Y.: Aerosol optical properties and its direct radiative
  forcing over Tibetan Plateau from 2006 to 2017, Particuology, 74, 64-73, https://doi.org/10.1016/j.partic.2022.05.007,
  2023.
- [75] Yang, J., Kang, S., and Ji, Z.: Critical contribution of south Asian residential emissions to atmospheric black
  carbon over the Tibetan plateau, Sci. Total Environ., 709, 135923, https://doi.org/10.1016/j.scitotenv.2019.135923,
  2020.
- 762 [76] Zhang, X., Ming, J., Li, Z., Wang, F., and Zhang, G.: The online measured black carbon aerosol and source 763 orientations in the Nam Co region, Tibet, Environ. Sci. Pollut. Res., 24, 25021-25033, doi: 764 10.1007/s11256.017.0165.1.2017
- 764 10.1007/s11356-017-0165-1, 2017.
- [77] Yang, J., Duan, K., Kang, S., Shi, P., and Ji, Z.: Potential feedback between aerosols and meteorological
  conditions in a heavy pollution event over the Tibetan Plateau and Indo-Gangetic Plain, Clim. Dyn., 48(9), 2901-2917,
  doi:10.1007/s00382-016-3240-2, 2017.
- [78] Wan, X., Kang, S., Wang, Y., Xin, J., Liu, B., Guo, Y., Wen, T., Zhang, G., and Cong, Z.: Size distribution of
  carbonaceous aerosols at a high-altitude site on the central Tibetan Plateau (Nam Co Station, 4730 m a.s.l.), Atmos.
  Res, 153, 155-164, doi:10.1016/j.atmosres.2014.08.008, 2015.
- [79] Li, C., Bosch, C., Kang, S., Andersson, A., Chen, P., Zhang, Q., Cong, Z., Chen, B., Qin, D., and Gustafsson, O.:
  Sources of black carbon to the Himalayan-Tibetan Plateau glaciers, Nat. Commun., 7, 12574, doi:10.1038/ncomms12574, 2016.
- [80] Lei, Y., Yang, K., Wang, B., Sheng, Y., Bird, B.W., Zhang, G., and Tian, L.: Response of inland lake dynamics
  over the Tibetan Plateau to climate change, Clim. Chang, 125, 281-290, doi:10.1007/210584-014-1175-3, 2014.
- [81] Zhu, G., Guo, H., Qin, D., Pan, H., Jia, W., and Ma, X.: Contribution of recycled moisture to precipitation in the
  monsoon marginal zone: Estimate based on stable isotope data, J. Hydrol., 569, 423-435,
  doi:10.1016/j.jhydrol.2018.12.014, 2019.
- [82] Boxe, C.: Nitrate photochemistry and interrelated chemical phenomena in ice[M]. California Institute ofTechnology, 2005.
- [83] Xu, R., Tie, X., Li, G., Zhao, S., Cao, J., Feng, T., and Long, X.: Effect of biomass burning on black carbon (BC)
- in South Asia and Tibetan Plateau: The analysis of WRF-Chem modeling, Sci. Total Environ., 645, 901-912,
   doi:10.1016/j.scitotenv.2018.07.165, 2018.
- [84] Neupane, B., Kang, S., Chen, P., Zhang, Y., Ram, K., Rupakheti, D., Tripathee, L., Sharma, C.M., Cong, Z., Li, C., Hou, J., Xu, M., and Thapa, P.: Historical black carbon reconstruction from the lake sediments of the
- 786 Himalayan-Tibetan Plateau, Environ. Sci. Tech., 53, 5641-5651, doi:10.1021/acs.est.8b07025, 2019.
- [85] Xu, K., Zhong, L., Ma, Y., Zou, M., and Huang, Z.: A study on the water vapor transport trend and water vapor sources of the Tibetan Plateau, Theor. Appl. Climatol., 140, 1031-1042, doi:10.1007/s00704-020-03142-2, 2020.
- [86] Xu, Y., Kang, S., Zhang, Y., and Zhang, Y.: A method for estimating the contribution of evaporative vapor from
  Nam Co to local atmospheric vapor based on stable isotopes of water bodies, Chinese Sci. Bull., 56(14), 1511-1517,
  doi:10.1007/s11434-011-4467-2, 2011.
- [87] Chen, P., Kang, S., Yang, J., Pu, T., Li, C., Guo, J., and Tripathee, L.: Spatial and temporal variations of gaseous and particle pollutants in six sites in Tibet, China, during 2016-2017, Aerosol Air Qual. Res., 19, 516-527, doi:10.4209/aaqr.2018.10.0360, 2019.
- [88] Wang, T., Xue, L., Brimblecombe, P., Lam, Y, Li, L. and Zhang, L.: Ozone pollution in China: A review of
  concentrations, meteorological influences, chemical precursors, and effects. Sci. Total Environ., 575, 1582–1596,
  doi:10.1016/j.scitotenv.2016.10.081, 2017.
- 798 [89] Pokharel, M., Guang, J., Liu, B., Kang, S., Ma, Y., Holben, B.N., Xia, X., Xin, J., Ram, K., Rupakheti, D., Wan,
- X., Wu, G., Bhattarai, H., Zhao, C., and Cong, Z.: Aerosol properties over Tibetan Plateau from a decade of AERONET measurements: Baseline, types, and influencing factors, J. Geophys. Res.: Atmos., 124, 13357-13374,
- 801 doi:10.1029/2019JD031293, 2019.
- 802 [90] Cong, Z., Kang, S., Smirnov, A., and Holben, B.: Aerosol optical properties at Nam Co, a remote site in central
- 803 Tibetan Plateau, Atmos. Res, 92, 42-48, doi:10.1016/j.atmosres.2008.08.005, 2009.

- [91] Qian, Y., Wang, H., Zhao, C., Zhao, C., Chen, S., Hu, X., and Kang, S.: Understanding third pole atmospheric
  dynamics and land surface processes and their associations with the cryosphere, air quality, and climate change, Adv.
  Atmos. Sci., 39, 1017-1020, doi:10.1007/s00376-022-2004-7, 2022.
- 807 [92] Xu, L., Liu, H., Du, Q., and Xu, X.: The assessment of the planetary boundary layer schemes in WRF over the 808 central Tibetan Plateau, Atmos. Res., 230, 104644, doi:10.1016/j.atmosres.2019.104644, 2019.
- [93] Yang, J., and Duan, K.: Effects of initial drivers and land use on WRF modeling for near-surface fields and
  atmospheric boundary layer over the northeastern Tibetan Plateau, Adv. Meteorol., 2016, 7849249,
  doi:10.1155/2016/7849249, 2016.
- 812 [94] Kleffmann, J., Gavriloaiei, T., Hofzumahaus, A., Holland, F., Koppmann, R., Rupp, L., Schlosser, E., Siese, M.,
- and Wahner, A.: Daytime formation of nitrous acid: A major source of OH radicals in a forest, Geophys. Res. Lett.,
   32(5), doi:10.1029/2005GL022524, 2005.
- [95] Su, H., Cheng, Y., Shao, M., Gao, D., Yu, Z., Zeng, L., Slanina, J., Zhang, Y., and Wiedensohler, A.: Nitrous
  acid (HONO) and its daytime sources at a rural site during the 2004 PRIDE-PRD experiment in China, J. Geophys.
  Page 113 D1/312 doi:10.1029/2007JD009060.2008
- 817 Res., 113, D14312, doi:10.1029/2007JD009060, 2008.
- 818 [96] Yang, Y., Li, X., Zu, K., Lian, C., Chen, S., Dong, H., Feng, M., Liu, H., Liu, J., Lu, K., Lu, S., Ma, X., Song, D.,
- Wang, W., Yang, S., Yang, X., Yu, X., Zhu, Y., Zeng, L., Tan, Q., and Zhang, Y.: Elucidating the effect of HONO
  and O<sub>3</sub> pollution by a case study in southwest China, Sci. Total Environ., 756, 144127,
  doi:10.1016/j.scitotenv.2020.144127, 2021.
- [97] Liu, T., Hong, Y., Li, M., Xu, L., Chen, J., Bian, Y., Yang, C., Dan, Y., Zhang, Y., Xue, L., Zhao, M., Huang, Z.,
  and Wang, H.: Atmospheric oxidation capacity and ozone pollution mechanism in a coastal city of southeastern China:
  analysis of a typical photochemical episode by an observation-based model, Atmos. Chem. Phys., 22, 2173-2190,
- 825 doi:10.5194/acp-22-2173-2022, 2022.
- [98] Xu, S., Wang, S., Xia, M., Lin, H., Xing, C., Ji, X., Su, W., Tan, W., Liu, C., and Hu, Q.: Observations by
  ground-based MAX-DOAS of the vertical characters of winter pollution and the influencing factors of HONO
  generation in Shanghai, China, Remote Sens., 13, 3518, doi:10.3390/rs13173518, 2021.
- [99] Hendrick, F., Müller, J.F., Clémer, K., Wang, P., De Mazière, M., Fayt, C., Gielen, C., Hermans, C., Ma, J.,
  Pinardi, G., Stavrakou, T., Vlemmix, T., Van Roozendael, M.: Four years of ground-based MAX-DOAS observations
  of HONO and NO2 in the Beijing area, Atmos. Chem. Phys., 14, 765–781, doi:10.5194/acp-14-765-2014, 2014.
- [100] Cui, L., Li, R., Fu, H., Li, Q., Zhang, L., George, C., and Chen, J.: Formation features of nitrous acid in the offshore area of the East China Sea, Sci. Total Environ., 682, 138-150, doi: 10.1016/j.scitotenv.2019.05.004, 2019.
- [101] Yang, J., Shen, H., Guo, M., Zhao, M., Jiang, Y., Chen, T., Liu, Y., Li, H., Zhu, Y., Meng, H., Wang, W., and
  Xue, L.: Strong marine-derived nitrous acid (HONO) production observed in the coastal atmosphere of northern
  China, Atmos. Environ., 244, 117948, doi: 10.1016/j.atmosenv.2020.117948, 2021.
- [102] Liu, Y., Nie, W., Xu, Z., Wang, T., Wang, R., Li, Y., Wang, L., Chi, X., and Ding, A.: Semi-quantitative
  understanding of source contribution to nitrous acid (HONO) based on 1 year of continuous observation at the
  SORPES station in eastern China, Atmos. Chem. Phys., 19, 13289–13308, doi: 10.5194/acp-19-13289-2019, 2019.
- [103] Jena, C., Ghude, S.D., Pfister, G.G., Chate, D.M., Kumar, R., Beig, G., Surendran, D.E., Fadnavis, S., and Lal,
  D.M.: Influence of springtime biomass burning in South Asia on regional ozone (O3): A model based case study,
  Atmos. Environ., 100, 37-47, doi:10.1016/j.atmosenv.2014.10.027, 2015.
- [104] Xing, L., Bei, N., Guo, J., Wang, Q., Liu, S., Han, Y., Pongpiachan, S., and Li, G.: Impacts of biomass burning
  in peninsular southeast Asia on PM<sub>2.5</sub> concentration and ozone formation in southeastern China during springtime-A
  case study, J. Geophys. Res.: Atmos., 126(22), e2021JD034908, doi:10.1029/2021JD034908, 2021.
- [105] Kumar, R., Naja, M., Pfister, G.G., Barth, M.C., Wiedinmyer, C., and Brasseur, G.P.: Simulations over South
  Asia using the Weather Research and Forecasting model with Chemistry (WRF-Chem): chemistry evaluation and
  initial results, Geosci. Model Dev., 5, 619–648, doi:10.5194/gmd-5-619-2012, 2012.
- [106] Sharma, A., Ojha, N., Pozzer, A., Mar, K.A., Beig, G., Lelieveld, J., and Gunthe, S.S.: WRF-Chem simulated surface ozone over south Asia during the pre-monsoon: effects of emission inventories and chemical mechanisms,
- 851 Atmos. Chem. Phys., 17, 14393–14413, doi: 10.5194/acp-17-14393-2017, 2017.
- [107] Cristofanelli, P., Bracci, A., Sprenger, M., Marinoni, A., Bonafè, U., Calzolari, F., Duchi, R., Laj, P., Pichon, J.
   M., Roccato, F., Venzac, H., Vuillermoz, E., and Bonasoni, P.: Tropospheric ozone variations at the Nepal Climate
- M., Roccato, F., Venzac, H., Vuillermoz, E., and Bonasoni, P.: Tropospheric ozone variations at the Nepal Climate
  ObservatoryPyramid (Himalayas, 5079 m a.s.l.) and influence of deep stratospheric intrusion events, Atmos. Chem.
  Phys., 10, 6537–6549, doi:10.5194/acp-10-6537-2010, 2010.
- [108] Chen, X. L., Ma, Y. M., Kelder, H., Su, Z., and Yang, K.: On the behaviour of the tropopause folding events
  over the Tibetan Plateau, Atmos. Chem. Phys., 11, 5113–5122, doi:10.5194/acp-11-5113-2011, 2011.
- 858 [109] Škerlak, B., Sprenger, M., and Wernli, H.: A global climatology of stratosphere–troposphere exchange using the
- 859 ERA-Interim data set from 1979 to 2011, Atmos. Chem. Phys., 14, 913–937, doi:10.5194/acp-14-913-2014, 2014.
- [110] Putero, D., Cristofanelli, P., Sprenger, M., Škerlak, B., Tositti, L., and Bonasoni, P.: STEFLUX, a tool for
  investigating stratospheric intrusions: application to two WMO/GAW global stations, Atmos. Chem. Phys., 16,
  14203–14217, doi:10.5194/acp-16-14203-2016, 2016.

- 863 [111] Fu, X., Wang, T., Zhang, L., Li, Q., Wang, Z., Xia, M., Yun, H., Wang, W., Yu, C., Yue, D., Zhou, Y., Zheng, 864 J., and Han, R.: The significant contribution of HONO to secondary pollutants during a severe winter pollution event 865
- in southern China, Atmos. Chem. Phys., 19, 1–14, doi: 10.5194/acp-19-1-2019, 2019.
- 866 [112] Ren, Y., Stieger, B., Spindler, G., Grosselin, B., Mellouki, A., Tuch, T., Wiedensohler, A., and Herrmann, H.: 867 Role of the dew water on the ground surface in HONO distribution: a case measurement in Melpitz, Atmos. Chem. 868 Phys., 20, 13069–13089, doi: 10.5194/acp-20-13069-2020, 2020.
- 869 [113] Crilley, L.R., Kramer, L.J., Pope, F.D., Reed, C., Lee, J.D., Carpenter, L.J., Hollis, L.D.J., Ball, S.M., and Bloss,
- 870 W.J.: Is the ocean surface a source of nitrous acid (HONO) in the marine boundary layer? Atmos. Chem. Phys., 21, 871 18213-18225, doi: 10.5194/acp-21-18213-2021, 2021.
- 872 [114] Li, S., Song, W., Zhan, H., Zhang, Y., Zhang, X., Li, W., Tong, S., Pei, C., Wang, Y., Chen, Y., Huang, Z.,
- 873 Zhang, R., Zhu, M., Fang, H., Wu, Z., Wang, J., Luo, S., Fu, X., Xiao, S., Huang, X., Zeng, J., Zhang, H., Chen, D.,
- 874 Gligorovski, S., Ge, M., George, C., and Wang, X.: Contribution of vehicle emission and NO<sub>2</sub> surface conversion to
- 875 nitrous acid (HONO) in urban environments: Implications from tests in a tunnel, Environ. Sci. Technol., 55(23),
- 876 15616-15624, doi:10.1021/acs.est.1c00405, 2021.
- 877 [115] Chai, J., Dibb, J.E., Anderson, B.E., Bekker, C., Blum, D.E., Heim, E., Jordan, C.E., Joyce, E.E., Kaspari, J.H., 878 Munro, H., Walters, W.W., and Hastings, M.G.: Isotopic evidence for dominant secondary production of HONO in 879
- near-ground wildfire plumes, Atmos. Chem. Phys., 21, 13077–13098, doi: 10.5194/acp-21-13077-2021, 2021.
- 880 [116] Cui, L., and Wang, S.: Mapping the daily nitrous acid (HONO) concentrations across China during 2006-2017 881 through ensemble machine-learning algorithm, Sci. Total Environ., 785, 147325, doi:10.1016/j.scitotenv.2021.147325, 882 2021.
- 883 [117] Cui, L., Li, R., Fu, H., Meng, Y., Zhao, Y., Li, Q., and Chen, J.: Nitrous acid emission from open burning of 884 major crop residues in mainland China, Atmos. Environ., 244, 117950, doi:10.1016/j.atmosenv.2020.117950, 2021.
- 885 [118] Su, H., Cheng, Y., Oswald, R., Behrendt, T., Trebs, I., Meixner, F.X., Andreae, M.O., Cheng, P., Zhang, Y., and
- 886 Poschl, U.: Soil nitrite as a source of atmospheric HONO and OH radicals, Science, 333(6049), 1616-1618, 887 doi:10.1126/science.1207687, 2011.
- 888 [119] Lin, F., Liu, C., Hu, X., Fu, Y., Zheng, X., Wang, R., Zhang, W., and Cao, G.: Characterizing nitric oxide 889 emissions from two typical alpine ecosystems, J. Environ. Sci., 77, 312-322, doi:10.1016/j.jes.2018.08.011, 2019.
- 890 [120] Gil, J., Kim, J., Lee, M., Lee, G., Lee, D., Jung, J., An, J., Hong, J., Cho, S., Lee, J., and Long, R.: The role of 891 HONO in O3 formation and insight into its formation mechanism during the KORUS-AQ Campaign, Atmos. Chem. 892 Phys. Disscu., doi: 10.5194/acp-2019-1012, 2019.
- [121] Wen, L., Chen, T., Zheng, P., Wu, L., Wang, X., Mellouki, A., Xue, L., and Wang, W.: Nitrous acid marine 893 894 boundary layer over eastern Bohai Sea, China: Characteristics, sources, and implications, Sci. Total Environ., 670, 895 282-291, doi:10.1016/j.scitotenv.2019.03.225, 2019.
- 896 [122] Lu, X., Wang, Y., Li, J., Shen, L., and Fung, J.C.H.: Evidence of heterogeneous HONO formation from aerosols 897 and the regional photochemical impact of this HONO source, Environ. Res. Lett. 13, 114002, 898 doi:10.1088/1748-9326aae492, 2018.
- 899 [123] Cui, L., Li, R., Zhang, Y., Meng, Y., Fu, H., and Chen, J.: An observational study of nitrous acid (HONO) in 900 Shanghai, China: The aerosol impact on HONO formation during the haze episodes, Sci. Total Environ., 630, 901 1057-1070, doi:10.1016/j.scitotenv.2018.02.063, 2018.
- 902 [124] Wang, S., Zhou, R., Zhao, H., Wang, Z., Chen, L., and Zhou, B.: Long-term observation of atmospheric nitrous
- 903 acid (HONO) and its implication to local NO<sub>2</sub> levels in Shanghai, China, Atmos. Environ., 77, 718–724,
- 904 doi:10.1016/j.atmosenv.2013.05.071, 2013.
- 905 [125] Meng, F., Qin, M., Tang, K., Duan, J., Fang, W., Liang, S., Ye, K., Xie, P., Sun, Y., Xie, C., Ye, C., Fu, P., Liu,
- 906 J., and Liu, W.: High-resolution vertical distribution and sources of HONO and NO<sub>2</sub> in the nocturnal boundary layer 907 in urban Beijing, China, Atmos. Chem. Phys., 20, 5071–5092, doi: 10.5194/acp-20-5071-2020, 2020.
- 908 [126] Zhang, W., Tong, S., Jia, C., Wang, L., Liu, B., Tang, G., Ji, D., Hu, B., Liu, Z., Li, W., Wang, Z., Liu, Y.,
- 909 Wang, Y., and Ge, M.: Different HONO sources for three layer at the urban area of Beijing, Environ. Sci. Technol., 910 54, 12870-12880, doi:10.1021/acs.est.0c02146, 2020.
- 911 [127] Fang, X., Li, T., Ban, C., Wu, Z., Li, J., Li, F., Cen, Y., and Tian, B.: A mobile differential absorption lidar for
- 912 simultaneous observations of tropospheric and stratospheric ozone over Tibet, Opt. Express, 27(4), 4126-4139, 913 doi:10.1364/OE.27.004126, 2019.
- 914 [128] Yu, J., Meng, L., Chen, Y., Zhang, H., and Liu, J.: Ozone profiles, precursors, and vertical distribution in urban 915 Lhasa, Tibetan Plateau, Remote Sens., 14(11), doi:10.3390/rs14112533, 2022.
- 916 [129] Zhang, J., Xia, X., and Wu, X.: First in situ UV profile across the UTLS accompanied by ozone measurement 917 over the Tibetan Plateau, J. Environ. Sci., 98, 71-76, doi:10.1016/j.jes.2020.05.020.
- 918 [130] Fisher, F. N.: Extinction of UV-visible radiation in wet midlatitude (maritime) snow: Implications for increased
- 919 NOx emission, J. Geophys. Res., 110, D21301, doi:10.1029/2005JD005963, 2005.
- 920 [131] Lin, W., Wang, F., Ye, C., Zhu, T.: Observation of strong NO<sub>x</sub> release over Qiyi Glacier, China. The 921 Cryosphere, doi.org/10.5194/tc-2021-32, 2021.

- [132] Ji, X., Liu, C., Wang, Y., Hu, Q., Lin, H., Zhao, F., Xing, C., Tang, G., Zhang, J., Wagner, T.: Ozone profiles
   without blind area retrieved from MAX-DOAS measurements and comprehensive validation with multi-platform
- 924 observations. Remote Sens. Environ., 284, 113339, doi.org/10.1016/j.res.2022.113339, 2023.
  - [133] Lin, H., Liu, C., Xing, C., Hu, Q., Hong, Q., Liu, H., Li, Q., Tan, W., Ji, X., Wang, Z., Liu, J.: Validation of
    water vapor vertical distributions retrieved from MAX-DOAS over Beijing, China. Remote Sens., 12, 3193,
    doi.org/10.3390/rs12193193, 2020.
  - [134] Xing, C., Xu, S., Song, Y., Liu, C., Liu, Y., Lu, K., Tan, W., Zhang, C., Hu, Q., Wang, S., Wu, H., Lin, H.: A new insight into the vertical differences in NO<sub>2</sub> heterogeneous reaction to produce HONO over inland and marginal seas. Atmos. Chem. Phys., 23, 5815-5834, doi.org/10.5194/acp-23-5815-2023, 2023.
  - [135] Rodgers, C. D.: Inverse methods for atmospheric sounding: theory and practice. Singapore-New
     Jersey-London-Hong: World Scientific Publishing; 2000.
  - [136] Wagner, T., Dix, B., FriedeBurg, C. V., Frieß, U., Sanghavi, S., Sinreich, R., Platt, U.: MAX-DOAS O<sub>4</sub>
     measurements: A new technique to derive information on atmospheric aerosols-Principles and information content. J.
  - 935 Geophys. Res.: Atmos., 109, D22205, doi.org/10.1029/2004jd004904, 2004.
  - [137] Serdyuchenko, A., Gorshelev, V., Weber, M., Chehade, W., Burrows, J. P.: High spectral resolution ozone
    absorption cross-sections-Part 2: Temperature dependence. Atmos. Meas. Tech., 7, 625-636,
    doi:10.5194/amt-7-625-2014, 2014.
  - [138] Wang, Y., Lampel, J., Xie, P., Beirle, S., Li, A., Wu, D., Wagner, T.: Ground-based MAX-DOAS observations
    of tropospheric aerosols, NO<sub>2</sub>, SO<sub>2</sub> and HCHO in Wuxi, China, from 2011 to 2014. Atmos. Chem. Phys., 17,
    2189-2215, doi.org/10.5194/acp-17-2189-2017, 2017.
  - 942 [139] Wang, Y., Apituley, A., Bais, A., Beirle, S., Benavent, N., Borovski, A., Bruchkouski, I., Chan, K. L., Donner,
  - 943 S., Drosoglou, T., Finkenzeller, H., Friedrich, M. M., Frieß, U., Garcia-Nieto, D., Gómez-Martín, L., Hendrick, F.,
  - Hilboll, A., Jin, J., Johnston, P., Koenig, T. K., Kreher, K., Kumar, V., Kyuberis, A., Lampel, J., Liu, C., Liu, H., Ma,
    J., Polyansky, O. L., Postylyakov, O., Querel, R., Saiz-Lopez, A., Schmitt, S., Tian, X., Tirpitz, J. L., Van Roozendeal,
  - M., Volkamer, R., Wang, Z., Xie, P., Xing, C., Xu, J., Yela, M., Zhang, C., Wagner, T.: Inter-comparison of
    MAX-DOAS measurements of tropospheric HONO slant column densities and vertical profiles during the CINDI-2
    campaign. Atmos. Meas. Tech., 13, 5087–5116, doi.org/10.5194/amt-13-5087-2020, 2020.
  - [140] Thalman, R., Volkamer, R.: Temperature dependent absorption cross-sections of O<sub>2</sub>-O<sub>2</sub> collision pairs between
    340 and 630 nm and at atmospherically relevant pressure. Phys. Chem. Chem. Phys., 15, 15371-15381,
    doi:10.1039/C3CP50968K, 2013.
  - 952 [141] Vandaele, A. C., Hermans, C., Simon, P. C., Carleer, M., Colin, R., Fally, S., Merienne, M. F., Jenouvrier, A., 953 Coquart, D.: Measurements of the NO<sub>2</sub> absorption cross-section from 42000 cm<sup>-1</sup> to 10000 cm<sup>-1</sup> (238–1000nm) at 954 220K and 294K. J. Quant. Spectrosc. Ra., 59, 171-184, doi:10.1016/S0022-4073(97)00168-4, 1998.
  - 220K and 294K. J. Quant. Spectrosc. Ra., 59, 171-184, doi:10.1016/S0022-4073(97)00168-4, 1998.
    [142] Stutz, J., Kim, E. S., Platt, U., Bruno, P., Perrino, C., Febo, A.: UV-visible absorption cross sections of nitrous
- 956 acid. J. Geophys. Res.: Atmos., 105, 14585-14592, doi:10.1029/2000JD900003, 2000.
- [143] Aliwell, S. R., Van Roozendael, M., Johnston, P. V., Richter, A., Wagner, T., Arlander, D. W., Burrows, J. P.,
  Fish, D. J., Jones, R. L., Tørnkvist, K. K., Lambert, J. C., Pfeilsticker, K., and Pundt, I.: Analysis for BrO in
  zenith-sky spectra: an intercomparison exercise for analysis improvement, J. Geophys. Res., 107, ACH 10-1–ACH
  10-20, https://doi.org/10.1029/2001JD000329, 2002.
- 961 [144] Vandaele, A. C., Hermans, C., Simon, P. C., Carleer, M., Colin, R., Fally, S., Mérienne, M. F., Jenouvrier, A.,
- and Coquart, B.: Measurements of the NO<sub>2</sub> absorption cross section from  $42000 \text{ cm}^{-1}$  to  $10000 \text{ cm}^{-1}$  (238–1000nm) at 220 K and 294 K, J. Quant. Spectrosc. Ra., 59, 171–184, doi:10.1016/S0022-4073(97)00168, 1998.
- 964 [145] Meller, R. and Moortgat, G. K.: Temperature dependence of the absorption cross sections of formaldehyde
  965 between 223 and 323 K in the wavelength range 225–375nm, J. Geophys. Res., 105, 7089–7101,
  966 doi:10.1029/1999JD901074, 2000.
- 967 [146] Volkamer, R., Spietz, P., Burrows, J., Platt, U.: High-resolution absorption cross-section of glyoxal in the
  968 UV-vis and IR spectral ranges, J. Photochem. Photobiol. A Chem., 172, 35–46,
- 969 doi:10.1016/j.jphotochem.2004.11.011, 2005.
- 970 [147] Rothman, L. S., Gordon, I. E., Barbe, A., Benner, D. C., Bernath, P. E., Birk, M., Boudon, V., Brown, L. R.,
- Campargue, A., Champion, J. P., Chance, K., Coudert, L. H., Dana, V., Devi, V. M., Fally, S., Flaud, J. M., Gamache,
- R. R., Goldman, A., Jacquemart, D., Kleiner, I., Lacome, N., Lafferty, W. J., Mandin, J. Y., Massie, S. T.,
  Mikhailenko, S. N., Miller, C. E., Moazzen-Ahmadi, N., Naumenko, O. V., Nikitin, A. V., Orphal, J., Perevalov, V. I.,
- Mikhailenko, S. N., Miller, C. E., Moazzen-Ahmadi, N., Naumenko, O. V., Nikitin, A. V., Orphal, J., Perevalov, V. I.,
  Perrin, A., Predoi-Cross, A., Rinsland, C. P., Rotger, M., Simeckova, M., Smith, M. A. H., Sung, K., Tashkun, S. A.,
- 975 Tennyson, J., Toth, R. A., Vandaele, A. C., Vander Auwera, J.: The HITRAN 2008 molecular spectroscopic database,
- 976 J. Quant. Spectrosc. Radiat. Transf., 110, 533–572, 2009.
- 977 [148] Fleischmann, O. C., Hartmann, M., Burrows, J. P., and Orphal, J.: New ultraviolet absorption cross-sections of
- BrO at atmospheric temperatures measured by time-windowing Fourier transform spectroscopy, J. Photoch. Photobio.
   A, 168, 117–132, 2004.
- 980 [149] Ward Jr, J. H.: Hierarchical grouping to optimize an objective function, J. Am. Stat. Assoc., 58, 236-244, 1963.
- [150] Wang, Y., Zhang, X., Arimoto, R.: The contribution from distant dust sources to the atmospheric particulate
   matter loading at XiAn, China during spring, Sci. Total Environ., 368, 875-883, 2006.

- 983 [151] Cheng, S., Pu, G., Ma, J., Hong, H., Du, J., Yudron, T., Wagner, T.: Retrieval of tropospheric NO<sub>2</sub> vertical column densities from ground-based MAX-DOAS measurements in Lhasa, a city on the Tibetan Plateau, Remote
- 985 Sens., 15, 4689, 2023a.

[152] Cheng, S., Ma, J., Zheng, A., Gu, M., Donner, S., Donner, S., Zhang, W., Du, J., Li, X., Liang, Z., Lv, J.,
Wagner, T.: Retrieval of O<sub>3</sub>, NO<sub>2</sub>, BrO and OCIO columns from ground-based zenith scattered light DOAS measurements in summer and autumn over the Northern Tibetan Plateau, Remote Sens., 13, 4242, 2021.

- 989 [153] Ma, J., Donner, S., Donner, S., Jin, J., Cheng, S., Guo, J., Zhang, Z., Wang, J., Liu, P., Zhang, G., Pukite, J.,
- 990 Lampel, J., Wagner, T.: MAX-DOAS measurements of NO<sub>2</sub>, SO<sub>2</sub>, HCHO, and BrO at the Mt. Waliguan WMO GAW
- global baseline station in the Tibetan Plateau, Atmos. Chem. Phys., 20, 6973-6990, 2020.
- 992 [154] Cheng, S., Cheng, X., Ma, J., Xu, X., Zhang, W., Lv, J., Bai, G., Chen, B., Ma, S., Ziegler, S., Donner, S.,
- Wagner, T.: Mobile MAX-DOAS observations of tropospheric NO<sub>2</sub> and HCHO during summer over the Three
- Rivers' Source region in China, Atmos. Chem. Phys., 23, 3655-3677, 2023b.
- 995 [155] Li, M., Mao, J., Chen, S., Bian, J., Bai, Z., Wang, X., Chen, W., Yu, P.: Significant contribution of lightning
- NO<sub>x</sub> to summertime surface O<sub>3</sub> on the Tibetan Plateau, Sci. Total Environ., 829, 154639, 2022.
   997
- 998