High-resolution physicochemical dataset of atmospheric aerosols over the Tibetan Plateau and its surroundings

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Abstract

Atmospheric aerosol in the Tibetan Plateau (TP) and its surroundings has attracted significant scientific interest in recent decades due to its notable impacts on regionally climatic and cryospheric changes, ecological and environmental securities, and the hydrological cycle. However, our understanding on the atmospheric aerosol in this remote region is highly limited by the scarcity of available dataset, owing to the extremely harsh natural conditions. This challenge has been mitigated in recent decades by establishing field observatories at typical sites within the TP and its surroundings. A continuous project initiated in 2015 aims to explore the properties and sources of atmospheric aerosols, as well as their regional differences, through multiple short-term intensive observations across this vast region utilizing a suite of high-time-resolution online instruments. This paper presents a systematic hourly scaled dataset of aerosol physicochemical and optical properties at eight sites across the TP and its surroundings derived from the project. It includes size-resolved chemical compositions of submicron aerosols, high-resolution mass spectra and sources of organic aerosols, size distributions of particle number concentrations, particle light scattering and absorption coefficients, particle light absorptions attributed to different carbonaceous substances including black carbon and brown carbon, and number concentrations of cloud condensation nuclei. In brief, atmospheric aerosols in these remote sites were all well-mixed and highly aged, reflecting their dominated regional transport sources. However, the southern TP region exhibited high contributions of carbonaceous organic aerosols, neutralized bulk submicron aerosols, and a relatively higher light absorption capacity, whereas in the northern TP region, secondary inorganic species were the main contributors to the overall acidic submicron aerosols. Beyond providing insights into the regional differences in aerosol sources and properties across the TP and its surroundings, the datasets will also benefit simulations of aerosol radiative forcing and evaluations of interactions among different Earth system components in numerical models in this region. The datasets are accessible through the National Cryosphere Desert Data Center, Chinese Academy of Sciences (https://doi.org/10.12072/ncdc.NIEER.db2200.2022; Xu, 2022).
1 Introduction

Tibetan Plateau (TP), with an average elevation exceeding 4,000 m above sea level (a.s.l.) and spanning a surface area of approximately 2.5 million square kilometers, stands as the highest plateau on the Earth. Its high-altitude mountain ranges, integral to one of the world’s most crucial cryospheric regions, have earned the TP the monikers “roof of the world”, “Third Pole”, and “Asian Water Tower” (Qiu, 2008; Yao et al., 2019). The TP and its surroundings play a pivotal role in influencing the global and regional climate systems, hydrological cycles, and cryospheric changes through its vast and complex topography and its function as a significant heat source (Duan and Wu, 2005; Yao et al., 2012; Chen et al., 2021). In recent decades, a major concern has focusing on the significant climatic warming and the rapid changes in the cryosphere of this region (Kang et al., 2010), which exhibits a warming trend that exceeding that of the Northern Hemisphere (0.34 vs. 0.29 °C/decade) (You et al., 2021; Zhou and Zhang, 2021).

Atmospheric aerosols, as one of the most complex and critical components in the atmosphere, significantly influence climatic warming and cryospheric changes in the TP regions. They exert crucial direct and indirect effects on atmospheric energy budget and the albedos of snow and ice surfaces, impacting Earth’s climate system (Xu et al., 2009; Kang et al., 2019b). Notably, light-absorbing carbonaceous aerosols (CAs) such as black carbon (BC) and brown carbon (BrC) directly absorb solar radiation, warming the atmosphere and contributing to positive forcing on Earth’s energy budget (Ramanathan et al., 2007; Kopacz et al., 2011). For instance, Li et al. (2018) demonstrated that BC causes significantly greater albedo reduction (~46%) and instantaneous radiative forcing (7–64 W m²) on aged snow surfaces of a TP glacier than mineral dust. Moreover, aerosol particles over the TP significantly affect ice cloud properties and cloud development through their semi-direct effects (Liu et al., 2019). Given the vast and remote nature of the TP, coupled with its complex topography, meteorology, and harsh environment, in-situ observation of atmospheric aerosols poses substantial challenges. Consequently, numerical model simulations based on reanalysis data have emerged as a predominant and crucial method in recent decades. Notable studies include Lau et al. (2006), who assessed the impact of atmospheric aerosols on the Asian summer monsoon intensification using the NASA finite-volume general circulation model; Kopacz et al. (2011), who explored the origin and radiative forcing...
of BC in the TP and Himalayas with the GEOS-Chem global chemical transport model; and Liu et al. (2015), who examined the transport of summer dust and anthropogenic aerosols using a three-dimensional aerosol transport–radiation model with satellite data inputs. Despite the significant insights gained from these simulations, *in-situ* observations of atmospheric aerosols in the TP are increasingly recognized as critical for evaluating and enhancing model accuracy in this remote area. The absence of *in-situ* aerosol data to refine models introduces considerable uncertainty into the results. Furthermore, while model simulations primarily focus on spatial distribution across broad regions, they often overlook temporal variations or the inherent evolution mechanisms at high temporal resolution, which *in-situ* observations could illuminate.

Recent advancements in observational techniques and instrumentation have enabled numerous *in-situ* measurement within the TP and its surroundings, aiming to delineate the physical, chemical, and optical properties of aerosols, along with their potential sources, transport pathways, and regional distributions. A detailed compilation of direct ambient aerosol measurements in the TP, employing a variety of observational methods and instruments, is presented in Table S1 in the supplementary material. Notably, *off-line* atmospheric filter sampling has emerged as a key *in-situ* aerosol collection method in the TP, favored for its low-cost and feasibility under the region’s harsh conditions and logistical constraints. This method has effectively captured the composition, size, light absorption properties, sources, and variations of ambient aerosols, including diverse CAs components such as BC, BrC, organic carbon (OC), water-soluble OC (WSOC), humic-like substances (HULS), and polycyclic aromatic hydrocarbons (PAHs) in the remote TP region (Cao et al., 2009; Zhao et al., 2013; Xu et al., 2014a; Zhang et al., 2014; Cong et al., 2015; Wan et al., 2015; Xu et al., 2015; Kang et al., 2016; Li et al., 2016b; Xu et al., 2020). Furthermore, *off-line* filter sampling has been instrumental in mapping the regional aerosol distribution across the TP, facilitated by its simplicity and the ability to conduct simultaneous observations at multiple locations (Li et al., 2016a; Chen et al., 2019; Kang et al., 2022). Despite these advancements, *off-line* filter sampling studies in the TP’s remote regions remain insufficient for current needs. The gathered data is generally fragmented and unsystematic, characterized by low temporal resolution, limited aerosol property parameters, and sparse data points. Typically, these studies have been localized to specific sites, utilizing single instruments with temporal resolutions ranging from days to weeks over brief periods. Such low temporal
resolution limits the accurate understanding of aerosols’ temporal evolution and underlying mechanisms, especially during rapid, short-term events. Additionally, integrating and comparing data across different research groups is challenging due to variations in research focuses, measured parameters, sampling methodologies, laboratory filter processing, and data analysis techniques. Moreover, despite the relative ease of off-line sampling, extensive areas of the TP still lack such observational efforts. To date, comprehensive research focusing on multiple aerosol parameters through real-time online consecutive measurements (with high temporal resolutions from minute to hour scales) at multiple sites remains a rarity in the TP.

Research on atmospheric aerosols in China has achieved significant advancements over the past decade. The Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) has become a widely used online instrument, enabling numerous studies to characterize the real-time, size-resolved chemical compositions and sources of submicron aerosols (Li et al., 2017; Zhou et al., 2020). However, its application in the TP has been limited, a situation attributed to the instrument’s demanding operational requirements and the area’s challenging observation conditions. Since 2015, our research group has initiated a continuous and systematic observation project, aiming to investigate the regional differences on aerosol sources and properties across the TP. This effort involves annual deployments of the HR-ToF-AMS alongside other high-resolution, real-time online instruments at various sites. Remarkably, our dataset represents the first and only collection that extensively covers a wide range of aerosol parameters (including physical, chemical, and optical properties and their diverse sources) across multiple geographic environments (e.g., urban, remote, high-altitude mountain, grassland, and subtropical forest) within the TP and its surroundings. This is achieved through real-time and high-time-resolution online observations, providing a comprehensive resource for understanding regional aerosol variations and serving as essential input for future aerosol radiative forcing simulations and Earth system interaction assessments. The structure of this paper is as follows: Sections 2 and 3 describe the observation sites, instrumental deployments, and data processing. Section 4 presents the high-time-resolution aerosol data, encompassing physical, chemical, and optical properties as well as source information. The limitations and the uniqueness of our dataset are discussed in Section 5.

2 Observation site descriptions
Between 2015 and 2022, intensive observations of atmospheric aerosol chemistry were carried out at eight sites across the TP and its surroundings. These sites comprise seven remote sites (QOMS, Motuo, NamCo, Ngari, Waliguan, LHG, and Bayanbululak) and one urban site (Lhasa), the latter serving as a contrast for comparative analysis. Figure 1 illustrates the geographical locations of these sites along with photographs during each observation. Table 1 details the specifics of each site including the sampling periods and the instruments deployed during each field campaign. The subsequent sections provide brief descriptions these sites, arranged geographically from the southern to the northern parts of the TP.

2.1 QOMS

The Qomolangma Station for Atmospheric and Environmental Observation and Research, Chinese Academy of Sciences (86.56°E, 28.21°N; 4276 m a.s.l.; abbreviated as QOMS in this study, with similar abbreviations for other sites hereafter), is situated in the Rongbuk valley basin on the northern slope of Mt. Everest. The climate in the northern slope of Mt. Everest has obvious seasonal variation, heavily influenced by the Indian monsoon system (Bonasoni et al., 2010; Cong et al., 2015). During the pre-monsoon season (typically March to May), dominant westerlies facilitates the long-range transport of atmospheric pollutants from South Asia, making QOMS an ideal high-altitude observatory on the south edge of the TP for examining transboundary pollutant transport into the plateau’s interior. In the summer monsoon season (June–August), prevailing southerly winds bring warm and wet airflow from the Indian Ocean, leading to increased humidity and precipitation in the plateau.

2.2 Motuo

Motuo County, located in the lower reaches of the Yarlung Tsangpo River on the southern slopes of the eastern Himalayas and Gangrigab Mountains in the southeast edge of the TP. The county, sited halfway up a mountain, enjoys a subtropical humid climate characterized by relatively high temperatures and abundant rainfall. With a small population of ~15,000, Motuo County remains one of the TP’s most pristine regions. The sampling site in Motuo (29.30°N, 95.32°E; 1305 m a.s.l.) was located atop a hill overlooking the Yarlung Tsangpo Grand Canyon. This vantage point makes it an ideal location for directly monitoring the transboundary transport of atmospheric pollutants and moisture from Southeast Asia and the Indian Ocean into the TP.
2.3 NamCo

The Nam Co Station for Multisphere Observation and Research, Chinese Academy of Sciences (NamCo; 90.95°E, 30.77°N; 4730 m a.s.l.) is a high-altitude observatory in the south-central part of the TP. Situated at the southeast shore of Nam Co Lake, the station is surrounded by a pristine and isolated area. The region experiences a typical semi-arid plateau monsoon climate, characterized by increased precipitation during the summer monsoon season. NamCo is a pivotal inland site within the TP, predominantly influenced by air masses from the south and west.

2.4 Ngari

The Ngari Station for Desert Environment Observation and Research, Chinese Academy of Science (Ngari; 79.70°E, 33.39°N; 4270 m a.s.l.) is located in the Rutog County within the Ngari Prefecture of the Tibet Autonomous Region, China. This area is at the southwestern edge of the TP and is characterized by its semi-arid climate, sparse vegetation, and intense solar radiation. As a key member of “high-cold region observation and research network for land surface processes & environment of China”, the Ngari station plays a crucial role in monitoring climate, hydrological, atmospheric, and ecological environmental changes in the TP’s western territories. Additionally, it contributes to understanding the interactions between the Indian monsoon system and the westerlies.

2.5 Waliguan

The Waliguan Baseline Observatory (Waliguan; 100.9°E, 36.28°N; 3816 m a.s.l.) is one of the twenty-nine baseline stations of Global Atmosphere Watch (GAW) under the World Meteorological Organization (WMO). Situated at the top of Mt. Waliguan, which rises approximately 600 m, the observatory is located in a pristine region with minimal human activity impact. Waliguan is represented as a key observatory on the northeastern edge of the TP, predominantly influenced by air masses from the northeast during the summer. This location is strategic for studying the transport and impact of air pollutants from industrial regions in northwestern China to the TP in this study.

2.6 LHG

The Qilian Observation and Research Station of Cryosphere and Ecologic Environment, Chinese Academy of Sciences (LHG; 39.50°N, 96.51°E; 4180 m a.s.l.), is located approximately 1km from the terminus of Laohugou Glacier No.12. This glacier is
among the largest mountain glacier in the northern slope of the western Qilian Mountains. LHG serves as another notable station in the northeastern TP, distinctly remote from human settlements. The climate in this region is predominantly arid and continental, influenced by the East-Asian monsoon in summer and the Westerlies in winter. A pronounced mountain-valley breeze during summer facilitates the upward transport of air masses from lower altitudes, making LHG an ideal site for background air mass sampling and for investigating the transport and potential impacts of air pollutants from surrounding areas.

2.7 Bayanbulak
The Bayanbulak National Basic Meteorological Station (Bayanbulak; 84.35° N, 42.83° E; 2454 m a.s.l.) is located in the Bayanbulak grassland, northwest of Hejing County in the Xinjiang Uygur Autonomous Region, China. Bayanbulak lies within an intermontane basin in the central Tianshan Mountains, surrounded by numerous snow mountains with altitudes more than 3000 m. The climate in Bayanbulak grassland is characterized by a typical temperate continental mountain climate, with an annual average precipitation ranging between 200 to 300 mm. Bayanbulak town experiences minimal human activities and traffic, maintaining its pristine environment.

2.8 Lhasa
Lhasa (29.65°N, 91.03°E; 3650 m a.s.l.) is the capital of the Tibet Autonomous Region, China, and located in the south-central part of the TP. The city lies in a broad river valley, surrounded by mountains that reach up to 5500 m, with the Lhasa River passing through the city from west to east. Our observation site is located in Binhe Park, adjacent to the Lhasa River. Notably, the Norbulingka scenic area, one of the main activity centers for local Tibetans celebrating their religious festivals such as the Sho Dun festival, is located ~1 km to the northwest of the sampling site. Moreover, the Potala Palace, the center of Tibetan Buddhism, is ~1.8 km to the northeast. Given Lhasa’s unique energy structure and the distinct living habits of the residents, comparative observations are carried out at this urban site. These studies aim to investigate the primary aerosol properties and sources, particularly from various residential combustion activities.

3 Online sampling, instrumental setup, and data processing
3.1 Online real-time aerosol sampling over the TP
Atmospheric aerosol observations were conducted at each site using a suite of real-time, high-resolution instruments. This instruments typically included a HR-ToF-AMS (Aerodyne Research Inc., Billerica, MA, USA) to determine the chemical composition (organic aerosol (OA), nitrate, sulfate, ammonium, and chloride) of non-refractory submicron aerosol (PM$_1$); A scanning mobility particle sizer (SMPS, model 3936, TSI Inc., Shoreview, MN, USA) was used to measure the size distribution and number concentration of submicron particles; A photoacoustic extinctionometer (PAX, DMT Inc., Boulder, CO, USA) was used for obtaining particle light absorption, scattering, and extinction coefficients (B$_{abs}$, B$_{scat}$, and B$_{ext}$) along with single scattering albedo (SSA) at 405 nm, as well as BC mass concentration; An Aethalometer (model AE33/AE31, Magee Scientific Corp., Berkeley, CA, USA) was used for acquiring the $B_{abs}$ across seven wavelengths (370–950 nm); A cloud condensation nuclei (CCN) counter (model CCN-100, DMT Inc., Boulder, CO, USA) measured CCN number concentrations at various water vapor supersaturations (SS). The measurement uncertainties for each instrument are difficult to quantify based on data from a single instrument. The uncertainty showing below are referred from other studies: <30% for HR-ToF-AMS (Jimenez et al., 2016) and SMPS, <40% for Aethalometer (Backman et al., 2017), <10% for PAX (Selimovic et al., 2018), and <25% for CCNC (Rose et al., 2008). Details on the specific instruments deployed and the sampling periods for each observation campaign are summarized in Table 1.

The HR-ToF-AMS was the cornerstone instrument for atmospheric aerosol chemistry observations across all field campaigns. The SMPS was deployed at QOMS, Motuo, LHG, and Lhasa, whereas the PAX was employed at QOMS, Motuo, Ngari, Waliguan, and Lhasa. The Aethalometer was utilized at QOMS, NamCo, and Waliguan, with the CCN-100 operational at Motuo, Waliguan, and LHG. Field observations in the southern, western, and central regions of the TP were predominantly conducted during the pre-monsoon season, aimed to study the transboundary transport of pollutants from South Asia into the TP, under the influence of Westerlies and the Indian monsoon. For instance, observations took place from 12 April to 12 May at QOMS, Motuo from 26 March to 22 May, NamCo from 31 May to 1 July, and Ngari from 1 Jun to 5 July. On the other hand, measurements in the remote regions of the northern TP and its surroundings were carried out during the summer to track aerosol transport from surrounding polluted areas, considering the effects of the Westerlies or the intensified East Asian monsoon.
Specifically, observations occurred from 1 to 31 July at Waliguan, from 4 to 29 August at LHG, and from 29 August to 26 September at Bayanbulak. The observation at Lhasa was conducted from 31 August to 26 September, focusing on capturing the peak atmospheric oxidation capacity during the summer.

3.2 Instrumental setup

Despite the variations in instrumentation across different observation campaigns, the core setup for sampling was largely consistent. Figure 1b illustrates the standard sampling configuration for each campaign typically involved housing all instruments within an air-conditioned trailer or room. The inlets were induced to the instruments from the roof with a cyclone (model URG-2000-30EH, URG Corp., Chapel Hill, NC, USA) in the front of the inlet to eliminate particles with an aerodynamic diameter ($D_{va}$) exceeding 2.5 µm. These fine particles were then passed through a Nafion dryer via 1/2-inch stainless steel tubing to dehumidify the airflows before being directed into the instruments for real-time analysis. For detailed information on the setup and methodology of the instruments, it can be found in our previous publications (Xu et al., 2018; Zhang et al., 2018; Zhang et al., 2019; Zhao et al., 2022).

3.3 Instrumental operation and data processing

The measurement principles, operation procedures, calibration methods, and data analysis for these instruments used in this study are thoroughly detailed in Texts S1–S4 of supplementary material. Here, we highlight only key descriptions and critical setting as follows: (1) The HR-ToF-AMS was operated at V-mode during most of the eight field campaigns to accommodate the relatively low signal-to-noise ratios due to low aerosol mass loading in the TP regions. (2) Particle size observations were not conducted during the NamCo and LHG campaigns due to a chopper malfunction in the HR-ToF-AMS. (3) Different relative ionization efficiency (RIE) values were used for ammonium and sulfate according to the ionization efficiency calibrations of HR-ToF-AMS in different campaigns. (4) Different size parameters were achieved according to the particle sizing calibrations in different campaigns. (5) Elemental ratios of OA such as oxygen-to-carbon (O/C), hydrogen-to-carbon (H/C), organic matter-to-organic carbon (OM/OC), and nitrogen-to-carbon (N/C), were determined using the improved method (Canagaratna et al., 2015). (6) A default collection efficiency (CE) value of 0.5 were employed to the HR-ToF-AMS measurements during the QOMS, NamCo, Ngari,
Waliguan, and Lhasa campaigns in consideration of their overall neutralized bulk submicron aerosols, whereas a composition-dependent CE (Middlebrook et al., 2012) value was adopted at Motuo, LHG, and Bayanbulak, where aerosols were slightly acidic. 

(7) Source apportionment of OA during all observations were performed by the positive matrix factorization (PMF) analysis. The details of the PMF solution determination for each site are not presented here but can be referenced in our previous publication for select campaigns (Xu et al., 2018; Zhang et al., 2018; 2019). (8) Only the chemical compositions of non-refractory PM$_1$ are reported for the Bayanbulak campaign due to the absence of BC observations. (9) The sample and sheath flow rates of SMPS were set at 0.3 and 3.0 L min$^{-1}$, respectively, at both QOMS and Lhasa, covering a particle size range between 14.6 and 661.2 nm in mobility diameter ($D_m$), whereas 0.5 and 5.0 L min$^{-1}$ at LHG and Motuo sites with a range of 10.9–495.8 nm in $D_m$. (10) Aethalometer measurements were corrected for filter-based loading and multiple scattering effects. A traditional absorption Ångström exponents (AAE) method (Zhang et al., 2021) was adopted to apportion total $B_{abs}$ into two parts of BC and BrC ($B_{abs,BC}$ and $B_{abs,BrC}$). (11) CCN number concentrations were measured at five different SS values of 0.2%, 0.4%, 0.6%, 0.8%, and 1.0% at a 30-min cycle during the Motuo, Waliguan, and LHG campaigns.

4 Aerosol properties, sources, and radiative forcing over the TP

4.1 Mass loading and chemical composition of submicron aerosols

Figure S1 presents the temporal variations of PM$_1$ chemical species (OA, nitrate, sulfate, ammonium, chloride, and BC) observed across the eight observations in the TP and its surroundings. The mass concentrations of PM$_1$ and its components exhibited distinct variations, with a few periods of high mass loading observed throughout each campaign’s sampling period. Despite variations in sampling years (2015–2022), seasons (March–September), and altitudes (1350–4730 m a.s.l.) across these sites, the distinct PM$_1$ mass concentrations and chemical compositions clearly illustrate the regional difference among these sites. On average, total PM$_1$ mass concentrations across the eight campaigns ranged from 1.9 to 9.1 µg m$^{-3}$ (Fig. 2 and Table 2). The highest mass concentration was observed at Waliguan, driven by the transport of anthropogenic aerosols and gaseous pollutants from urban centers in northwestern China. In contrast, the lowest values were observed at NamCo and Bayanbulak, reflecting their background and pristine environmental conditions. The average PM$_1$
mass level across the TP and its surroundings was comparable to those observed at other high-altitude, coastal, forest, and remote background sites globally (0.46–15.1 µg m$^{-3}$; Table 3), yet remains significantly lower than those observed at densely urban (34.4–71.5 µg m$^{-3}$) and suburban (21.4–44.9 µg m$^{-3}$) areas in other parts of China (Li et al., 2017). This suggests the predominantly clean atmospheric conditions in the remote and high-altitude regions of the TP.

The chemical compositions of PM$_1$ also exhibited significant regional differences across the TP (Fig. 2), highlighting varied aerosol sources in different areas of the TP. At five sites (QOMS, Motuo, NamCo, Ngari, and Lhasa) located in the southern, western, or central TP, OA and BC together accounted for as high as 64.9–85.7% of the total PM$_1$ mass (Table 2). This high contribution was largely attributed to the frequent transport of biomass-burning emissions from South and Southeast Asia to the TP during the pre-monsoon season (Bonasoni et al., 2010; Cong et al., 2015; Zhang et al., 2018), along with significant local biomass burning from religious activities in Lhasa (Cui et al., 2018; Zhao et al., 2022). In contrast, at the three northern sites (Waliguan, LHG, and Bayanbulak), inorganic species (sulfate, nitrate, and ammonium; referred to as SNA) accounted for more than 60% of the total PM$_1$. Sulfate was the most significant component of SNA (38.1–46.0%), aligning with observations from another high-altitude site in the northeastern TP (Menyuan; 28%) and various rural and remote sites (19–64%) in East Asia (Du et al., 2015). The pronounced SNA contributions, particularly sulfate, in the northern TP and its surroundings, were mainly related to the regional transport of anthropogenic aerosols and gaseous precursor from nearby urban areas as well as important in-cloud aqueous reactions during the transportation to the mountains (Zhang et al., 2019).

4.2 Bulk acidity, size distribution, and diurnal variation of submicron aerosols

Particle phase acidity significantly influences aerosol physicochemical properties, affecting hygroscopicity, toxicity, and heterogeneous reactions. The bulk acidity of submicron aerosols was evaluated at each site following the method in Zhang et al. (2007b) and Schueneman et al. (2021) using AMS measurements. A detailed description of this method can be found in Text S5 of the supplementary material or our previous publications (Zhang et al., 2018; Zhang et al., 2019). Notably, we observed clear regional variations between the southern and northern TP, largely due to differences in aerosol sources and composition (Fig. 3). Linear regression slopes at
QOMS, NamCo, Ngari, and Lhasa (located in the southern, western, or central TP) were fitted to be 1.2, 1.11, 0.98, and 1.18, respectively, indicating the submicron aerosols at these sites were generally neutralized, occasionally showing an excess of ammonium. The result is consistent with previous findings on high ammonia availability from agriculture emissions in the South Asia (Van Damme et al., 2015). Moreover, as reported in our previous publications, atmospheric aerosols at QOMS and NamCo were significantly influenced by biomass-burning emissions from South and Southeast Asia during the pre-monsoon season (Xu et al., 2018; Zhang et al., 2018), and Lhasa experienced intense biomass fuel burning during frequent religious festivals (Zhao et al., 2022). In contrast, submicron particles at the remaining four sites, particularly LHG and Bayanbulak in the north, were overall acidic, with regression slopes ranging from 0.73 to 0.86, where sulfate was a major PM$_1$ component (46.0% and 41.6%). Similar findings of acidic submicron aerosol particles have also been observed at Menyuan and LHG in the northern TP in previous studies (Du et al., 2015; Xu et al., 2015), mainly related to the transport of enriched SNA species or their gaseous precursors from the industrial areas in northwestern China.

The size distributions of non-refractory PM$_1$ chemical species, obtained from HR-ToF-AMS measurement, provide valuable insights into aerosol sources, oxidation degrees, mixing states, formation, transformation, and growth mechanisms as well as their impacts on CCN activity. Typically, size distributions for SNA species and oxidized OA peaked in the accumulation mode (~400–600 nm in $D_{va}$), as a result of secondary formation processes. In contrast, fresh organics from primary emission sources exhibit smaller size (Zhang et al., 2005b; Aiken et al., 2009). In this study, we focus on organics and the combined three SNA species to highlight regional variations in size distributions across the TP. As shown in Figs. 4a and S2 and Table 2, the peak diameters of OA and SNA size distributions varied significantly, from 584.4 and 634.5 nm at Ngari to a smaller size of 228.1 and 250.0 nm at Lhasa, respectively. This variation suggests the distinctly different sources and aging processes of atmospheric aerosols across the TP, particularly between those high-altitude remote sites and urban sites. For instance, bulk PM$_1$ at QOMS was reported to be internally well-mixed and aged, attributed to long-range transport from biomass-burning emissions in South Asia (Zhang et al., 2018), whereas local primary sources, including cooking, traffic exhausts, and biomass burning, totally accounted for more than 60% of the total OA in urban Lhasa (Zhao et
al., 2022). The crucial influence of aerosol sources on size distributions is further supported by the correlation between the mode size and O/C ratios of OA ($R^2 = 0.74$) (Fig. 4a).

The diurnal variations of PM$_1$ chemical compositions are typically influenced by multiple factors, including meteorological conditions (e.g., planetary boundary layer (PBL) height, wind direction and speed, temperature, relative humidity), various primary emission sources (e.g., vehicle exhausts during traffic rush hours, cooking emissions, and coal combustion emissions for heating), and distinct formation mechanisms (e.g., daytime photochemical oxidation, nighttime heterogeneous reactions, and gas-particle partitioning of secondary species). A comprehensive understanding of these diurnal variations is crucial for exploring the dynamic evolution of aerosol compositions and identifying the primary drivers (source, meteorology, or secondary formation) behind the variations in different chemical species.

Distinct diurnal patterns in the total PM$_1$ mass concentrations were observed across different field campaigns (Fig. 4b). At remote sites of QOMS, LHG, NamCo, and Waliguan, located in valleys or atop mountains, variations were largely governed by mountain-valley wind circulation and change in PBL height. For instance, QOMS exhibited a distinct diurnal pattern with continuously decreasing concentrations during the daytime, but relatively higher concentrations at night. The minimum occurred at around ~15:00 in this valley site, likely due to strong afternoon glacier winds and a higher PBL (Zhang et al., 2018). Conversely, LHG and NamCo experienced lower PM$_1$ concentrations from night to early morning, with increase in the afternoon, attributed at LHG to up-slope wind transport and at NamCo to aerosols descending from higher layers and enhanced afternoon transport from the west (Xu et al., 2018). A complex diurnal pattern of PM$_1$ was observed at Waliguan, influenced by diffusion conditions, wind directions, and air mass sources, including afternoon air masses from the northeast, which likely carried industrial pollutants (Zhang et al., 2019). At Motuo, the diurnal pattern of PM$_1$ was relatively stable, with two weak peaks linked to local combustion activities in the late morning and evening. Ngari exhibited relatively higher nighttime loadings and lower daytime loadings, mainly due to the variations of PBL height. Bayanbulak, on the other hand, had relatively low and stable PM$_1$ mass throughout the entire day due to its background location. In contrast, urban Lhasa displayed two pronounced peaks correlating with primary emissions during morning and evening rush.
hours (Zhao et al., 2022). Although the diurnal pattern of PM$_1$ were mainly shaped by
mountain-valley winds and PBL height in those remote sites, and primary emissions in
the urban site in this study, secondary formation processes, including photochemical
oxidation and aqueous-phase reactions were also played a key role in the formation of
inorganic and organic aerosol species. This could be evidenced by the afternoon peaks
of oxygenated OA (OOA) components observed across almost all the sites, which were
commonly formed by photo-chemical processes (Xu et al., 2018; Zhang et al., 2018;
Zhang et al., 2019; Zhao et al., 2022).

4.3 High-resolution mass spectrum and elemental ratios of organic aerosol

The high-resolution mass spectrum (HRMS) and elemental ratios of OA were
determined to identify the possible sources, formation and evolution mechanisms, and
oxidation states at each site. A direct comparison of the average O/C ratios from the
eight field campaigns was presented in Fig. 5a. It is apparent that the O/C ratios at the
remote sites of QOMS, Motuo, NamCo, Ngari, Waliguan, and LHG typically reached
or exceeded 1.0, indicating highly oxidized OA. In contrast, Bayanbulak exhibited a
lower O/C ratio of 0.69, and the urban site of Lhasa showed an even lower ratio of 0.44.
These variations in O/C ratios across sites primarily reflect differences in OA sources
and aging processes. Remote sites in the TP were generally received well-mixed and
aged OA due to long-range transport (Xu et al., 2018; Zhang et al., 2018; Zhang et al.,
2019). Meanwhile, local emissions from activities like cooking, traffic, and biomass
burning significantly contribute to the OA in urban Lhasa, resulting in a comparatively
low O/C ratio (Zhao et al., 2022). This pattern of higher O/C ratios at remote sites and
lower ratios at urban sites were observed in previous findings across China, such as O/C
ratios of 0.98, 1.11, and 1.16 observed at Mt. Wuzhi (Zhu et al., 2016), Mt. Yulong
(Zheng et al., 2017), and LHG (Xu et al., 2015), respectively, versus urban sites where
O/C ratios typically fell below 0.5 at most urban sites (Zhou et al., 2020). The Van
Krevelen diagram, which plots H/C versus O/C ratios to illustrate changes in OA
elemental composition due to atmospheric aging, shows an overall slope of $-0.66$ for
the bulk OA across all campaigns. This result is comparable to slopes of $-0.58$ and
$-0.47$ obtained in previous studies, further illustrating common pathways in OA aging
(Chen et al., 2015; Zhou et al., 2020).

The average HRMSs of OA between the remote site (Waliguan) and the urban site
(Lhasa) were directly compared to investigate the difference in ionic compositions (Fig.
Waliguan was selected for comparison due to its representation of overall highly aged OA, a characteristic shared with other remote sites (Fig. S3). The HRMS of OA at Waliguan and Lhasa displayed significant differences. At Waliguan, the $m/z$ 44, predominantly composed of CO$_2^+$ and a key marker for OOA, was the most prominent peak (18%) in the OA HRMS. The CO$_2^+$ and its related ions (CO$^+$, H$_2$O$^+$, HO$^+$ and O$^+$) together contributed over 41% of the total OA signals. Additionally, two oxygenated ion fragments (C$_x$H$_y$O$_1^+$ and C$_x$H$_y$O$_2^+$) accounted for as much as 66% of the total OA signals (Fig. 5c), suggesting the highly oxygenated nature of OA at this remote site. In contrast, the OA HRMS at Lhasa was remarkably similar to those observed in urban environments, with significant contributions from four $m/z$ values at 43, 55, 57, and 60. These ions are recognized as markers for less oxidized OA or primary emissions related to traffic, cooking, and biomass burning (Zhang et al., 2005a; Alfarra et al., 2007; He et al., 2010), making up a significant contribution to the OA signals in Lhasa. Specifically, non-oxygenated ion fragments (C$_x$H$_y^+$) contributed as much as 64.5% of the total OA in Lhasa, whereas the oxygenated fragments accounted for only 33.6%. This pattern of fresh ion fragments in the OA HRMS in Lhasa is comparable to those measured at other urban cities, such as 56% and 59% in Lanzhou (Xu et al., 2014b; Xu et al., 2016), 51.2% in Nanjing (Wang et al., 2016), and 51.2% in New York (Sun et al., 2011).

### 4.4 OA components from PMF source apportionment

Source apportionments of OA were performed using PMF analysis on OA HRMS data for each field campaign. Figure 6 presents the average mass contributions of OA components from the selected 2–4 factor solutions across eight field campaigns, while Figure S4 details the specific HRMS signatures for each OA component. In regions with limited local emissions but significant influence from regional transport, such as NamCo, LHG, and Bayanbulak, two secondary OA factors with different oxidation degrees, namely a less oxidized OOA (LO-OOA) and a more oxidized OOA (MO-OOA), were identified. For instance, during the NamCo campaign, the MO-OOA and LO-OOA accounted for 59.0% and 41.0% of the total OA mass, with average O/C ratios of 0.96 and 0.49, respectively. The Bayanbulak campaign exhibited a similar result, with MO-OOA (average O/C of 1.12) and LO-OOA (average O/C of 0.55) accounting for 66.3% and 33.7% of the OA mass, respectively. Contrastingly, the LHG campaign revealed a different pattern, with 24.9% MO-OOA and 75.1% LO-OOA, albeit with
higher O/C ratios of 1.29 and 1.08, respectively. Note that the properties of each OOA factor could be different across the locations in the TP despite the same name. Additionally, biomass-burning-related OA (BBOA) was also a prevalent component in the TP. At QOMS, the OA was composed by 42.4% MO-OOA, 43.9% BBOA, and 13.9% nitrogen-containing OA (NOA), with average O/C ratios of 1.34, 0.85, and 1.10, respectively. The high O/C ratio and significant contributions of BBOA and NOA at QOMS were linked to biomass burning emissions transported from South Asia during the pre-monsoon season (Cong et al., 2015; Zhang et al., 2018; Kang et al., 2019a). At Waliguan, the OA was composed by 34.4% MO-OOA, 40.4% relatively aged BBOA (agBBOA), 18.3% BBOA, and 6.9% hydrocarbon-like OA (HOA), with average O/C ratios of 1.42, 1.02, 0.69, and 0.33, respectively. The agBBOA exhibited an enhanced contribution to OA as the OA mass concentration increased, ranging from ∼10% to 70% when OA mass varied from <1.0 µg m$^{-3}$ to 7 µg m$^{-3}$ (Zhang et al., 2019). High contributions of BBOA at Waliguan were associated with regional transport of biomass burning emissions from areas in the northeast (Zhang et al., 2019). At Ngari, the OA was composed by 43.7% MO-OOA, 28.5% LO-OOA, and 27.8% BBOA, with average O/C ratios of 1.43, 1.00, and 0.56, respectively. In contrast, the Motuo exhibited OA components of 36.9% MO-OOA, 46.9% LO-OOA, and 16.2% BBOA, with O/C ratios of 1.30, 1.11, and 0.25, respectively. The lower BBOA contribution and O/C ratio at Motuo suggest a weaker local biomass burning emissions. At urban Lhasa, four OA factors were identified including an OOA with O/C ratio of 0.54 and three primary OA components, i.e., BBOA (O/C of 0.13), cooking-related OA (COA, O/C of 0.12), and HOA (O/C of 0.11). These components were markedly different from those at the above remote sites with the three primary OA components accounting for more than 60% of the total OA, suggesting the abundant primary aerosol sources from the residential activities. In addition, the BBOA contribution increased significantly (up to 36%) during a major local festival in Lhasa, suggesting the crucial aerosol source from biomass burning during religious activities in the city (Zhao et al., 2022).

In summary, our study identified diverse OA components with varying O/C ratios at different sites, indicating the heterogeneity of sources and oxidation states of OA across the TP regions.

### 4.5 Number concentrations of submicron aerosols and cloud condensation nuclei

The measurement of particle number size distribution (PNSD) was useful for studying
the formation and growth mechanisms of aerosol particles in the atmosphere. Figure 7a shows the high-resolution temporal variations of the PNSDs during four field campaigns (QOMS, Motuo, LHG, and Lhasa), revealing significant variability in number concentrations and size distribution patterns across the different sites. On average, the total number concentration was 709.3 and 3994.4 cm$^{-3}$ at QOMS and Lhasa, respectively, while it was 1639.2 and 1462.0 cm$^{-3}$ at Motuo and LHG. Notably, the variations in particle number concentrations were not consistent with mass concentrations measured from the HR-ToF-AMS at the four sites (Table 2). For instance, although the PM$_1$ mass concentration at Lhasa was comparable to that at QOMS (4.7 versus 4.4 µg m$^{-3}$), the number concentration at Lhasa was more than five times higher than that at QOMS. This inconsistency was mainly related to the difference on size distribution at different sites. As mentioned above, submicron aerosols at QOMS were predominantly secondary due to long-range transport from South Asia and characterized by accumulation mode size. In contrast, Lhasa exhibited fresher aerosols, emitted from local residential activities and characterized by Aitken mode size. The variation in submicron aerosol sizes across the TP was further evidenced by the peak diameters in the average mass and number size distributions (Figs. 4a and 7b). For instance, the average OA mass size distribution peaked at 510.2 and 430.5 nm in $D_{va}$ for QOMS and Motuo, respectively. Meanwhile, the average number size distributions at these two sites had peak at 109.4 and 131.0 nm in $D_{nm}$. In contrast, Lhasa displayed significantly smaller peak diameters of only 228.1 nm in $D_{va}$ and 28.9 nm in $D_{nm}$.

New particle formation (NPF) events were observed at several sites in our study. Typically, an NPF event is characterized by a rapid burst in nucleation mode followed by the subsequent growth into larger particles, as defined as banana-shaped temporal developments in the PNSD (Dal Maso et al., 2005). Figure 7a displays the cases of banana-shaped patterns in the PNSD, which were frequently observed at urban Lhasa. Throughout the 27-day Lhasa campaign, a total of 10 NPF events were observed (Zhao et al., 2022). In contrast, such banana-shaped pattern in the PNSD was relatively rare at the other three remote sites (QOMS, Motuo, and LHG), which might be related to their predominance of long-range transported aerosol with overall highly-aged states and limited gaseous precursors.

Cloud condensation nuclei (CCN) is a distinct class of atmospheric aerosol particles which could be activated as cloud droplets at a certain supersaturated water vapor
condition and played important roles in cloud formation, precipitation, climate change, and regional hydrological cycle (Andreae and Rosenfeld, 2008). Across the TP field campaigns, CCN measurements were conducted at three sites: Motuo, Waliguan, and LHG. The temporal variations of CCN number concentrations at each SS exhibited a similar trend with the total number concentration from the SMPS measurement and the PM$_1$ mass concentration from the HR-ToF-AMS measurement during each campaign. On average, the CCN number concentrations at Motuo were 974.0, 1142.6, 1240.1, 1296.5, and 1337.9 cm$^{-3}$ at SS level of 0.2%, 0.4%, 0.6%, 0.8%, and 1.0%, respectively. At Waliguan, relatively comparable average values of 233.7, 857.8, 1138.7, 1313.1, and 1407.0 cm$^{-3}$ were observed at corresponding SS levels. In contrast, LHG exhibited significantly lower average CCN concentration of 120.5, 340.1, 417.8, 468.0, and 504.5 cm$^{-3}$ at the same SS levels, respectively (Table 2). The lower CCN number concentrations at LHG compared to Waliguan and Motuo were consistent with its lower PM$_1$ mass loading. Comparing with other regions, the CCN number concentrations at the three TP sites were almost an order of magnitude lower than those observed in polluted urban environments or from specific combustion emissions. For instance, CCN concentrations reached 12963 cm$^{-3}$ (SS = 0.70%) in Wuqing, 9890 cm$^{-3}$ (SS = 0.86%) in Beijing (Deng et al., 2011; Gunthe et al., 2011), 7913 cm$^{-3}$ (SS = 0.70%) at Panyu in the Pearl River Delta, as well as 11565 cm$^{-3}$ (SS = 0.87%) and 10000 cm$^{-3}$ (SS = 0.80%) during unique biomass burning plumes (Rose et al., 2010; Zhang et al., 2020). However, the CCN values from our study were comparable to those measured at eight remote marine sites in the South China Sea (228–2150 cm$^{-3}$ at SS = 0.87%) and the amazon rain forest (941 cm$^{-3}$ at SS = 0.74%) (Pöhlker et al., 2016; Atwood et al., 2017). These comparisons again highlight the overall clean atmospheric condition in the TP.

4.6 Aerosol optical properties and light absorptions from BC and BrC

In this study, the parameters of $B_{\text{scat}}$, $B_{\text{abs}}$, and SSA of fine particles at 405 nm were observed at the field campaigns of QOMS, Motuo, Waliguan, Ngari, and Lhasa, to explore the variations in aerosol optical properties across the TP. On average, the $B_{\text{scat}}$ and $B_{\text{abs}}$ at 405 nm during the five campaigns were 121.9, 44.9, 36.3, 8.9, and 2.1 Mm$^{-1}$ and 10.8, 7.0, 4.1, 3.6, and 1.9 Mm$^{-1}$, respectively. These values yielded average SSA values of 0.89, 0.83, 0.86, 0.67, and 0.52, correspondingly (Fig. 8a and Table 2). The $B_{\text{scat}}$ and $B_{\text{abs}}$ values at the TP sites were significantly lower than those reported in various urban areas in China, such as 459.5 and 47.2 Mm$^{-1}$ at 630 nm in Beijing (Xie
et al., 2019), 272 and 31 Mm$^{-1}$ at 532 nm in Xi'an (Zhu et al., 2015), and 418 and 91 Mm$^{-1}$ at 540 nm in Guangzhou (Andreae et al., 2008), again suggesting the overall clean atmospheric condition in the TP. Although the PM$_1$ mass concentrations at QOMS was comparable to or even lower than those at the other four sites, QOMS exhibited the highest $B_{\text{scat}}$, $B_{\text{abs}}$, and SSA values. This discrepancy may be due to variations in aerosol chemical compositions and their mass scattering and absorbing efficiencies. In contrast, Lhasa exhibited a significantly lower SSA compared to the other four remote sites, suggesting a prevalence of fresh aerosols in the urban environment. On the other hand, aerosols at the four remote sites were highly aged, leading to significant photobleaching of BrC chromophores and an obvious decrease in their light absorptivity.

Real-time online measurements of particle $B_{\text{abs}}$ at seven wavelengths ranging from 370 to 950 nm were also conducted using an aethalometer at QOMS, NamCo, and Waliguan, respectively, to explore regional variations in aerosol absorption properties across the TP. Overall, the muti-wavelength $B_{\text{abs}}$ decreased significantly with the increasing wavelength during the three measurement campaigns, with fitted AAE values of 1.73, 1.28, and 1.12, respectively (Fig. 8b). The average $B_{\text{abs}}$ at the shortest wavelength of 370 nm was 13.40, 3.25, and 2.66 Mm$^{-1}$ at the three sites, respectively (Table 2). Despite a relatively low PM$_1$ mass concentrations at QOMS, the $B_{\text{abs}}$ at 370 nm was five times higher than that at Waliguan, mainly due to a higher contribution of light-absorbing aerosol components in the southern TP. Specifically, OA and BC together accounted for nearly 80% of the total PM$_1$ at QOMS, whereas this contribution decreased to only 37.5% at Waliguan. The obviously higher AAE at QOMS also suggested a dominant light-absorbing contribution from BrC or the significant lensing effect of coated BC (Zhang et al., 2021). The inserted plots in Fig. 8b illustrate significant decreases in particle $B_{\text{abs},\text{BC}}$ and $B_{\text{abs},\text{BrC}}$ to total $B_{\text{abs}}$ with increasing wavelength, yet their contributions to total $B_{\text{abs}}$ ($f_{\text{abs},\text{BC}}$ and $f_{\text{abs},\text{BrC}}$) varied inversely. BC was the primary light-absorbing component across all the three sites, contributing 66.9%, 78.7%, and 77.6% to the total $B_{\text{abs}}$ at 370 nm at QOMS, NamCo, and Waliguan sites, respectively; its contribution increased apparently with longer wavelengths (Table 2). Conversely, BrC showed more significant contributions to total $B_{\text{abs}}$ at shorter wavelengths. For instance, the average $B_{\text{abs},\text{BrC}}$ at 370 nm were 4.42, 0.69, and 0.60 Mm$^{-1}$ at the three sites, respectively, ultimately contributing 33.1%, 21.3%, and 22.4% to the total $B_{\text{abs}}$. The significantly higher values of total $B_{\text{abs}}$, $B_{\text{abs},\text{BC}}$, $B_{\text{abs},\text{BrC}}$, and
4.7 Estimation of aerosol radiative forcing in the different TP regions

Atmospheric aerosols play a significant role in impacting Earth’s climate systems through affecting solar radiation and exerting a positive forcing on the energy budget (Bond and Bergstrom, 2006). In this study, aerosol direct radiative forcings (DRF) caused by BC, organic carbon (OC), and water-soluble ions (WSIs) are estimated, respectively, using Santa Barbara DISORT (Discrete Ordinate Radiative Transfer) Atmospheric Radiative Transfer (SBDART) model (Ricchiazzi et al., 1998). A detailed introduction and operation of this model are described in Text S6 in the supplementary material. Since the model’s performance is evaluated and calibrated by comparing the values with measurements from Aethalometer and PAX results, the aerosol DRF estimations are limited to QOMS, NamCo, and Waliguan. Furthermore, these three sites are located in the southern, central, and northern regions of the TP, respectively, which allows for an exploration of regional variations in aerosol DRF across the TP.

Figure 9 presents the results of DRFs attributed to BC, OC, and WSIs across the three campaigns. BC demonstrated a pronounced warming effect at the top of the atmosphere, with average DRF values of $+2.5 \pm 0.5$, $+2.1 \pm 0.1$, and $+1.9 \pm 0.1$ W m$^{-2}$ during the QOMS, Waliguan, and NamCo campaigns, respectively. In contrast, BC exhibited a noticeable cooling effect at the earth’s surface, with average DRF values of $-4.7 \pm 0.8$, $-4.1 \pm 0.2$, and $-3.7 \pm 0.1$ W m$^{-2}$ across the three campaigns. The combination of these two effects resulted in significantly high net atmospheric forcing by BC, amounting to $+7.3 \pm 1.2$, $+6.2 \pm 0.3$, and $+5.6 \pm 0.2$ W m$^{-2}$ for the three campaigns. These results suggest the important radiative effect of BC in the TP, especially in the southern region influenced by long-range transported biomass burning emission from South Asia. In contrast, OC and WSIs exhibited cooling effects at both the top of the atmosphere and the Earth’s surface, characterized by negative and relatively low average DRFs.

Consequently, the net atmospheric forcings for OC and WSIs were notably lower compared to BC across the three campaigns, with values of $+2.0 \pm 1.2$, $+0.7 \pm 0.2$, and $+0.9 \pm 0.7$ W m$^{-2}$ for OC, and $+1.9 \pm 0.8$, $+1.4 \pm 0.6$, and $+1.2 \pm 0.2$ W m$^{-2}$ for WSIs at QOMS, Waliguan and NamCo, respectively. Interestingly, at QOMS, the average atmospheric DRF of OC accounted for 27.3% of that of BC, whereas at Waliguan and
NamCo, the fractions were only 11.1% and 15.7%, respectively. The higher atmospheric DRF observed at QOMS suggests a dominant contribution from light-absorbing BC and BrC aerosols compared to Waliguan and NamCo.

It was worth noting that the simulations of DRF effects in this study were only conducted at three specific sites over limited periods. Future research should focus on long-term comprehensive measurements and DRF simulations across the entire TP to enhance our understanding of aerosol impacts on regional climate.

4.8 Long-range transport of aerosols from surrounding areas

To further understand the potential sources and specific transport pathways of aerosols at each site, particularly for those remote sites, three- or five-days air mass back trajectories were calculated during each measurement period at an ending height of 500 m above ground level every 6h using the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Draxler and Rolph, 2003). The cluster analysis on the trajectories was based on the total spatial minimum variance method. Figure 10 displays the average backward trajectory clusters across all eight field campaigns.

In general, distinct air mass sources were identified across the TP. The five sites (QOMS, Motuo, Lhasa, NamCo, and Ngari) located in the southern or south-central part of the TP generally showed air masses from south or southwest with different transport distances and pathways during their measurement periods. For instance, during the QOMS campaign, 38% of the air masses originated from the west, covering a long distance, while another 40% originated from the southwest, covering a shorter distance. In the Motuo campaign, two major clusters both originated from the southwest with different distances (77% at shorter distance compared to only 13% at a longer distance). Similarly, during the NamCo campaign, two major clusters with comparable contributions (37% and 34%) and transport distances but different pathways were identified from the south. In the Ngari campaign, air masses showed similar transport distances, with 56% originating from southwest and 26% from the south. These air mass clusters originating from the south of the TP generally traverse polluted regions in South Asia such as the Indo-Gangetic Plain, Nepal, and Bangladesh, carrying significant amounts of pollutants, particularly biomass-burning emissions into the inland of the TP. In contrast, air masses at northern sites were primarily influenced by Westerlies and East Asian monsoon during summer season. In the campaigns of Bayanbulak, air
masses originated from the west with different distances, i.e., 69% in relatively shorter
distance versus 18% in a longer distance. During the LHG campaign, the air masses
originated from the northwest of the site with 63% in longer distance but 27% in shorter
distance. For the Waliguan campaign, the air mass originated from two distinct
directions. Most of air masses (57%) came from the northeast of the site with a shorter
distance, while the remaining air masses originated from the west and northwest with
longer distances.

In summary, significant variations in air mass sources and transport pathways were
observed across the TP, particularly between southern and northern regions. These
differences play a crucial role in shaping the different physiochemical and optical
properties of aerosols across the TP regions.

5 Dataset limitations and applications

Our dataset was compiled from eight short-term intensive field observations across the
TP utilizing a suite of high-resolution online instruments. However, it is important to
note that our dataset does have certain limitations that proved to be quite challenging to
address in these remote regions.

The primary limitation stems from the short and inconsistent measurement periods
across different observational years and seasons at different sites, impeding robust
comparisons of aerosol properties across the TP. This limitation also hampers the ability
to ascertain long-term and seasonal characteristics. The harsh natural environments,
challenging weather conditions, limited logistical support and instruments, and
stringent instrumental requirements (such as the necessity for comprehensive field
stations with stable power supply) presented significant challenges during our field
observations in these remote TP regions. It is worth noting that online HR-ToF-AMS
observations, such as the one we conducted, are predominantly short-term and intensive
observations carried out worldwide due to the instability issues and its challenging
maintenance required for long-term measurements. The short-term intensive
measurement is enough to capture and characterize the dynamic evolution of aerosol
properties at a high-time-resolution (Jimenez et al., 2009; Li et al., 2017). Long-term
high-time-resolution observation utilizing HR-ToF-AMS have been rarely conducted
thus far, even in urban environment with relatively favorable observational conditions
and logistic support compared to our remote TP sites. Consequently, performing
continuous long-term observations or simultaneous comparison at multiple sites in these high-altitude remote TP regions, without stable power supply, is exceedingly challenging. Furthermore, assessing the representativeness of each observation for the spatial scale is particularly challenging due to the limited number of observatories across the TP. Actually, these observatories have been strategically established based on the representation of specific geographic and climatic features.

In addition to the limitations above, our team has made significant efforts to conduct this comprehensive observation project over the past decade, aiming to study the regional differences in aerosol sources and properties across the TP. The dataset generated from our project represents the first and exclusive high-time-resolution dataset focusing on atmospheric aerosol physicochemical and optical properties, covering most region of the TP. The applications of this dataset in atmospheric science are multifaceted. Firstly, the high-time-resolution observations offer crucial advantages in understanding the rapid evolution and diurnal variations of aerosol properties during a short period or special event. Furthermore, these observations are invaluable for model simulation and verification, providing a wealth of data points that can be utilized for assessing aerosol loading, chemical composition, size distribution, and other parameters essential for model accuracy and validation. Such advantages are not achievable with traditional off-line samplings, which typically exhibit low time resolutions ranging from days to weeks. Secondly, the eight sites encompassed in our project effectively represent a wide range of the TP. This is particularly noteworthy considering the limited availability of observatories on the TP. These sites facilitate comparisons of aerosol sources and properties among different regions. Thirdly, our observations encompass a wide range of aerosol physical, chemical, and optical parameters, including aerosol mass loadings, chemical compositions, size distribution, diurnal variations, number concentrations, light scattering and absorption coefficients, and so on. This comprehensive dataset plays a crucial role in fostering a profound understanding of aerosol properties in the TP.

Overall, it is noteworthy that our online observational aerosol datasets, focusing on multiple parameters with at least hourly-scale resolution at various sites across the TP, are the only ones reported to date.

6 Data availability
The high-resolution online measurement datasets, encompassing aerosol physical, chemical, and optical properties over the Tibetan Plateau and its surroundings in our observation project have been released and are now available for download from the National Cryosphere Desert Data Center (https://doi.org/10.12072/ncdc.NIEER.db2200.2022; Xu, 2022). These datasets are provided in an Excel file comprising eight worksheets. The first sheet of the Excel file contains a concise description of the dataset, including the dataset name, observation stations, sampling periods, online instruments used, and corresponding references. The remaining seven sheets present the high-resolution measurement data obtained from the online instruments employed during the eight campaigns. These instruments include HR-ToF-AMS, SMPS, PAX, aethalometer, and CCN-100.

7 Conclusions

This study presents a comprehensive dataset encompassing aerosol physicochemical and optical properties, with a particular focus on high-resolution size-resolved chemical characteristics and sources of submicron aerosols, conducted at eight different sites of the TP and its surroundings. The datasets collected offer insights into temporal and diurnal variations, size distribution of PM$_1$ chemical compositions, HRMS and chemical components of OA, particle light scattering and absorption coefficients, and CCN number concentrations at different supersaturations in different campaigns.

The datasets offer valuable insights into regional variations in aerosol properties and sources. In the southern TP region, atmospheric aerosols were primarily influenced by biomass burning emissions transported from South Asia, resulted in high mass contributions (>70%) of CAs and overall neutralized PM$_1$, as well as an enhanced light absorption capability of aerosols. In contrast, in the northern TP, secondary inorganic species, particularly sulfate, contributed significantly to total PM$_1$ due to the regional transport of anthropogenic aerosol and gaseous precursor emissions from urban areas in northwestern China. Furthermore, in contrast to the well-mixed, highly-aged, and regionally transported aerosols observed in the remote sites, atmospheric aerosols in the urban Lhasa were mainly originated from local primary sources such as cooking, traffic vehicle exhausts, and biofuel combustion during the residential activities.

Consequently, these aerosol particles were relatively fresh, characterized by small size and low oxidation degree, but exhibited a high frequency of NPF origins.
Appendix A: Main Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>TP</td>
<td>Tibetan Plateau</td>
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<tr>
<td>HR-ToF-AMS</td>
<td>high-resolution time-of-flight aerosol mass spectrometer</td>
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<td>SMPS</td>
<td>scanning mobility particle sizer</td>
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<tr>
<td>PAX</td>
<td>photo-acoustic extintiometer</td>
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<td>CCN</td>
<td>cloud condensation nuclei</td>
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<tr>
<td>SS</td>
<td>supersaturation</td>
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<td>PM$_1$</td>
<td>submicron aerosol</td>
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<td>BC</td>
<td>black carbon</td>
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<tr>
<td>BrC</td>
<td>brown carbon</td>
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<td>OA</td>
<td>organic aerosol</td>
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<tr>
<td>SNA</td>
<td>sulfate, nitrate, and ammonium</td>
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<tr>
<td>$D_m$</td>
<td>mobility diameter</td>
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<tr>
<td>$D_{va}$</td>
<td>aerodynamic diameter</td>
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<tr>
<td>CE</td>
<td>collection efficiency</td>
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<tr>
<td>HRMS</td>
<td>high-resolution mass spectrum</td>
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<tr>
<td>PBL</td>
<td>planetary boundary layer</td>
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<tr>
<td>O/C</td>
<td>oxygen-to-carbon ratio</td>
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<tr>
<td>H/C</td>
<td>hydrogen-to-carbon ratio</td>
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<tr>
<td>N/C</td>
<td>nitrogen-to-carbon ratio</td>
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<tr>
<td>OM/OC</td>
<td>organic matter-to-organic carbon ratio</td>
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<tr>
<td>PMF</td>
<td>positive matrix factorization</td>
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<tr>
<td>OOA</td>
<td>oxygenated organic aerosol</td>
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<tr>
<td>LO-OOA</td>
<td>less oxidized oxygenated organic aerosol</td>
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<tr>
<td>MO-OOA</td>
<td>more oxidized oxygenated organic aerosol</td>
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<tr>
<td>BBOA</td>
<td>biomass-burning-related organic aerosol</td>
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<tr>
<td>agBBOA</td>
<td>aged biomass-burning-related organic aerosol</td>
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<td>NOA</td>
<td>nitrogen-containing organic aerosol</td>
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<tr>
<td>HOA</td>
<td>traffic-related hydrocarbon-like organic aerosol</td>
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<tr>
<td>COA</td>
<td>cooking-related organic aerosol</td>
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<tr>
<td>PNSD</td>
<td>particle number size distribution</td>
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<tr>
<td>NPF</td>
<td>new particle formation</td>
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<tr>
<td>$B_{\text{scat}}$</td>
<td>light scattering coefficient</td>
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<tr>
<td>$B_{\text{abs}}$</td>
<td>light absorption coefficient</td>
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<tr>
<td>$B_{\text{ext}}$</td>
<td>light extinction coefficient</td>
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<td>SSA</td>
<td>single scattering albedo</td>
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<td>AAE</td>
<td>absorption Ångström exponents</td>
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<td>$B_{\text{abs},BC}$</td>
<td>light absorption coefficient from BC</td>
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<td>$B_{\text{abs},BrC}$</td>
<td>light absorption coefficient from BrC</td>
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<td>OC</td>
<td>organic carbon</td>
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<td>WSIs</td>
<td>water-soluble ions</td>
</tr>
<tr>
<td>DRF</td>
<td>direct radiative forcing</td>
</tr>
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</table>

Author Contributions. JX designed the study, XZ, WZ, and JX wrote the manuscript. JX and SK organized and supervised the field measurement campaigns, JX, XZ, WZ, LZ, MZ, JS, JShi, YL, CX, YT, KL, XG, and QZ conducted the field measurements, JX, XZ, WZ, and YT analyzed the data. All authors reviewed and commented on the final form of the manuscript.
Competing interests. The authors declared that they have no competing interests.

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Van Damme, M., Erismann, J. W., Clarisse, L., Dammers, E., Whitburn, S., Clerbaux, C., Dolman, A. J.,


**Figures**

**Figure 1.** (a) Geographical locations of the observation sites (see Table 1 for full name and characteristics of each site) in the Tibetan Plateau and its surroundings in this study (The geographical base map is created with ArcGIS). Fieldwork photographs illustrate the real observation conditions and surroundings at each site. (b) The normal sampling setups of instruments during the online aerosol observations.
Figure 2. Regional distribution of average mass concentrations (values marked in the central of each pie chart with unit of µg m⁻³) and chemical compositions (percentage values around each pie chart) of submicron aerosols (PM₁) during the eight online aerosol field measurements in the Tibetan Plateau and its surroundings (The geographical base map is created with ArcGIS). The concentration at each site is presented in ambient condition.
Figure 3. Regional difference of bulk acidity of submicron aerosols based on the scatterplot analysis and linear regression of measured $\text{NH}_4^+$ versus predicted $\text{NH}_4^+$ during the eight aerosol field measurement campaigns in the Tibetan Plateau and its surroundings (The geographical base map is created with ArcGIS).
Figure 4. (a) Average size distributions of organic mass concentrations during six field measurement campaigns in the Tibetan Plateau and its surroundings. (b) Diurnal variations of total PM$_1$ mass concentrations during the eight field measurement campaigns in the Tibetan Plateau and its surroundings. Insert graph in (a) is the scatter plot of peak diameters in their size distributions versus the average O/C ratio of organics.

Figure 5. (a) Box plots of the average O/C ratios and (b) Van Krevelen diagram of H/C versus O/C among the eight field measurement campaigns in this study. (c) The average HRMSs of OA colored with different ion categories during the Waliguan and Lhasa measurement campaigns. The whiskers of boxes indicate the 90th and 10th percentiles, the upper and lower boundaries of boxes indicate the 75th and 25th percentiles, the lines in the boxes indicate the median values, the markers indicate the mean values, and similarly hereinafter.
Figure 6. Regional distribution of OA components from PMF source apportionment during the eight online aerosol field measurements in the Tibetan Plateau and its surroundings (The geographical base map is created with ArcGIS). Values marked in the central of each pie chart are average OA mass with unit of µg m$^{-3}$, while the percentage values around the pie chart are the mass contributions of each OA component. The O/C ratio of each OA component is also marked around each pie chart.
Figure 7. (a) Temporal variations of the size distributions of particle number concentrations during the aerosol field measurement campaigns at QOMS, Motuo, LHG, and Lhasa sites. (b) The average size distribution of particle number concentration during entire measurement period at each site.
Figure 8. Box plots of (a) the average particle light scattering coefficient ($B_{\text{scat}}$), light absorption coefficient ($B_{\text{abs}}$), and single scattering albedo (SSA) during the five aerosol field measurement campaigns at QOMS, Motuo, Waliguan, Ngari, and Lhasa sites, and (b) the particle $B_{\text{abs}}$ at seven wavelengths measured by aethalometers at QOMS, NamCo, and Waliguan sites. The dashed lines in the boxes in (b) show the power-law fit of the average $B_{\text{abs}}$ as a function of wavelength. The inserted plots in (b) are the apportioned contributions of BC and BrC to total $B_{\text{abs}}$ at different wavelengths.

Figure 9. Box-plots of the modelled direct radiative forcing (DRF) at the top of the atmosphere (TOA), the atmosphere (ATM), and the earth’s surface (SUR) caused by black carbon (BC), organic carbon (OC), and water-soluble ions (WSIs) during the QOMS, NamCo, and Waliguan campaigns.
Figure 10. Air mass backward trajectory clusters during the eight field campaigns in the Tibetan Plateau and its surroundings in our study (The geographical base map is created with Igor Pro). Only major trajectory clusters during each field campaign are displayed with the contributions marked correspondingly.
### Table 1. Detailed information about the full name and geographic characteristic of observation station, sample period, online instruments, and corresponding references during each aerosol field measurement campaigns over the Tibetan Plateau and its surroundings in this study.

<table>
<thead>
<tr>
<th>Station</th>
<th>Full Station Name</th>
<th>Lat. (°N)</th>
<th>Long. (°E)</th>
<th>Alt. (m)</th>
<th>Sample Period</th>
<th>Online Instruments</th>
<th>References</th>
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<td>QOMS</td>
<td>Qomolangma Station for Atmospheric and Environmental Observation and Research, Chinese Academy of Sciences</td>
<td>28.36</td>
<td>86.95</td>
<td>4276</td>
<td>12 April to 12 May 2016</td>
<td>√</td>
<td>√</td>
</tr>
<tr>
<td>Motuo</td>
<td>Motuo County, Linzhi City, Tibet Autonomous Region, China</td>
<td>29.30</td>
<td>95.32</td>
<td>1305</td>
<td>26 Mar to 22 May 2021</td>
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<td>√</td>
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<td>NamCo</td>
<td>Nam Co Station for Multisphere Observation and Research, Chinese Academy of Sciences</td>
<td>30.77</td>
<td>90.95</td>
<td>4730</td>
<td>31 May to 1 July 2015</td>
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<td>Ngari</td>
<td>Ngari Station for Desert Environment Observation and Research, Chinese Academy of Sciences</td>
<td>33.39</td>
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<td>1 Jun to 5 Jul 2022</td>
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<td>√</td>
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<td>Waliguan</td>
<td>China Global Atmospheric Watch Baseline Observatory, Mount Waliguan Base</td>
<td>36.28</td>
<td>100.90</td>
<td>3816</td>
<td>1 July to 31 July 2017</td>
<td>√</td>
<td>√</td>
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<td>LHG</td>
<td>Qilian Observation and Research Station of Cryosphere and Ecologic Environment, Chinese Academy of Sciences</td>
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<td>96.51</td>
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<td>4 August to 29 August 2020</td>
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<tr>
<td>Bayanbuluk</td>
<td>Bayanbulak Town, Hejing County, Bayingolin Mongolian Autonomous Prefecture, Xinjiang Uyghur Autonomous Region, China</td>
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<td>84.35</td>
<td>2454</td>
<td>29 August to 26 September 2021</td>
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<td>Lhasa</td>
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<td>29.65</td>
<td>91.03</td>
<td>3650</td>
<td>31 August to 26 September 2019</td>
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Table 2. Summary of the average values measured with various instruments during the eight aerosol field measurement campaigns in the TP and its surroundings in this study.

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<th>Measurement items</th>
<th>QOMS</th>
<th>Motuo</th>
<th>NamCo</th>
<th>Ngari</th>
<th>Waliguan</th>
<th>LHG</th>
<th>Bayanbulak</th>
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<td>4.1</td>
<td>9.1</td>
<td>3.0</td>
<td>1.9(^*)</td>
<td>4.7</td>
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<td>57.0</td>
<td>68.0</td>
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<td>2.0</td>
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<td>Ammonium</td>
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<td>8.6</td>
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<td>O/C</td>
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<td>B(_{abs,BC,370}) (Mm(^{-1}))</td>
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<td>f(<em>{B</em>{abs,BrC,370}})%</td>
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<td>f(<em>{B</em>{abs,BC,370}})%</td>
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<td>CCN number conc. (SS 0.2%)</td>
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\(^*\)only non-refractory PM\(_1\) is reported at Bayanbulak due to the absence of BC observation.
Table 3. Summary of the average PM$_1$ mass concentrations ($\mu$g m$^{-3}$) measured by the Aerodyne AMSs at various high-altitude and remote sites worldwide.

<table>
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<th>Observation Sites</th>
<th>Latitude (°N)</th>
<th>Longitude (°E)</th>
<th>Altitude (m a.s.l.)</th>
<th>PM$_1$ mass ($\mu$g m$^{-3}$)</th>
<th>References</th>
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<tr>
<td>QOMS, China</td>
<td>28.36</td>
<td>86.95</td>
<td>4276</td>
<td>4.4</td>
<td>This study &amp; Zhang et al. (2018)</td>
</tr>
<tr>
<td>Motuo, China</td>
<td>29.30</td>
<td>95.32</td>
<td>1305</td>
<td>5.7</td>
<td>This study</td>
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<tr>
<td>NamCo, China</td>
<td>30.77</td>
<td>90.95</td>
<td>4730</td>
<td>2.0</td>
<td>This study &amp; Xu et al. (2018)</td>
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<td>79.70</td>
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<td>This study</td>
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<td>Waliguan, China</td>
<td>36.28</td>
<td>100.90</td>
<td>3816</td>
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<td>96.51</td>
<td>4180</td>
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<tr>
<td>Bayanbulak, China</td>
<td>42.83</td>
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<td>2454</td>
<td>1.9$^a$</td>
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<td>Lhasa, China</td>
<td>29.65</td>
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<td>3650</td>
<td>4.7</td>
<td>This study &amp; Zhao et al. (2022)</td>
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<td>4730</td>
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<td>Menyuan, China</td>
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<td>Mt. Bachelor, USA</td>
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<td>Mt. Whistler, Canada</td>
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<td>-122.95</td>
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<td>Sun et al. (2009)</td>
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<td>Mt. Cimone, Italy</td>
<td>44.18</td>
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<td>Puy de Dôme, France</td>
<td>45.77</td>
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<td>1465</td>
<td>5.58</td>
<td>Freney et al. (2011)</td>
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<td>Sub-Antarctic Bird Island</td>
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$^a$only non-refractory PM$_1$ is reported at Bayanbulak due to the absence of BC observation.