Measurement report: Vertical profiling of particle size distributions over Lhasa, Tibet: Tethered balloon-based in-situ measurements and source apportionment

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Abstract. In-situ measurements of vertically resolved particle size distributions based on a tethered balloon system were carried out for the first time in the highland city of Lhasa over the Tibetan Plateau in summer 2020, using portable optical counters for the size range of 0.124–32 μm. The vertical structure of 112 aerosol profiles was found to be largely shaped by the evolution of the boundary layer (BL), with a nearly uniform distribution of aerosols within the daytime mixing layer and a sharp decline with the height in the shallow nocturnal boundary layer. During the campaign, the average mass concentration of particulate matters smaller than 2.5 μm in aerodynamic diameter (PM₂.₅) within the BL was around 3 μg m⁻³, almost four times of the amount in the free troposphere (FT), which was rarely affected by surface anthropogenic emissions. Though there was a lower level of particle mass in the residual layer (RL) than in the BL, a similarity in particle mass size distributions (PMSDs) suggested that particles in the RL might be of the same origin as particles in the BL. This was also in consistence with the source apportionment analysis based on the PMSDs. Three distinct modes were observed in the PMSDs for the BL and the RL. One mode was exclusively coarse particles up to roughly 15 μm and peaked around 5 μm. More than 50% of total particle mass was often contributed by coarse mode particles in this area, which was thought to be associated with local dust
resuspension. The mode peaking over 0.5–0.7 μm was representative of biomass burning on religious holidays and was found to be most pronounced on holiday mornings. The contribution from the religious burning factor rose from about 25% on non-holidays to nearly 50% on holiday mornings. The mode dominated by particles smaller than 0.3 μm was thought to be associated with combustion related emissions and/or secondary aerosol formation. In the FT coarse mode particles only accounted for less than 10% of the total mass and particles larger than 5 μm were negligible. The predominant submicron particles in the FT might be related to secondary aerosol formation and the aging of existed particles. To give a full picture of aerosol physical and chemical properties and better understand the origin and impacts of aerosols in this area, intensive field campaigns involving measurements of vertically resolved aerosol chemical compositions in different seasons would be much encouraged in the future.

1 Introduction

Lhasa, the provincial capital of Tibet, lies almost in the heart of the sparsely populated Tibetan Plateau, which is the highest plateau in the world with an average altitude of approximately 4320 m (Zhang et al., 2021). As a result of its unique topography, the Tibetan Plateau plays a vital role in East Asian Summer Monsoon, thereby in the regional and global climate (He et al., 2019; Chiang et al., 2020). As the most urbanized and populated highland city in Tibet, Lhasa is quite suitable for exploring the impact of anthropogenic activities on atmospheric components over this remote region.

In the past decade, the population in Lhasa has increased from about 0.55 million in 2010 to nearly 0.87 million in 2020, according to the newly published data from the Seventh National Population Census (http://www.xizang.gov.cn/zwgk/zfsj/ndtjgb/202105/t20210520_202889.html, last access on 18 June 2021). Rapid urbanization and economic growth, also the ever thriving tourism, have given rise to a marked increase in residential, industrial and traffic emissions. As a religious and cultural center, the widespread religious burning was also an important source for
various air pollutants (Cui et al., 2018b; Wei et al., 2019a). In the low oxygen-containing atmosphere (molecules per volume) at an altitude of about 3650 m, incomplete combustion could further enhance emissions of air pollutants such as black carbon and organic aerosols. The probable photochemically active atmosphere due to strong solar radiation at such a high altitude and low latitude might facilitate the formation of secondary aerosols and gaseous pollutants like ozone (Lin et al., 2008), though the extent to which secondary production is enhanced remains uncertain considering pressure-dependent reaction rate constants (Atkinson, 2000). In addition, the topography of Lhasa being located in the Lhasa River Valley and surrounded by high mountains prevents effective dilution of air pollutants. Consequently, serious air pollution comparable to the level in more urbanized and developed cities at lower altitudes has been observed (Ran et al., 2014; Duo et al., 2018; Li et al., 2019). However, it was demonstrated that air quality in Lhasa has been slightly improved in 2017 compared with previous years, due to the implementation of Action Plan on Prevention and Control of Air Pollution all over China since 2013 (Yin et al., 2019).

Previous studies on aerosols in Lhasa were quite limited. One particular focus was the chemical composition of single particle (Zhang et al., 2001a; Duo et al., 2015), bulk aerosol (Cong et al., 2011; Gong et al., 2011; Liu et al., 2013; Chen et al., 2018) and size-segregated aerosols (Zhang et al., 2001b; Wan et al., 2016; Cui et al., 2018a; Wei et al., 2019a, 2019b) near the ground. Results from these studies revealed that vehicular exhaust, religious activities involved incense burning and biomass burning, and the suspension of mineral dust were major sources. It was also pointed out from the results of the backward trajectory analysis that local emissions dominated during the monsoon season (Wei et al., 2019b). Aerosol optical properties such as multi-wavelength aerosol absorption near the ground (Zhu et al., 2017) and aerosol optical depth (Bai et al., 2000; Zhu et al., 2019) have also been investigated. However, the particle size distribution, as a critical microphysical property, has rarely been studied. Using a portable optical particle counter, Cui et al. (2018b) measured particle size distributions (14 size bins) within the range of 0.14–3 μm for three weeks in summer 2016 in urban Lhasa and emphasized the significance of religious burning to fine particles. In-situ measured vertical profiles of particle size distributions with a rather coarse size resolution were
obtained from three balloon measurements in 1999 over Lhasa, but were only used to explore an enhancement in fine particle concentrations near the tropopause (Tobo et al., 2007). The vertical structure of particle size distributions and its temporal variation within the planetary boundary layer has not yet been examined. To get a better knowledge on the current status of aerosol microphysical properties in Lhasa and to advance our understanding on the influence of anthropogenic activities, in-situ measurements of vertically resolved particle size distributions with a high vertical resolution are much needed.

In this study, the vertical structure and temporal variability of aerosol profiles were explored, based on in-situ measurements of particle size distributions within 1 km above the ground at a suburban site in Lhasa, using optical counters attached to a tethered balloon. Source apportionment based on measured particles size distributions was further performed to identify possible sources for aerosols in different layers.

2 Measurements

2.1 The field site

The city of Lhasa lies in an east-west oriented valley along the Lhasa River, surrounded by mountains up to an elevation of about 5500 m (Fig. 1a), as shown by SRTM data V4 from Jarvis et al. (2008). The field site is on the Najin campus of Tibet University (29.64° N, 91.18° E) in a suburban area of Lhasa, approximately 5 km to the east of the urban centre (Fig. 1b). As the commercial and religious centre, the urban area features crowded streets and heavy traffic, partly due to the thriving tourism and partly due to frequent religious activities. Religious burnings are constantly practiced and especially vigorous on religious ceremonies in the Potala Palace and the neighbouring famous temples, such as Jokhang Temple and Ramoche Temple, which are all located in this area and attended by many worshippers and tourists throughout the year. Comparatively, the suburb is thinly populated with much less traffic. Thus emissions from residential, vehicular and religious sources are expected to be much weaker in the surrounding area of the site than in the downtown.
2.2 Instruments and data

Measurements were conducted in an open space, being kept about 90 m away from the nearest building, in order to ensure the safety of launching the tethered balloon system. The 60 m³ helium-filled tethered balloon, with a payload capacity of roughly 10 kg at an elevation of 3650 m, was operated by an electric winch that controlled the ascent and descent rate of the fish-shaped balloon. It was scheduled to launch the balloon to at least 800 m above the ground every 3 hours from 07:00 to 22:00 (in Beijing time) during 8-28 August 2020. The actual number of launches and the actual maximum height reached by each flight were subject to unavailable flight permits and unfavorable weather conditions (precipitation and/or strong winds). In total, 56 launches were successfully performed for the entire campaign (Fig. S1a). Most flights of the tethered balloon reached above 600 m, with 3 launches reaching 450–500 m, 15 reaching 500–600 m, 17 reaching 600–700 m, 15 reaching 700–800 m, and 6 reaching 800–850 m. In particular, strong winds were often encountered in the late afternoon, resulting in a lack of observations during this time period (Fig. S1b).

A light-weighted optical counter, the Portable Optical Particle Spectrometer (POPS, Handix Scientific), was attached to the tethered balloon for in-situ measurements of vertical profiles of atmospheric aerosols. Details about the principle on which the instrument operates were given in Gao et al. (2016). The POPS was calibrated before the campaign using ammonium sulfate...
aerosols (for diameters smaller than 1 μm) and polystyrene latex spheres (for diameters larger than 1 μm). The established relationship between the scattering signal and the particle size was used to obtain 42 logarithmically equal size bins over the size range of 0.124~2.55 μm. Ambient aerosols were sufficiently dried before entering the POPS by a homemade silica gel-filled diffusion drier. The flow rate of the instrument was regularly checked during the campaign and used for data correction.

Particle number size distributions (PNSD, \(n(\log D_p)=dN/d\log D_p\)) were obtained with a temporal resolution of 1 s, and integrated over the whole size range to provide total number concentrations \((N_a)\). Particle mass size distributions (PMSD, \(m(\log D_p)=dM/d\log D_p\)) were calculated from PNSDs assuming spherical particles with a density of 1.7 g cm\(^{-3}\) (Sloane et al., 1991; Tao et al., 2014). Mass concentrations of particulate matters smaller than 1 μm and 2.5 μm in aerodynamic diameter (PM\(_1\) and PM\(_{2.5}\)) were obtained under the assumption that the optically equivalent diameter could be considered equal to geometric diameter. The effective diameter \((D_e)\) was the ratio of the third-order moment to the second-order moment of particle size distributions. Similarly, the effective diameter for submicron particles \((D_{e,<1μm})\) and for super-micron particles \((D_{e,1~2.5μm})\) were respectively calculated for particles with the diameter smaller than 1 μm and in the range of 1~2.5 μm.

A portable aerosol spectrometer (Model 11-C, GRIMM Aerosol Technik GmbH & Co. KG) was also employed for 24 launches to concurrently measure the PNSD for dry particles within the size range of 0.25~32 μm. The measuring time interval was set to be 6 s, for which 31 size channels were classified. Sizing of aerosols by the instrument was verified before the campaign for several selected diameters using polystyrene latex spheres. The PNSD measured by GRIMM 11-C was combined with the PNSD simultaneously measured by the POPS, using weighting factors in the overlapped size range, namely, from the lower size limit of GRIMM 11-C \((D_{p,L})\) to the upper size limit of the POPS \((D_{p,U})\). The weighting factors for GRIMM 11-C were derived as \(w_{11-C}(\log D_p)=(\log D_{p,L}-\log D_p)/(\log D_{p,U}-\log D_{p,L})\), while for the POPS they were defined as \(w_{POPS}(\log D_p)=(\log D_{p,U}-\log D_p)/(\log D_{p,U}-\log D_{p,L})\).
Vertical profiles of meteorological parameters, including pressure (MS5540B, Intersema Sensoric SA), temperature (T) and relative humidity (RH) (SHT7x, Sensirion), as well as winds (homemade sensor based on a slotted optical switch (OPB610, TT Electronics) and an electronic compass module (ZCC211N, Shanghai Zhichuan Electronic Tech Co., Ltd)) were measured for each flight with a temporal resolution of 1 s. Potential temperature (θ) and specific humidity (q), two conservative quantities, were calculated following the equations in Ran et al. (2016). Besides, surface meteorological parameters in August were available from an Automatic Weather Station roughly 4.5 km to the northwest of the site in urban Lhasa and were averaged into hourly data, except that winds were recorded as 2-min averages at the beginning of each hour.

Profiles of aerosol and meteorological parameters were all processed into 10 m-averaged data for the subsequent analysis. A mixed layer (ML) for daytime profiles or a nocturnal boundary layer (NBL) for nighttime profiles, mentioned altogether as the boundary layer (BL) in what follows, could be identified for 72 profiles in the dataset of totally 112 profiles. For the remaining 40 profiles, the flight path was entirely within the daytime ML. The height of the BL, denoted as $H_m$ hereafter, was estimated from profiles of aerosols and meteorological parameters based on the gradient method (Seibert et al., 2000; Kim et al., 2007; Ferrero et al., 2010; Ran et al., 2016). For profiles with the top of the ML above the flight limit, $H_m$ was taken as the maximum height reached by the tethered balloon for further statistical analysis. The normalized height ($H_{nor}$) was then calculated as $H_{nor} = height / H_m - 1$ (Ferrero et al., 2014). For several profiles, the free troposphere (FT) that was identified as the layer above the BL without the presence of elevated aerosol layers or the residue from aerosols in the daytime after the collapse of the ML, as well as the residual layer (RL) could also be identified. In addition, the time mentioned anywhere in this study was Beijing time, which was around two hours earlier than local solar time.

### 2.3 Positive Matrix Factorization (PMF)

A bilinear factor-based receptor model, Positive Matrix Factorization (PMF), was used for source apportionment of particle mass in this study based upon observed PMSDs (Paatero and Tapper, 1994; Paatero, 1997). By minimizing an objective
function $Q$, a key parameter to review the distribution of the components and to estimate the stability of the solution (Brown et al., 2015), the measured matrix was decomposed into factor profiles ($F$) and factor contributions ($G$) with non-negativity constraints. The input matrix, comprising PMSDs (22 size bins within 0.124–15 μm) averaged for the ML, the RL and the FT, and the lower 10-m averaged PMSDs in the NBL, was used in 20 random PMF runs (using EPA PMF 5.0 software) for 2 to 5 factors. In total, there were 48 PMSDs for the ML, 23 PMSDs for the RL and 10 PMSDs for the FT. The components in the PMF runs were separated to keep one signal rather significant from those dominated by noise on account of a criterion of the signal-to-noise ratio (S/N) (Amato et al., 2009). The components with the S/N less than 2 were removed from further analysis, while components with the S/N above 2 were set as “strong” species (Paatero and Hopke, 2003). The stability of converged $Q$ values was found for all runs. By estimating diagnostic errors, the 3-factor result was found to be the optimal solution, with all factors mapped in 100% of the Bootstrap run, no swap of Displacement and no swapping case of Bootstrap-Displacement. The results of the scaled residuals for the 3-factor outcome were within an acceptable range of ($-3$, 3) (Juntto and Paatero, 1994). The measured and PMF-simulated total particle mass concentrations were significantly correlated ($R^2=0.90$).

3 Results and discussion

3.1 An overview of meteorological conditions

Under the impact of the Asian Summer Monsoon, Lhasa normally experienced a rainy season from May to September (Ding, 2007; Ran et al., 2014). However, in August 2020 when the campaign took place, two periods with distinctly different conditions of water vapor were found, as could be clearly seen from $q$ near the ground measured in urban Lhasa (Fig. 2a). The sharp reduction in $q$ during the daytime of 17 August was an indicator of changing from a humid Period I (averagely 11.1 g kg$^{-1}$) to a relatively dry Period II (averagely 8.1 g kg$^{-1}$). Period I was also characterized as rainy with the precipitation amounting to 78.5 mm and quite cloudy for most of the time, while there was plentiful sunshine and almost no precipitation during Period
II (Fig. 2a). Rainfall mostly occurred in the evening and at night (20:00-08:00), accounting for 84% of the total amount in that month. The two periods differed not much in $T$, which ranged from 8.8–26.8 °C in Period I and from 8.6–25.0 °C in Period II with an average of about 17.0 °C for both periods. In contrast, RH averaged nearly 62% in Period I, but only 45% in Period II (Fig. 2b), corresponding well to the difference in $q$. The temperature shared a similar pattern on each day, except that the maximum value in the daytime apparently lowered on days with an overcast sky. Unlike the diurnal variation of $T$ being a peak in the late afternoon (around 18:00) and a valley in the early morning (around 09:00) without much differences between the two periods, average RH reached a maximum in the early morning of about 79%±11% for period I and 64%±6% for Period II, as well as a minimum in the late afternoon of about 43%±11% for period I and 28%±11% for Period II. Surface winds were largely under 4m s$^{-1}$ and dominated by easterly and westerly winds (Fig. 2c).

Vertical profiles of meteorological parameters observed at the site were categorized into five time periods of the day as give in Table S1, and were presented along $H_{Nor}$, if either a ML or a NBL could be identified, for a straightforward comparison between the two periods (Fig. S2). The amount of moisture was obviously higher in Period I than in Period II for all time periods, being consistent with what had been revealed from continuous surface measurements. The conservative quantities, $\theta$ and $q$, were generally uniform in the ML for daytime profiles, whereas $T$ decreased and RH increased with increasing height, respectively. For profiles collected during 07:00-08:00 and 20:00-23:00, air temperature inversions in the NBL were recognized for several cases. Averagely, there was a slight increase in $\theta$ and decrease in $q$ along with increasing height. Accordingly, wind speed apparently increased with increasing height. A further examination on the probability distributions of wind speed and wind direction indicated that winds were mainly below 4 m s$^{-1}$ both within and above the BL throughout the campaign, also being dominated by easterly and westerly winds as near the ground (Fig. S3). Stronger easterly winds exceeding 4 m s$^{-1}$ occurred above the BL, even up to nearly 7 m s$^{-1}$ in Period II. It should be kept in mind that data points available for calculating time-period averages at some heights above the BL, corresponding to $H_{Nor}$ larger than 0, might make
up only a part of the total number and might thus introduce misleading details. Caution should thereby be taken about statistically drawing conclusions from the characteristics of height-normalized profiles across the BL and at different heights above the top of the BL. The last but not the least, the analysis regarding meteorological parameters, especially winds, was somewhat limited to measurements under a relatively mild condition when the tethered balloon were able to be launched.

![Figure 2](https://doi.org/10.5194/acp-2021-810)

**Figure 2** The time series of (a) precipitation in blue and specific humidity \( q \) in green, (b) temperature \( T \) in red and relative humidity \( \text{RH} \) in black, (c) wind speed and wind direction (in colours), based on hourly data in August from an Automatic Weather Station in urban Lhasa.
3.2 The vertical structure and temporal variability of aerosol profiles

3.2.1 Vertical distributions and diurnal variations of aerosol properties

The evolution of the BL was an important influencing factor in shaping the vertical structure of atmospheric aerosols, as being already pointed out by some previous studies (Ferrero et al., 2010; Rant et al., 2016; Zhang et al., 2017). During the campaign, heights of the shallow NBL were largely below 200 m around sunrise/sunset and at night (Fig. S4). In Period I, \( H_m \) could be determined for most profiles before midday around 14:00, possibly resulting from the development of the ML being suppressed under usually cloudy conditions. In the afternoon, the ML often developed higher than the maximum height reached by the tethered balloon. Unfortunately, intensive observations covering a day from the early morning until the night were unavailable, either because of unobtainable aviation permission or bad weather. Nevertheless, relatively frequent measurements on 12 August (marked by black cross in Fig. S4) revealed, in a more realistic way as compared with what the overall picture could tell, the gradual increase of \( H_m \) in the overcast morning, from below 100 m before sunrise to about 700 m towards noon. The depth of the ML on that day was expected to be far above the flight limit of 500 m in the afternoon, since it turned to be quite sunny and cloudless overhead after midday. As for Period II, the ML should have developed very quickly after sunrise on those sunny days, thus \( H_m \) were often already high above the flight limit when aviation permissions were granted in the late morning and the afternoon.

Normalized by \( H_m \), average vertical profiles of aerosol properties in five time periods of the day were illustrated in Fig. 3, exhibiting a similarity between Period I and II. Profiles of \( N_a \) and particle mass concentrations around sunrise (07:00-08:00) and since evening (20:00-23:00) resembled an exponential decay in the stable NBL, which favored the accumulation of air pollutants near the ground. In the daytime, profiles were characterized by a nearly uniform distribution of aerosols within the ML, except for profiles collected during 08:00-12:00 in Period II with the characteristics of nighttime profiles. This misleading feature was attributed largely to profiles on religious holidays that were frequently encountered in Period II with the impact of
strong emissions from religious burning in the morning, which will be discussed in detail in Sect. 3.3. With the evolution of the ML, stronger dilution effects led to a decline in surface $N_a$ and particle mass concentrations in the daytime, compared to that in the early morning and at night. However, surface $N_a$ and particle mass concentrations were found to be the highest during 08:00-12:00 in both periods. The reasons for this phenomenon might be enhanced emissions from more anthropogenic activities after sunrise but still weak dispersion inside a ML, where vertical convection had not been fully developed. Above the BL, the number of data points used for averaging varied with $H_{Nor}$, depending on $H_m$ and the maximum height of each flight. To avoid being misled by insufficient data, averaged data were only adopted at heights where the data availability exceeded 75%. An evident reduction in $N_a$ and particle mass concentrations above the BL was observed by comparison with that within the BL. For both periods, ratios of PM$_1$ to PM$_{2.5}$ mass concentrations (PM$_1$/PM$_{2.5}$) were relatively uniform within the BL even for NBL-type profiles, despite distinctly different aerosol amount along the normalized height in the NBL. The daytime PM$_1$/PM$_{2.5}$ ratios were relatively larger than ratios in the early morning and at night. Accordingly, $D_e$ was found to be relatively smaller during the day.

Table 1 listed particle parameters in the BL, the RL and the FT. For the BL category, averages within the ML and 10-m averages near the ground for the NBL were adopted. Inside the well mixed layer, the surface level and the ML-averaged level were close to each other. Averaged $N_a$ within the BL was about 338±162 cm$^{-3}$ in Period I and 409±201 cm$^{-3}$ in Period II. The mean PM$_{2.5}$ mass concentration within the BL was 2.7±1.7 μg m$^{-3}$ in Period I, with a fraction of PM$_1$ in PM$_{2.5}$ to be about 81±6%. In Period II, PM$_{2.5}$ mass concentrations within the BL averaged 3.6±2.4 μg m$^{-3}$, with PM$_1$ accounting for approximately 73±9%. The lower level of aerosols in Period I might be ascribed to frequent rainfalls and associated efficient removal processes. Submicron particles took up a smaller proportion in PM$_{2.5}$ during Period II, together with a larger average $D_e$ of 0.35 and a larger average $D_{e,1-2.5\mu m}$ of 1.76, indicating more large particles in Period II than in Period I. Though average $D_{e,<1\mu m}$
in the two periods was comparable with each other, a wider range was found in Period II, showing the large variability in particle size distributions among profiles collected at different times.

Figure 3 Average vertical profiles of (a) particle number concentration \(N_a\), (b) PM_{1.0} mass concentration, (c) \(PM_{2.5}\) mass concentration, (d) the ratio of PM_{1.0} to \(PM_{2.5}\) mass concentration, and (e) the effective diameter \(D_e\) over the whole size range of the POPS, along the normalized height \(H_{Nor}\) during Period I. Profiles for different time periods are shown in different colours, with dots representing average values and error bars representing standard deviations at each of the \(H_{Nor}\). The number of data points used for averaging at each \(H_{Nor}\) was more than about 75% of the number of profiles during each time period, as given in parentheses.

Average vertical profiles measured during Period II are displayed in the same way in (f)-(j).

It was found that average \(PM_{2.5}\) mass concentrations within the BL during this campaign \((3.1\pm2.1 \mu g m^{-3})\) were far below what had been observed near the ground in August 2016 in urban Lhasa, where the high levels often exceeded 20 \(\mu g m^{-3}\) and were sometimes up to more than 40 \(\mu g m^{-3}\), with the overall average to be 11\(\pm2.2 \mu g m^{-3}\) over the size range of 0.14–3 \(\mu m\) (Cui et al., 2018b). The significant difference of particle mass concentrations between the previous and the current study might reflect different emission strengths surrounding the two sites. Being located in the downtown area and adjacent to several temples, the observational site in the previous study was greatly affected by various strong local emissions, such as traffic, residential...
and religious activities-related emissions, while the suburban site in this study was under the influence of much weaker nearby emissions and more likely to represent an average condition over this area. Furthermore, the continuous monitoring of PM$_{2.5}$ mass concentrations from 2013 to 2017 at 6 sites across the city of Lhasa revealed a slight decrease in 2017 (Yin et al., 2019). This was demonstrated to be partly attributed to measures taken to prevent and control air pollution in Lhasa since 2013. Actually, the location of the site in this study was on the same campus of Tibet University as one site (LHASA-XZ) in Yin et al. (2019). Considering that monthly mean PM$_{2.5}$ mass concentration of all 6 sites for August was reported as about 15 μg m$^{-3}$ and the LHASA-XZ site was not the cleanest one among the others, it might suggest an apparent reduction in PM$_{2.5}$ mass concentrations with the continuing implementation of air quality policies.

Table 1 Particle parameters within the BL, in the RL and in the FT for Period I and II, given as average value ± standard deviation and the range in parentheses. The number of profiles for each category is also given in parentheses in the second row.

|                | BL | RL | FT |
|----------------|----|----|----|
|                | Period I (56) | Period II (56) | Period I (0) | Period II (29) | Period I (14) | Period II (10) |
| $N_a$ (cm$^{-3}$) | 338±162 | 409±201 | / | 299±101 | 89±37 | 81±23 |
|                | (77~812) | (95~1051) | / | (123~575) | (47~156) | (44~112) |
| PM$_1$ (μg m$^{-3}$) | 2.1±1.3 | 2.6±1.7 | / | 1.6±0.6 | 0.5±0.2 | 0.5±0.3 |
|                | (0.4~5.5) | (0.5~7.8) | / | (0.6~3.1) | (0.3~1.1) | (0.2~0.9) |
| PM$_{2.5}$ (μg m$^{-3}$) | 2.7±1.7 | 3.6±2.4 | / | 2.0±0.7 | 0.7±0.3 | 0.7±0.4 |
|                | (0.5~7.8) | (0.6~10.9) | / | (0.9~4.0) | (0.3~1.3) | (0.3~1.1) |
| PM$_1$/PM$_{2.5}$ (%) | 81±6 | 73±9 | / | 78±5 | 79±5 | 76±9 |
|                | (62~90) | (48~93) | / | (69~86) | (71~87) | (63~89) |
| $D_e$ (μm) | 0.30±0.06 | 0.35±0.09 | / | 0.31±0.03 | 0.29±0.04 | 0.34±0.07 |
|                | (0.20~0.47) | (0.21~0.61) | / | (0.26~0.37) | (0.24~0.36) | (0.24~0.46) |
| $D_{e<1μm}$ (μm) | 0.23±0.03 | 0.23±0.04 | / | 0.22±0.01 | 0.22±0.01 | 0.23±0.03 |
|                | (0.19~0.31) | (0.18~0.35) | / | (0.20~0.24) | (0.20~0.24) | (0.18~0.27) |
| $D_{e,1-2.5μm}$ (μm) | 1.71±0.15 | 1.76±0.13 | / | 1.72±0.08 | 1.58±0.11 | 1.62±0.18 |
|                | (1.34~2.46) | (1.26~2.00) | / | (1.56~1.87) | (1.40~1.76) | (1.32~1.78) |

Profiles in Period II were mainly collected in the early morning and at night, when elevated aerosol layers were often encountered, though it was hard to tell whether those were only residual layers that could easily form above the NBL after sunset and remain until the BL evolved to an enough height in the next day, or there could also be contributions from
transported plumes. Here, we generally denoted such elevated layers as residual layers (RL). No RL were observed during Period I. In Period II, the average particle number and mass concentrations in the RL were respectively around 75% and 55% of that within the BL. In the FT, average particle number and mass concentrations in the two periods were generally comparable with each other, and were almost less than 25% of the amount within the BL. In general, fractions of PM$_{1}$ in PM$_{2.5}$ within the BL were close to those in the RL and the FT for each period.

Undoubtedly, air quality over the Tibetan Plateau has been increasingly influenced by anthropogenic activities and emissions during the processes of urbanization and economic growth in the past several decades, especially in relatively thickly populated cities like Lhasa. However, the level of $N_a$ over Lhasa was still much lower than what has been observed in more populated and developed regions. Aircraft measurements of particles with the diameter ranging from 0.1 μm to 3.0 μm using a passive cavity aerosol spectrometer probe (PCASP) over the Loess Plateau, which is adjacent to the Tibetan Plateau, revealed that $N_a$ in the BL often exceeded 1000 cm$^{-3}$ and $N_a$ above the BL reached well above 200 cm$^{-3}$ for most flights during the summer campaign 2020 (Cai et al., 2021). Influenced more by anthropogenic emissions, $N_a$ over the Loess Plateau were more than 3 times of those obtained within and above the BL over Lhasa. Though a slightly larger average $D_e$ of around 0.4 μm was observed over the Loess Plateau, no apparent dependence of $D_e$ on altitude was found in both locations. Aircraft measurements that also employed PCASP in non-winter seasons during 2005 and 2006 over Beijing in the polluted North China Plain showed that the surface level of $N_a$ ranged from 1000 cm$^{-3}$ (10th percentile) to 10000 cm$^{-3}$ (90th percentile) and averaged as high as 6600 cm$^{-3}$, more than 10 times of that over Lhasa (Liu et al., 2009). Aerosol profiles measured by a tethered balloon-born OPC over Milan in Italy, which is also located in a river valley (the Po Valley) but in a region much more industrialized and populated than the Lhasa River Valley, exhibited an average $N_a$ of 35±2 cm$^{-3}$ in the diameter range of 0.3–20 μm above the BL in summer (Ferrero et al., 2010). Given the same size range, a lower level of particle number concentrations was found in the FT over Lhasa, with the average $N_a$ to be 12±8 cm$^{-3}$. 
3.2.2 Vertically resolved particle mass size distributions

Particle mass size distributions (PMSD) measured by the POPS were categorized into two periods and averaged for three layers (Fig. 4). In general, size-resolved particle mass concentrations were the highest at all diameters within the BL, while much lower in the FT. Nevertheless, a similarity was found among all PMSDs. Plainly, a distinct mode below around 0.23 μm existed for all layers during both periods, though its relative strength to the rest part of the PMSD considerably differed among different layers and periods. For this mode, more particles were observed towards the lower end of the size range in Period II, possibly implying stronger secondary formation of fine particles favored by the fine weather during that period. Particles larger than 1 μm apparently increased with the diameter for all PMSDs, suggesting another mode in coarse particles. The normalized averages of PMSDs that were merged from the POPS and GRIMM 11-C measurements over the size range of 0.124–32 μm, which were only available for 48 profiles in Period II, did exhibit a mode peaking around 2.5 μm in the FT and around 5 μm in the other two layers (Fig. S5). Particles as large as roughly 15 μm were observed within the BL and in the RL. In contrast, particles with the diameter larger than 5 μm accounted only for a negligible part of the PMSD in the FT. It was also noted, from both the average PMSD (Fig. 4) and the normalized average PMSD (Fig. S5), that there was a third mode ranging over 0.5–0.7 μm within the BL and in the RL, though relatively less pronounced for the later one, whereas no such an apparent mode was observed in the FT. These features revealed that under most circumstances particles in the RL were of the same origin as particles in the BL, whereas particles in the FT should be rarely affected by local anthropogenic emissions near the ground.
Figure 4 Average particle mass size distributions \((dM/d\log D_p)\) for different layers during the two periods as indicated by different markers.

The vertical distribution of PMSDs within the size range of 0.124–32 \(\mu m\) for each profile in Period II was explored in more detail. It was noteworthy that the mode over 0.5–0.7 \(\mu m\) was found to be more distinct on the mornings of religious holidays compared with that at other times. In order to conveniently make comparisons among different cases, the dataset was grouped into subsets of data collected on non-holidays and holidays. In total, there were 22 aerosol profiles collected on non-holidays and 26 on the four religious holidays (19, 20, 26, 28 August). The subset of data from holidays was further divided into five cases, four of which respectively comprised measurements on the morning of each religious holiday. One case, consisting of data collected on holidays except in the morning, was denoted as holidays* to distinguish from the category named holidays that covered all profiles measured on religious holidays. The average profile of \(D_{e,<1\mu m}\) on non-holidays and holidays* were found to be close to each other, with an average of roughly 0.21±0.01 \(\mu m\) and 0.23±0.02 \(\mu m\) through the vertical direction, respectively (Fig. 5a). However, the average vertical distribution of \(D_{e,<1\mu m}\) on the morning of each religious holiday, as
separately displayed in different colors, was significantly different from the categories of non-holidays and holidays*. In the BL, a remarkably larger $D_{e,<1\mu m}$ was observed in the morning on all religious holidays than that on both non-holidays and holidays*. The largest $D_{e,<1\mu m}$ was nearly 0.35 μm on 19 August, the Shakyamuni Buddha Day and also the beginning of Shodun Festival, one of the most ceremonious traditional festivals in Tibet. This could plausibly explain the aforementioned larger range of $D_{e,<1\mu m}$ averaged within the BL in period II than in Period I, since emissions from religious activities on religious holidays enlarged the effective diameter and only one religious holiday (8 August) was encountered in Period I. Above the BL, $D_{e,<1\mu m}$ for the four holiday morning cases generally fell in the range of $D_{e,<1\mu m}$ on non-holidays and holidays, except that $D_{e,<1\mu m}$ on the morning of 19 August was apparently larger than others. As for coarse particles in the size range of 1~10 μm, no obvious difference was observed between non-holidays and holidays (Fig. 5b).

Figure 5 (a) Average vertical profiles of $D_{e,<1\mu m}$ for different cases: solid black dots for non-holidays, magenta circles for religious holidays but excluding samples in the mornings, various colored solid dots for religious holiday mornings. (b) Average vertical profiles of $D_{e,1~10\mu m}$ for non-holidays (black dots) and holidays (red dots). The number of profiles collected during the period of each case was given in the parentheses.
A further examination was taken on average PMSDs within 50 m above the ground for each category, considering that in the NBL or in an early morning ML measurements near the ground would better help elucidate the impact religious activities brought to PMSDs. A marked mode peaking around 0.6 μm and amounting to about 18 μg m\(^{-3}\) was observed in the PMSDs on 19 and 28 August (Fig. 6). Though much lower around 5 μg m\(^{-3}\) on 20 and 26, the peak of this mode was still almost double of that on non-holidays and holidays*. The existence of the accumulation mode over 0.5–0.7 μm on religious holidays was consistent with the findings from surface aerosol measurements in August 2016, and was demonstrated to be characteristic of emissions from incense burning and biomass burning for religious ceremonies (Cui et al., 2018b and references therein). As a widespread religious custom, incense burning was commonly performed in the daily life of local people, not only in temples but also at home. The burning of several types of wood branches and herbs as well as butter lamps in temples was also traditional religious activities. The PMSDs clearly showed that these emissions were largely enhanced on holidays especially in the mornings and formed a much strengthened accumulation mode.
3.3 Potential sources of particles in different layers

Three source factors were identified from the PMF analysis of the PMSDs combined from the POPS and GRIMM 11-C measurements in Period II (Fig. 7a). The first factor (factor 1) revealed a potential source that predominantly contributed to particles smaller than 0.3 μm, probably being associated with local emissions from fossil fuel combustions and/or secondary aerosol formation. The second factor (factor 2) showed a broad peak over about 0.3~0.7 μm, with also a considerable contribution from particles with the diameter extending from 0.7 μm to around 2.5 μm. Considering the apparent enhancement of particle mass in the size range of 0.5~0.7 μm on religious holiday mornings, factor 2 was taken to be mainly representative of aerosols released from religious activities such as burning incense, cypress branches, herbs, and butter lamp (Cui et al., 2018b and references therein). The third factor (factor 3) was almost exclusively composed of particles in the coarse mode (> 1 μm). This factor might be attributed to suspended dust particles from unpaved roads, construction sites, and the surrounding mountains. Indoor particles previously collected at a temple in Lhasa exhibited a bimodal particle mass distribution, with one peak around 0.4~0.7 μm and one peak in the coarse mode around 5 μm (Cui et al., 2018a). It was speculated that factor 3 might also involve a certain contribution from religious burning and/or residential biomass burning. However, concentrations contributed by factor 3 did not increase as expected with the rising concentrations contributed by factor 2 on holiday mornings, when assuming that coarse mode particles observed at the site were partly generated from religious activities (Fig. S6). Also residential burning of biomass including cow dung and plants, as an important source in the past for energy, was negligible nowadays and mostly replaced by a mixture of fossil fuel and renewable energy sources such as solar, wind, geothermal and hydroelectric power.
On average, particles in the BL and the RL shared a similar makeup of potential sources (Fig. 7b), suggesting that particles in the two layers were probably of the same origin. The contribution by suspended dust particles took up a portion of more than 50%, while the contribution by religious burning was comparable with the contribution by combustion related or secondary formation source with a fraction of roughly 20%~25%. Particularly, the results for the BL were separated into one part on holiday mornings and the other under other conditions, namely, non-holidays and holidays* as defined above, since particles in the BL was mostly influenced by emissions near the ground. It was found that on non-holidays and holidays* the makeup of potential sources was close to that during the whole Period II, while religious burning contributed about 45% on holiday mornings and exceeded the other two factors. However, the relative relationship of the two factors other than religious burning...
was similar under both conditions, with the contribution by combustion/secondary formation nearly 40\% of the contribution by dust suspension, suggesting these two sources stayed relatively stable during the campaign. An examination on contributions from the three factors at different time periods on non-holidays revealed the predominant contribution by dust suspension and no obvious diurnal variation of the makeup of potential sources was found. Unlike in the BL and the RL where the significant importance of coarse mode particles was found, the contribution by large particles that were possibly associated with dust suspension was below 10\% in the FT. Factor 1 and 2 contributed approximately 40\% and 50\% in the FT. Although factor 2 was identified as a religious burning factor for the BL and the RL, this might not be the case in the FT, as the contribution from factor 2 did not differ so much on holiday mornings and at other times. Given the FT was generally much less influenced by emissions near the ground, factor 1 and 2 may be related more to secondary aerosol formation and the aging of existed particles. Measurements of aerosol chemical compositions along the vertical direction in the future might help entangle this issue.

### 4 Summary

In this study, vertical profiles of particle size distributions within 1 km were measured by a POPS and a GRIMM 11-C attached to a tethered balloon in summer 2020 at a suburban site in Lhasa. The variability in the vertical structure and temporal features of parameters such as aerosol number and mass concentrations, the effective diameter and particle mass size distributions were examined. Possible sources for aerosols in different layers were investigated by the PMF analysis.

The vertical distribution of aerosol properties was found to be largely governed by the diurnal variation of the boundary layer. Generally speaking, aerosols uniformly distributed in the daytime BL and sharply declined in the NBL. Usually average particle number and mass concentrations in the FT were less than 25\% of the amount in the BL. For the humid Period I under the influence the Asian Summer Monsoon, a lower level of aerosols and also less large particles in the BL were found than
that in the relatively dry Period II, possibly due to frequent rainfalls and associated efficient removal by wind. More emissions from religious activities in Period II might also contribute to the differences between the two periods. In contrast, both the total amount and the size distribution of particle mass in the FT for the two periods were comparable. Residual layers with elevated aerosol were often encountered in Period II, with particle mass concentrations averagely around 55% of that in the BL. The PMSDs in the RL shared a similar pattern as that in the BL, suggesting they were probably of the same origin. The source apportionment analysis revealed that the factor associated with suspended dust in the coarse mode contributed more than 50% for the BL and the RL. However, a distinct peak over 0.5–0.7 μm was observed on religious holiday mornings, the same time periods when the contribution by the identified religious burning factor with a broad peak over about 0.3–0.7 μm was found to exceed the contribution by dust suspension. In the FT, submicron particles dominated and the contribution by coarse mode particles was below 10%. More investigations based on ground-level long-term monitoring and vertical measurements of aerosol chemical and physical properties will help better identify the origin and evolution processes of aerosols in this area.

Data availability

All data presented in this paper can be accessed by contacting corresponding authors, Zhaoze Deng (dengzz@mail.iap.ac.cn) and Liang Ran (shirleyrl@mail.iap.ac.cn).

Author contribution

JB and LR proposed the study. LR, ZD and ZB designed and conducted the field campaign. YL participated in the field campaign. LR and ZD processed the data. JL and YW performed the Positive Matrix Factorization modelling. DZ provided surface meteorological data. LR visualized the data and wrote the manuscript. ZD and YW participated in several discussions and provided valuable suggestions. All authors reviewed the manuscript carefully.
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References

Amato, F., Pandolfi, M., Escrig. A., Querol, X., Alastuey, A., Pey, J., Perez, N., and Hopke, P. K.: Quantifying road dust resuspension in urban environment by Multilinear Engine: A comparison with PMF2, Atmos. Environ., 43, 2770-2780, doi:10.1016/j.atmosenv.2009.02.039, 2009.

Atkinson, R.: Atmospheric chemistry of VOCs and NOx, Atmos. Environ., 34, 2063-2101, doi:10.1016/S1352-2310(99)00460-4, 2000.

Bai, Y. B., Shi, G. Y., Tamura, K., Shibata, T., Iwasaka, Y., Shin’ichi, K., and Tamio, T.: Aerosol optical properties derived from simultaneous sunphotometer and aureolemeter measurements in Lhasa, J. Environ. Sci., 12, 439-443, 2000.

Brown, S. G., Eberly, S., Paatero, P., and Norris, G. A.: Methods for estimating uncertainty in PMF solutions: Examples with ambient air and water quality data and guidance on reporting PMF results, Sci. Tot. Environ., 518-519, 626-635, doi:10.1016/j.scitotenv.2015.01.022, 2015.

Cai, Z. X., Li, Z. Q., L, P. R., Li, J. X., Sun, H. P., Yang, Y. M., Gao, X., Ren, G., Ren, R. M., and Wei, J.: Vertical distributions of aerosol microphysical and optical properties based on aircraft measurements made over the Loess Plateau in China, Atmos. Environ., in press, doi:10.1016/j.atmosenv.2021.118888, 2021.
Chen, P. F., Kang, S. C., Li, C. L., Li, Q. L., Yan, F. P., Guo, J. M., Ji, Z. M., Zhang, Q. G., Hu, Z. F., Tripathee, L., and Sillanpää, M.: Source apportionment and risk assessment of atmospheric polycyclic aromatic hydrocarbons in Lhasa, Tibet, China, Aerosol Air Qual. Res., 18, 1294-1304, doi:10.4209/aaqr.2017.12.0603, 2018.

Chiang, J. C. H., Kong, W., Wu, C. H., and Battisti, D. S.: Origins of East Asian Summer Monsoon seasonality, J. Climate, 33, 7945-7965, doi: 10.1175/JCLI-D-19-0888.1, 2020.

Cong, Z. Y., Kang, S. C., Luo, C. L., Li, Q., Huang, J., Gao, S. P., and Li, X. D.: Trace elements and lead isotopic composition of PM$_{10}$ in Lhasa, Tibet, Atmos. Environ., 45, 6210-6215, doi:10.1016/j.atmosenv.2011.07.060, 2011.

Cui, L. L., Duo, B., Zhang, F., Li, C. L., Fu, H. B., and Chen, J. M.: Physiochemical characteristics of aerosol particles collected from the Jokhang Temple indoors and the implication to human exposure, Environ. Pollut., 236, 992-1003, doi: 10.1016/j.envpol.2017.10.107, 2018a.

Cui, Y. Y., Liu, S., Bai, Z. X., Bian, J. C., Li, D., Fan, K. Y., McKeen, S. A., Watts, L. A., Ciciora, S. J., and Gao, R. S.: Religious burning as a potential major source of atmospheric fine aerosols in summertime Lhasa on the Tibetan Plateau, Atmos. Environ., 181, 186-191, doi:10.1016/j.atmosenv.2018.03.025, 2018b.

Ding, Y. H.: The variability of the Asian Summer Monsoon, J. Meteor. Soc. Japan, 85B, 21-54, doi:10.2151/jmsj.85B.21, 2007.

Duo, B., Zhang, Y. C., Kong, L. D., Fu, H. B., Hu, Y. J., Chen, J. M., Li, L., and Qiong, A.: Individual particle analysis of aerosols collected at Lhasa City in the Tibetan Plateau, J. Environ. Sci., 29, 165-177, doi:10.1016/j.jes.2014.07.032, 2015.

Duo, B., Cui, L. L., Wang, Z. Z., Li, R., Zhang, L. W., Fu, H. B., Chen, J. M., Zhang, H. F., and Qiong, A.: Observations of atmospheric pollutants at Lhasa during 2014-2015: Pollution status and the influence of meteorological factors, J. Environ. Sci., 63, 28-42, doi:10.1016/j.jes.2017.03.010, 2018.
Ferrero, L., Perrone, M. G., Petraccone, S., Sangiorgi, G., Ferrini, B. S., Lo Porto, C., Lazzati, Z., Cocchi, D., Bruno, F., Greco, F., Riccio, A., and Bolzacchini, E.: Vertically-resolved particle size distribution within and above the mixing layer over the Milan metropolitan area, Atmos. Chem. Phys., 10, 3915-3932, doi:10.5194/acp-10-3915-2010, 2010.

Ferrero, L., Castelli, M., Ferrini, B. S., Moscatelli, M., Perrone, M. G., Sangiorgi, G., D’Angelo, L., Rovelli, G., Moroni, B., Scardazza, F., Mocnik, G., Bolzacchini, E., Petitta, M., and Cappelletti, D.: Impact of black carbon aerosol over Italian basin valleys: high-resolution measurements along vertical profiles, radiative forcing and heating rate, Atmos. Chem. Phys., 14, 9641-9664, doi:10.5194/acp-14-9641-2014, 2014.

Gao, R. S., Telg, H., McLaughlin, R. J., Ciciora, S. J., Watts, L. A., Richardson, M. S., Schwarz, J. P., Perring, A. E., Thornberry, T. D., Rollins, A. W., Markovic, M. Z., Bates, T. S., Johnson, J. E., and Fahey, D. W.: A light-weight, high-sensitivity particle spectrometer for PM$_{2.5}$ aerosol measurements, Aerosol Sci. Technol., 50, 88-99, doi:10.1080/02786826.2015.1131809, 2016.

Gong, P., Wang, X. P., and Yao. T. D.: Ambient distribution of particulate- and gas-phase n-alkanes and polycyclic aromatic hydrocarbons in the Tibetan Plateau, Environ. Earth Sci., 64, 1703-1711, doi:10.1007/s12665-011-0974-3, 2011.

He, C., Wang, Z. Q., Zhou, T. J., and Li, T.: Enhanced latent heating over the Tibetan Plateau as a key to the enhanced East Asian Summer Monsoon circulation under a warming climate, J. Climate, 32, 3373-3388, doi:10.1175/JCLI-D-18-0427.1, 2019.

Jarvis, A., Reuter, H. I., Nelson, A., and Guevara, E.: Hole-filled seamless SRTM data V4, International Centre for Tropical Agriculture (CIAT), available from http://srtm.csi.cgiar.org, 2008.

Juntto, S. and Paatero, P.: Analysis of daily precipitation data by positive matrix factorization, Environmetrics, 5, 127-144, doi:10.1002/env.3170050204, 1994.
Kim, S. W., Yoon, S. C., Won, J. G., and Choi, S. C.: Ground-based remote sensing measurements of aerosol and ozone in an urban area: a case study of mixing height evolution and its effect on ground-level ozone concentrations, Atmos. Environ., 41, 7069-7081, doi:10.1016/j.atmosenv.2007.04.063, 2007.

Li, C. L., Han, X. W., Kang, S. C., Yan, F. P., Chen, P. F., Hu, Z. F., Yang, J. H., Ciren, D. J., Gao, S. P., Sillanpää, M., Han, Y. M., Cui, Y. Y., Liu, S., and Smith, K. R.: Heavy near-surface PM$_{2.5}$ pollution in Lhasa, China during a relatively static winter period, Chemosphere, 214, 314-318, doi:10.1016/j.chemosphere.2018.09.135, 2019.

Lin, W., Zhu, T., Song, Y., Zou, H., Tang, M., Tang, X., and Hu, J.: Photolysis of surface O$_3$ and production potential of OH radicals in the atmosphere over the Tibetan Plateau, J. Geophys. Res., 113, D02309, doi:10.1029/2007JD008831, 2008.

Liu, J. W., Li, J., Lin, T., Liu, D., Xu, Y., Chaemfa, C., Qi, S. H., Liu, F. B., and Zhang, G.: Diurnal and nocturnal variations of PAHs in the Lhasa atmosphere, Tibetan Plateau: Implication for local sources and the impact of atmospheric degradation processing, Atmos. Res., 124, 34-43, doi:10.1016/j.atmosres.2012.12.016, 2013.

Liu, P. F., Zhao, C. S., Zhang, Q., Deng, Z. Z., Huang, M. Y., Ma, X. C., and Tie, X. X.: Aircraft study of aerosol vertical distributions over Beijing and their optical properties, Tellus, 61B, 756-767, doi:10.1111/j.1600-0889.2009.00440.x, 2009.

Paatero, P.: Least squares formulation of robust non-negative factor analysis, Chemom. Intell. Lab. Syst., 37, 23-35, doi:10.1016/S0169-7439(96)00044-5, 1997.

Paatero, P. and Hopke, P. K.: Discarding or downweighting high-noise variables in factor analytic models, Anal. Chim. Acta, 490, 277-289, doi:10.1016/S0003-2670(02)01643-4, 2003.

Paatero, P. and Tapper, U.: Positive matrix factorization: A non-negative factor model with optimal utilization of error estimates of data values, Environmetrics, 5, 111-126, doi:10.1002/env.3170050203, 1994.

Ran, L., Lin, W., Deji, Y., La, B., Tsering, P., Xu, X., and Wang, W.: Surface gas pollutants in Lhasa, a highland city of Tibet - current levels and pollution implications, Atmos. Chem. Phys., 14, 10721-10730, doi:10.5194/acp-14-10721-2014, 2014.
Ran, L., Deng, Z. Z., Xu, X. B., Yan, P., Lin, W. L., Wang, Y., Tian, P., Wang, P. C., Pan, W. L., and Lu, D. R.: Vertical profiles of black carbon measured by a micro-aethalometer in summer in the North China Plain, Atmos. Chem. Phys., 16, 10441-10454, doi:10.5194/acp-16-10441-2016, 2016.

Seibert, P., Beyrich, F., Gryning, S. E., Joffre, S., Rasmussen, A., and Tercier, P.: Review and intercomparison of operational methods for the determination of the mixing height, Atmos. Environ., 34, 1001-1027, doi:10.1016/S1352-2310(99)00349-0, 2000.

Sloane, C. S., Watson, J., Chow, J., Pritchett, L., and Richards, L. W.: Size-segregated fine particle measurements by chemical species and their impact on visibility impairment in Denver, Atmos. Environ., 25, 1013-1024, doi:10.1016/0960-1686(91)90143-U, 1991.

Tao, J., Zhang, L. M., Cao, J. J., Hsu, S.-C., Xia, X. A., Zhang, Z. S., Lin, Z. J., Cheng, T. T., and Zhang, R. J.: Characterization and source apportionment of aerosol light extinction in Chengdu, southwest China, Atmos. Environ., 95, 552-562, doi:10.1016/j.atmosenv.2014.07.017, 2014.

Tobo, Y., Iwasaka, Y., Shi, G. Y., Kim, Y. S., Ohashi, T., Tamura, K., and Zhang, D. Z.: Balloon-borne observations of high aerosol concentrations near the summertime tropopause over the Tibetan Plateau, Atmos. Res., 84, 233-241, doi:10.1016/j.atmosres.2006.08.003, 2007.

Wan, X., Kang, S. C., Xin, J. Y., Liu, B., Wen, T. X., Wang, P. L., Wang, Y. S., and Cong, Z. Y.: Chemical composition of size-segregated aerosols in Lhasa city, Tibetan Plateau, Atmos. Res., 174-175, 142-150, doi: 10.1016/j.atmosres.2016.02.005, 2016.

Wei, N. N., Ma, C. L., Liu, J. W., Wang, G. H., Liu, W., Zhuoga, D. Q., Xiao, D. T., and Yao, J.: Size-segregated characteristics of carbonaceous aerosols during the monsoon and non-monsoon seasons in Lhasa in the Tibetan Plateau, Atmosphere, 10, 157, doi:10.3390/atmos10030157, 2019a.
Wei, N. N., Xu, Z. Y., Liu, J. W., Wang, G. H., Liu, W., Zhuoga, D. Q., Xiao, D. T., and Yao, J.: Characteristics of size distributions and sources of water-soluble ions in Lhasa during monsoon and non-monsoon seasons, J. Environ. Sci., 82, 155-168, doi:10.1016/j.jes.2019.02.017, 2019b.

Yin, X. F., de Foy, B., Wu, K. P., Feng, C., Kang, S. C., and Zhang, Q. G.: Gaseous and particulate pollutants in Lhasa, Tibet during 2013-2017: Spatial variability, temporal variations and implications, Environ. Pollut., 253, 68-77, doi:10.1016/j.envpol.2019.06.113, 2019.

Zhang, D. Z., Iwasaka, Y., and Shi, G. Y.: Soot particles and their impacts on the mass cycle in the Tibetan atmosphere, Atmos. Environ., 35, 5883-5894, doi:10.1016/S1352-2310(01)00391-0, 2001a.

Zhang, K., Wang, D. F., Bian, Q. G., Duan, Y. S., Zhao, M. F., Fei, D. N., Xiu, G. L., and Fu, Q. Y.: Tethered balloon-based particle number concentration, and size distribution vertical profiles within the lower troposphere of Shanghai, Atmos. Environ., 154, 141-150, doi:10.1016/j.atmosenv.2017.01.025, 2017.

Zhang, X. Y., Arimoto, R., Cao, J. J., An, Z. S., and Wang, D.: Atmospheric dust aerosol over the Tibetan Plateau, J. Geophys. Res., 106, 18471-18476, doi:10.1029/2000JD900672, 2001b.

Zhang, Y. L., Li, B. Y., Liu, L. S., Zheng, D.: Redetermine the region and boundaries of Tibetan Plateau, Geograph. Res., 40, 1543-1553, doi:10.11821/dlyj020210138, 2021. (in Chinese)

Zhu, C. S., Cao, J. J., Hu, T. F., Shen, Z. X., Tie, X. X., Huang H., Wang, Q. Y., Huang, R. J., Zhao, Z. Z., Močnik, G., and Hansen, A. D. A.: Spectral dependence of aerosol light absorption at an urban and a remote site over the Tibetan Plateau, Sci. Total Environ., 590-591, 14-21, doi:10.1016/j.scitotenv.2017.03.057, 2017.

Zhu, J., Xia, X. A., Che, H. Z., Wang, J., Cong, Z. Y., Zhao, T. L., Kang, S. C. Zhang, X. L., Yu, X. N., and Zhang, Y. L.: Spatiotemporal variation of aerosol and potential long-range transport impact over the Tibetan Plateau, China, Atmos. Chem. Phys., 19, 14637-14656, doi:10.5194/acp-19-14637-2019, 2019.