Measurement report: Dust and anthropogenic aerosols’ vertical distributions over northern China dense aerosols gathered at the top of the mixing layer

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Abstract. Over the past decades, northern China has been suffering from persistent air pollution caused by both fine and coarse atmospheric particles. Although there are plenty of theoretical and observational studies on aerosols in northern China, most of them only consider total aerosol concentrations and focus on heavy pollution episodes; the long-term vertical distributions of dust (coarse) and anthropogenic aerosols (fine) and their relationships with the mixing layer height (MLH) have not been revealed. In this study, the dust and anthropogenic aerosols’ mass concentration and the MLH were retrieved by polarization Raman lidar over Beijing from May 2019 to February 2022. We found that large amounts of anthropogenic aerosols accumulate at the top of the mixing layer, which is most noticeable in summer, with monthly mean mass concentration up to 57 µg m⁻³. It is mainly influenced by the southward transport in the upper air, where the atmosphere is relatively stable and moist, favoring hygroscopic growth of particles. Dust mass concentration is discontinuous in the vertical direction, not only on the ground but also in lofted layers that reach up to several kilometers. The heights of these lofted dust layers exhibited apparent seasonal dependence, with the height of the main dust layer gradually ascending from 1.1 km to about 2.5 km from April to June and below 3 km from October to December. In addition, there is a significant negative correlation between bottom anthropogenic aerosols’ mass concentration and the MLH, and an inverse function fit is more suitable to characterize this relationship, while the relationship between bottom
dust mass concentration and the MLH is insignificant. These results will enhance our understanding of the sophisticated interactions between dust and anthropogenic aerosols, the MLH, and regional transport in northern China. It will also help to refine atmospheric chemistry models and improve surface prediction capabilities.

1 Introduction

For the past decades, atmospheric particle pollution has attracted a great deal of public concern in China, because it can decrease visibility and have substantial adverse effects on human health (Liang et al., 2022). Also, coarse and fine particles considerably affect the Earth’s climate through direct and indirect effects of aerosols, and there are considerable differences in the effects of the aerosol types and particles’ vertical distributions on the Earth’s climate (Su et al., 2020; Liu et al., 2022a, b, 2021a). Beijing used to be one of the most polluted cities in China, suffering from persistent atmospheric particulate pollution caused by atmospheric fine particles (anthropogenic aerosols) and coarse particles (Asian dust) (Gui et al., 2022; Zheng et al., 2015; Song et al., 2023; Huang et al., 2014; Xiao et al., 2023). Beijing is surrounded by the Yanshan and Taihang to the north and west, respectively, and the flat North China Plain (NCP) to the south, where atmospheric dispersion is relatively poor. Multiple factors contribute to severe air pollution in Beijing; strong anthropogenic emissions due to rapid urbanization and industrialization are the crucial reasons (Su et al., 2020). In addition, the influence of the Mongolian cyclone, which predisposes East Asian dust to be lifted into the free troposphere (FT) and transported over long distances to the east, also affects Beijing’s air quality (Gui et al., 2022; Sun et al., 2001; Xiao et al., 2023). Therefore, the observed aerosol layer in Beijing is usually a mixture of multiple types (natural and anthropogenic).

Previous studies have also shown that meteorological conditions are the key factors controlling air pollution in Beijing (Zhong et al., 2019, 2018; Guo et al., 2014), with weak southerly winds, high relative humidity (RH) levels, a shallow mixing layer height (MLH), and stable atmospheric stratification associated with severe anthropogenic air pollution, while strong northwest wind is closely related to dust aerosol pollution (Wang et al., 2021c, 2020). Among these meteorological factors, the MLH is one of the most critical, as its depth substantially determines the total volume of pollutant dispersion and mixing (Miao et al., 2018). The MLH affects surface aerosol concentrations, which are associated with vertical mixing of aerosols, and can impact the dilution of surface air pollutants through various interaction and feedback mechanisms. The MLH can also significantly affect the aerosol vertical distributions, as most air pollutants tend to accumulate within the mixing layer (ML) (Su et al., 2020). The development of the MLH also accelerates the hygroscopic growth of fine particles within the ML, which is conducive to enhance the surface air pollution (Cheng et al., 2016; Zhang et al., 2023a; Tang et al., 2016). On the other hand, aerosols also have significant radiative feedbacks to the MLH, surface dimming and upper-level heating; the magnitude and sign of the radiative feedback depends on the aerosol type and distribution height of aerosols (Huang et al., 2018; Ding et al., 2016). In a previous comprehensive review, Li et al. (2017) presented sufficient evidence for aerosol and MLH interactions and described their determinants. Thus, in order to more accurately estimate the formation, growth, and dissipation of air pollution and its interactions with the MLH, it is necessary to observe the vertical distribution of aerosol type and its optical properties at high spatial and temporal resolution.

Although field surface observations or column measurements provide detailed optical, chemical, and microphysical characteristics of particles, they do not capture vertical distribution characteristics of aerosol types and their optical properties (Holben et al., 1998). Plenty of numerical simulations and ground-based remote sensing or meteorological radiosonde techniques have been used to acquire aerosol vertical distributions over a short time in Beijing (Zhong et al., 2019; Guo et al., 2016, 2021; Tang et al., 2016; Su et al., 2020; Liu et al., 2021b, 2023), primarily focusing on severe air pollution episodes, which emphasizes the scarcity of long-term continuous measurements with high spatial and temporal resolution. Moreover, although there are more theoretical and observational studies on MLH characteristics and their correlations with surface aerosols (Zhong et al., 2019, 2018; Guo et al., 2016, 2021; Miao et al., 2015, 2018; Tao et al., 2014; Su et al., 2020), most of them only consider the total aerosol concentrations, and the vertical structure and MLH relationships for coarse particles (dust) and fine particles (anthropogenic aerosols) have not been revealed. This has led to a lack of knowledge about the different aerosol types’ vertical distribution and their relationship with the MLH. This lack of knowledge hinders the understanding of the long-range transport, mixing and stratification processes of aerosols, leading to uncertainties in the assessment of the aerosol climatic effects and meteorological chemistry models. Therefore, a comprehensive investigation of the dust and anthropogenic aerosols’ vertical distributions is urgently needed to facilitate mixing processes of dust with anthropogenic aerosols, as well as quantitative studies of the environmental contributions of different aerosol types.

The PRL (polarization Raman lidar) technique provides a reliable tool to explore the vertical distribution characteristics of dust and anthropogenic aerosols with high spatial
and temporal resolution (Ansmann et al., 2012; Tesche et al., 2011a, b, 2009a, b). Moreover, the PRL is also suitable for retrieving the evolution of the MLH (Sicard et al., 2006; Flamant et al., 1997). In this study, we summarize nearly 3 years of PRL observations in Beijing from May 2019 to February 2022. The annual cycle of dust and anthropogenic aerosols’ vertical distributions and their relationship with the MLH in Beijing are clarified. The results of the PRL measurements presented here can help to fill gaps in the knowledge of the different aerosol types’ vertical distribution and their relationship with the MLH, where limited long-term PRL observations are available. In addition, our observations can also establish a dataset of aerosol optical parameters such as the particle depolarization ratio (PDR) and the lidar ratio (LR) in megacities in northern China. The data can be used to support basic data analysis for spaceborne lidar missions such as CALIPSO (Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations) (Winker et al., 2009), ADIN (Atmospheric LAser Doppler IInstrument) (Witschas et al., 2020), and ATLID (Atmospheric Lidar) (Illingworth et al., 2015), upgrading the accuracy of regional terrestrial and global satellite lidar inversions.

The objectives of our study are to (1) present a long-term dataset of aerosol LR, PDR and aerosol types in northern China, (2) elucidate the annual cycle of the dust and anthropogenic aerosols’ vertical distributions in Beijing, and (3) reveal the relationship between the MLH and the dust and anthropogenic aerosols’ vertical distributions. These results will help improve our understanding of the complex interactions between dust and anthropogenic aerosols, the MLH, and regional transport over northern China. They will also help refine atmospheric chemistry models and improve surface air pollution monitoring and prediction capabilities.

2 Materials and methodology

In this section, we specify the instrument, materials, and methods employed throughout the study. Section 2.1 introduces the PRL system, Sect. 2.2–2.5 describes the auxiliary data, and the polarization lidar photometer networking (POLIPHON) method for retrieving dust and anthropogenic aerosols’ mass concentration is described in Sect. 2.6. Aerosol Robotic Network (AERONET) aerosol optical parameters were the input parameters of the POLIPHON method, and surface PM$_{10}$ and PM$_{2.5}$ mass concentrations are used for validation of POLIPHON method results. The Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model and reanalysis data are used for auxiliary analysis.

2.1 Polarization Raman lidar

PRL was deployed at the Beijing Meteorological Observation Center (BMOC; 39.80°N, 116.47°E) from 22 May 2019 and 20 February 2022 (Fig. 1); it is close to Beijing’s 5th Ring Road in the south, with dense traffic and high traffic flow. The PRL was placed in an air-conditioned room equipped with an uninterruptible power supply (UPS) and dehumidifier to detect aerosols over Beijing in continuous mode through a roof skylight, with a temporal resolution of half an hour, approximately 7 min of data collection, and a 23 min interval. The PRL receive system receives 532 nm perpendicular and parallel polarization backscattered signal, 355 nm Mie backscattered signal, and 387 nm nitrogen Raman backscattered signal in both analog and photon counting modes, with a spatial resolution of 7.5 m. More detailed information about the PRL system can be found in our previous study (Wang et al., 2020).

PRL can independently measure the extinction coefficient (EXT$_{355}$) and backscatter coefficient (BAC$_{355}$) at 355 nm to determine the LR at 355 nm (LR$_{355}$). Since the signal-to-noise (SNR) ratio of the Raman channel is quite low, we used a 3 h average PRL backscatter signal profile to retrieve LR$_{355}$ at night (18:00–06:00 local time) and a 12 h average PRL backscatter signal profile to retrieve LR$_{355}$ during the day. In addition, a 4-point spatial accumulation followed by a 15-point spatial smoothing was applied on the average PRL backscatter signal profile. Also, an overlap function (Wandinger and Ansmann, 2002) was employed to determine the LR$_{355}$ for the lower layers. The uncertainty of the overlap correction is too high below 0.25 km; data below 0.25 km were not used for subsequent analysis. The elastic backscatter signal was used to retrieve the EXT$_{532}$ (Fernald, 1977)}
From January 2018 to January 2019, we conducted a study on the aerosol classification of the NCP in farmland about 120 km north of the BMOC (Wang et al., 2021c). We used particle backward trajectories and PRL observations to summarize the aerosol optical parameters in the NCP, which is mainly influenced by two aerosol types; natural dust from the northwest prevailed in spring (March, April, May) and autumn (September, October, November), and anthropogenic aerosols from the NCP were mainly concentrated in summer (June, July, August) and winter (December, January, February). The PDR was found to be the most relevant parameter for aerosol type, and it was used to classify the aerosols. When the PDR is greater than 23 %, it is classified as dust aerosol, and anthropogenic aerosols are determined when the PDR is less than 9 %. When the PDR was between 9 % and 23 %, it was a mixture of dust and anthropogenic aerosols (also called polluted dust). The PRL aerosol classification results are in good agreement with CALIPSO aerosol classification results. The reason the PDR can be used to distinguish between pure dust and polluted dust is that dust mixes with smoke or other anthropogenic aerosols during transport, making the mixed aerosol particles more spherical and resulting in a smaller PDR. The classification of aerosols by the PDR is also widely done in other cities in China (Zhang et al., 2019b; Huang et al., 2015). Therefore, this research still uses the above PDR threshold to classify aerosols.

2.2 AERONET aerosol optical parameters

The sun-photometer-derived aerosol optical parameters were acquired from AERONET (https://aeronet.gsfc.nasa.gov/, last access: 14 April 2023) (Holben et al., 1998; Che et al., 2015, 2019), and data from four sites were collected, namely “Beijing” (39.97° N, 116.38° E), “Beijing-CAMS” (39.93° N, 116.32° E), “Beijing_PKU” (39.99° N, 116.31° E), and “XiangHe” (39.75° N, 116.96° E). The four AERONET sites around Beijing are shown in Fig. 1. In our study, AOD at 500 nm is utilized to evaluate the representativeness of PRL measurements, and coarse- and fine-mode AOD at 500 nm and coarse- and fine-mode volume concentrations are employed to convert EXT profiles to mass concentration profiles. The monthly average AOD at 500 nm and the conversion factor were obtained by analyzing the sun-photometer-derived aerosol optical parameters around Beijing from May 2019 to February 2022.

2.3 Surface PM$_{10}$ and PM$_{2.5}$ mass concentration

The surface PM$_{10}$ and PM$_{2.5}$ mass concentrations from 22 May 2019 and 20 February 2022 were downloaded from the Beijing Municipal Ecological and Environmental Monitoring Center (http://beijingair.sinaapp.com/, last access: 14 April 2023). We used the closest observation site to the BMOC (observation number 1005A; 39.97° N, 116.47° E), about 15 km north of the BMOC. The surface PM$_{10}$ and PM$_{2.5}$ mass concentrations from 22 May 2019 and 20 February 2022 were downloaded from the Beijing Municipal Ecological and Environmental Monitoring Center. The surface PM$_{10}$ and PM$_{2.5}$ mass concentrations from 22 May 2019 and 20 February 2022 were downloaded from the Beijing Municipal Ecological and Environmental Monitoring Center.
PM$_{2.5}$ mass concentrations were screened before comparison. Firstly, outliers of PM$_{2.5}$ greater than PM$_{10}$ were discarded. Then, PM$_{10}$ and PM$_{2.5}$ mass concentrations were filtered using the Z score (Barrero et al., 2015). This was achieved by comparing each concentration to the variability of its adjacent points and time series. The filter criteria of the Z score must satisfy three conditions at the same time: (1) the absolute Z score is greater than 4, (2) the increment of the previous value is greater than 9, and (3) the ratio of the value to its central third-order moving average was greater than 2.

2.4 HYSPLIT backward trajectory

The air mass backward trajectories were calculated by the HYSPLIT model, developed by the National Oceanic and Atmospheric Administration (NOAA) and the Global Data Assimilation System (GDAS 1°) (http://www.ready.noaa.gov, last access: 14 April 2023) (Draxler and Hess, 1998). Considering the significant differences in aerosol mass concentration and types distribution at different altitudes, three trajectory arrival heights were set, 0.1, 2, and 4 km. Hourly, 2d back trajectories were generated by TrajStat software, which employed HYSPLIT for trajectory calculation. Then, the backward trajectories were clustered from two to four classification directions utilizing MeteoInfo software (http://www.meteothink.org/downloads/index.html, last access: 14 April 2023); detailed information can be found in Wang et al. (2009).

2.5 ERA5 and MERRA reanalysis dataset

In order to filter dust and anthropogenic aerosols’ mass concentration profiles at high-RH conditions (> 85%), and to investigate the reliability of the PRL-derived MLH, variables such as RH and boundary layer height (BLH) are needed from the ERA5 reanalysis data of the European Centre for Medium-Range Weather Forecasts (ECMWF). In recent years, several studies have assessed the accuracy of the ERA5 reanalysis data provided by ECMWF based on radiosonde. For example, Luo et al. (2020) found that the mean RH difference below 500 hPa between ERA5 and radiosondes was within 10%. Guo et al. (2021) found that ERA5 BLH was optimal with the radiosonde BLH by comparing BLH using radiosonde data and various reanalysis data such as ERA5 and MERRA-2 (Modern-Era Retrospective Analysis for Research and Applications, Version 2). The above results show that the RH and BLH of ERA5 reanalysis data have excellent accuracy and have been widely used in various studies. The ERA5 RH profile employed in this paper is divided into 15 layers below 500 hPa, with a temporal and spatial resolution of 1 h and 0.25° × 0.25°, respectively. To ensure spatial consistency, the RH and BLH data closest to the PRL measurement site were used. In addition, the aerosol optical property data obtained from MERRA-2, including AOD, dust AOD, and black carbon AOD, were also employed in this study, provided by the National Aeronautics and Space Administration (NASA) and the Global Modeling and Assimilation Office (GMAO) (Gelaro et al., 2017).

2.6 Polarization lidar photometer networking (POLIPHON) method

In this study, the POLIPHON method was applied to retrieve dust and anthropogenic aerosols’ mass concentration (Sugimoto et al., 2003; Shimizu, 2004; Tesche et al., 2009a, 2011b; Ansmann et al., 2012, 2019). Assuming that dust and anthropogenic aerosols are externally mixed, there are two steps to obtain dust and anthropogenic aerosols’ mass concentration with the POLIPHON method. In the first step, the contributions of dust and anthropogenic aerosols to the total BAC$_{532}$ are separated. Then, EXT$_{532}$ of dust and anthropogenic aerosols is converted into mass concentration (Ansmann et al., 2012; Mamouri and Ansmann, 2014). The monthly average conversion factors of dust and anthropogenic aerosols are presented in Fig. 2a. The detailed derivation of POLIPHON method and relevant input parameters involved in the POLIPHON method are summarized in the Supplement and Table S1, respectively.

Uncertainties in POLIPHON method are discussed in detail in previous studies (Freudenthaler et al., 2009; Tesche et al., 2009b, 2011a, b; Mamouri and Ansmann, 2014; Mamouri et al., 2015; Wang et al., 2021b; Ansmann et al., 2011; Bravo-Aranda et al., 2015), mainly consisting of two components: the input empirical parameters in Table S1 and the uncertainties of PRL-derived BAC$_{532}$ and PDR$_{532}$. In general, the POLIPHON method estimates an uncertainty of 40% (Ansmann et al., 2011). BravoAranda et al. (2015) compared the POLIPHON method directly with field aircraft measurements and found a relative difference of less than 30%. In our measurements, the uncertainties of BAC$_{532}$ and PDR$_{532}$ below 5 km are 2%–15% and 2%–20%, respectively. The uncertainty of the retrieved mass concentration is calculated by the law of error propagation. Each input parameter in Table S1 varies within a specified range of values (here we set it as 1 standard deviation), while the other input parameters are set to a fixed value. The sensitivity of the parameters in Table S1 is tested one by one. Combining the uncertainty of BAC$_{532}$, PDR$_{532}$, LR, mass density, and conversion factor of dust and anthropogenic aerosols, the uncertainty of mass concentration below 5 km is between 15% and 70%.

It must be cautioned that the POLIPHON method is based on the significant difference in particle shape between the two aerosol mixtures. At higher RH, the dust and anthropogenic aerosols’ mass concentration separated by POLIPHON method may not be reliable. Firstly, dust and anthropogenic aerosols’ mass densities vary with RH, depending on the actual composition of the aerosol mixture. Secondly, the increase in RH may lead to an increase in the
conversion factor. Most importantly, the shape of the originally non-spherical particles altered accordingly at high RH. All these changes will lead to an increase in the error of the POLIPHON method. Therefore, scenarios with higher RH (> 85 %) must be excluded when retrieving the dust and anthropogenic aerosols’ mass concentration (Ansmann et al., 2012, 2011). The RH profile from ERA5 reanalysis data was applied to exclude PRL-derived dust and anthropogenic aerosols’ mass concentration profiles under high-RH conditions from May 2019 to February 2022.

To ensure the reliability of the dust and anthropogenic aerosols’ mass concentration profiles, the PRL-derived particle mass concentration at 0.25 km was compared with the surface PM$_{10}$ and PM$_{2.5}$ mass concentration. Anthropogenic aerosols’ mass concentration at 0.25 km was positively correlated with surface PM$_{2.5}$ mass concentration, with a Pearson correlation $R = 0.63$ ($p < 0.01$; Fig. 2). The total aerosol mass concentration at 0.25 km was compared with the surface PM$_{10}$ mass concentration and had a Pearson correlation coefficient $R = 0.67$ ($p < 0.01$). Due to the influence of the water vapor effect (Wiegner and Gasteiger, 2015), it is difficult to get completely consistent results in the comparison. Moreover, PM$_{2.5}$ and PM$_{10}$ were measured at the ground level 15 km north of BMOC, while dust and anthropogenic aerosols’ mass concentration are data at an altitude of 0.25 km above the BMOC. Overall, it can be seen from the significant positive correlation in Fig. 2 that the POLIPHON method can reliably estimate the dust and anthropogenic aerosols’ mass concentration in Beijing.

3 Results

The data presented here were obtained in Beijing, China, by PRL, sun-photometer, and reanalysis data over the period from 22 May 2019 to 20 February 2022. Since PRL data were missing from 16 November 2020 to 29 May 2021 due to laser failure, the sun-photometer observations are used to demonstrate the representativeness of PRL measurements, by statistical comparison of sun-photometer and PRL AOD. Once the representativeness of the PRL data has been demonstrated, the annual cycle of the AOD and MLH can be analyzed and discussed, as well as the mean vertical profile of the dust and anthropogenic aerosols retrieved by the POLIPHON method.

3.1 Annual cycle of AOD

Figure 3a is intended to demonstrate the representativeness of PRL measurements on a statistical basis. This is achieved by comparing the annual cycle of PRL-derived AOD and sun-photometer-measured AOD; the error bar in PRL-derived AOD is calculated as 1 standard deviation. Figure 3a qualitatively and quantitatively shows an excellent agreement between the two datasets, with a Pearson correlation and root mean square error (RMSE) of 0.96 ($p < 0.01$) and 0.018, respectively. The Ångström exponent was employed to convert the sun-photometer-measured AOD at 500 nm to AOD at 532 nm. Although the number of PRL measurements was statistically low, the monthly AOD was similar for both datasets. This result reinforces the representativeness of PRL measurements because it shows that the range of the PRL-derived monthly AOD set is comparable to that of the sun photometer. The difference in the monthly
mean AOD between the sun photometer and the PRL ranges from 0.002 to 0.14, with the largest difference occurring in March, where the PRL-derived AOD is 0.14 (18%) smaller than the sun photometer’s AOD, which may be due to the absence of PRL measurements in March 2021. Two mega dust events occurred in March 2021 in Beijing, with the recorded peak AOD at 500 nm larger than 2.5 measured by the sun photometer; the satellite data registered these two massive dust events as the most intense episode at the same time in history over the past 20 years (Gui et al., 2022).

The PRL-derived annual mean AOD is 0.50 ± 0.29 at 532 nm. The annual variation of PRL-derived AOD showed a bimodal feature, with the first peak in March (0.61 ± 0.29), the second peak in July (0.81 ± 0.37), and a minimum AOD in December (0.32 ± 0.12); sun-photometer observations also reveal a similar annual cycle. The high-loading aerosols in July and March are primarily due to long-range transport air pollutants such as Asian dust (more frequent in March) and anthropogenic aerosols (more frequent in July) (Wang et al., 2021c). Figure 3b shows the PRL-derived AOD in the ML and FT. ML AOD is obtained by integrating the EXT profile from the ground to the top of the MLH, while FT AOD is obtained by integrating the EXT profile from the top of the MLH to 5 km. The error bar associated with the monthly mean AOD shows the 1 standard deviation. There are three interesting features in Fig. 3b as follows:

- FT AOD ranges from 0.14–0.35, implying the presence of aerosol layer above the ML all year round, and FT AOD increases significantly in March and July, which emphasizes the frequent long-range transport of dust and anthropogenic aerosols in spring and autumn.

- The annual cycle of the ML AOD and FT AOD is almost identical (Pearson correlation $R = 0.90$, $P < 0.01$), exhibiting a bimodal shape with the primary peak (0.44 ± 0.23 in ML and 0.35 ± 0.26 in FT) and secondary peak (0.34 ± 0.17 in ML and 0.28 ± 0.20 in FT) in July and March, respectively. This is a very intriguing result because ML AOD and FT AOD may be regulated by the same mechanisms, such as the regional transport of aerosols. Also note that ML AOD is consistently higher than FT AOD, with more aerosols concentrated within the ML.

- The annual cycle of AOD observed by the PRL and sun photometer showed the highest level in summer and the lowest in winter, while the surface observation showed diametrically contrary conclusions (Fig. 4). Due to the difference in atmospheric diffusion capacity, high surface PM$_{2.5}$ concentrations typically occur in winter and the lowest in summer. This implies a high load of aerosols in the upper air of Beijing, which will be discussed in detail in Sect. 3.3.2. An opposite annual cycle of air pollution at the surface and in the whole layer also provides motivation for further investigation of the potential relationship between surface PM$_{2.5}$ levels and columnar AOD.

### 3.2 Seasonal variability of the MLH, PDR$_{532}$, and LR$_{355}$

The MLH illustrates the relationship between the intensity, duration, and extent of air pollution. The maximum MLH in Beijing usually appears at 15:00 local time (LT), so this study focuses on analyzing the characteristics of the MLH at 15:00 LT (Wang et al., 2021a). To investigate the reliability of the PRL-derived MLH, we used the daily and monthly BLH reanalysis data at 15:00 LT from ERA5 to obtain a linear fit with the PRL-derived MLH. The daily and monthly Pearson correlation coefficient between two sequences is 0.78 and

Figure 4. Surface monthly mean PM$_{2.5}$ mass concentration collected from the Beijing Municipal Ecological and Environmental Monitoring Center and AOD at 500 nm measured by the sun photometer from May 2019 to February 2022.
The MLH in winter usually ranges from 0.81 to 1.31 km, and the high and fluctuating MLHs in March, April, and May are mainly attributed to the dynamics of strong winds, although the surface temperature was not high. Cloud and rain prevailed in July and August, especially in July, and less surface solar radiation and lower wind speed (Guo et al., 2016; Wang et al., 2021a) depress the elevation of the daytime MLH. The MLH reaches its second peak (1.57 ± 0.81 km) in September due to the strong solar radiation (Miao et al., 2015), while the height is significantly lower than the MLH in May. In November and December, the weak thermal convection and wind speed result in a relatively lower MLH.

The PDR$_{532}$ and LR$_{355}$ seasonal, ML, and FT density distribution from May 2019 to February 2022 are shown in Fig. 6. The broad distributions of seasonal, ML, and FT PDR$_{532}$ and LR$_{355}$ indicate the occurrence of very diverse aerosol conditions, including complex aerosol mixtures of anthropogenic and natural aerosol types and compositions. According to Cui et al. (2022), the major fine atmospheric particles sources in Beijing include residential coal, residential biomass, industrial combustion, and industrial process and vehicles’ emission, and in addition to all local sources, long-range transport of east Asian dust and anthropogenic pollution from the NCP also contributes to the observed complexity of aerosol mixtures (Wang et al., 2016). As seen in Fig. 6, the PDR$_{532}$ ranges from values in pure anthropogenic aerosol pollution conditions (0 % to 9 %) to typical values in heavy dust outbreak conditions (up to 39 % at 532 nm). Also, the values of LR$_{355}$ are between 19 and 96 sr and cluster around 46–61 sr. All mean values and 1 standard deviations of the PRL-derived PDR$_{532}$ and LR$_{355}$ are summarized in Table 1. The maximum and minimum values of PDR$_{532}$ occurred in spring (8.2 % ± 6.0 %) and winter (4.5 % ± 3.4 %), respectively, and the maximum and minimum LR$_{355}$ values occurred in winter (55.0 ± 11.4 sr) and autumn (50.2 ± 11.6 sr). The value of FT LR$_{355}$ in the summer is slightly larger than that in ML, while in winter, the value of ML LR$_{355}$ is considerably higher than that in FT. The values of ML LR$_{355}$ in winter and FT LR$_{355}$ in summer are about 55 sr, which is a typical value for anthropogenic aerosols. Moreover, the values of ML PDR$_{532}$ in winter and FT PDR$_{532}$ in the summer are 6.1 % ± 5.8 % and 3.8 % ± 2.2 %, respectively, which indicates that anthropogenic aerosols accumulate in the ML in winter and the FT in summer.

### 3.3 Annual cycle of aerosol vertical profiles

The POLIPHON method allows the conversion of aerosol EXT profiles into dust and anthropogenic aerosols’ mass concentration profiles. Vertical profiles of the monthly mean aerosol parameters are shown in Fig. 7, including EXT$_{532}$, PDR$_{532}$ POLIPHON-method-retrieved anthropogenic aerosol and dust mass concentration, LR$_{355}$, and the fraction of dust mass concentration. These monthly mean

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**Figure 5.** Seasonal density distribution of (a) the PRL-derived MLH and (b) the annual cycle of the PRL-derived MLH and BLH in ERA5 reanalysis data. Note that the MLH and BLH are the monthly mean values at 15:00 LT. The box-and-whisker plots show the 5th, 25th, 50th, 75th, and 95th percentiles, and the dots represent the monthly mean MLH.
profiles are the average of a certain number of mutually un-correlated individual profiles, some of which may contain aerosols only in the ML, while others may contain several aerosol layers in/above the ML. The number of points averaged at different heights is shown in Fig. S1b, excluding cases where the monthly average number of points is less than 1000.

### 3.3.1 EXT$_{532}$, PDR$_{532}$, and LR$_{355}$

The EXT$_{532}$ profile exhibits a clear seasonal variation (Fig. 7a). In autumn and winter, the lowest aerosol layer is shallower and has a higher value between 0.19 and 0.32 km$^{-1}$, and then the EXT$_{532}$ drops sharply with the increase in altitude. The variation of EXT$_{532}$ profiles in autumn and winter is mainly explained by meteorological conditions, the lower height of the inversion layer (usually less than 1.5 km), and the ML that is weakly developed (Fig. 5); particles are hard to transport to the upper level. The EXT$_{532}$ profile in summer is highest in the range of 0.36–0.96 km, with values between 0.27–0.42 km$^{-1}$, and then EXT$_{532}$ decreased slowly with increasing altitude. In spring, due to strong wind speed (Guo et al., 2016), the aerosol was lifted to a higher height and then gently decreased. The monthly mean EXT$_{532}$ profile shows that the highest mean EXT$_{532}$ is found in July (within the height range of 0.36–0.96 km), and the monthly mean EXT$_{532}$ in July gradually increases from 0.33 km$^{-1}$ at 0.25 km to a maximum value of 0.43 km$^{-1}$ at 0.56 km. EXT$_{532}$ was lowest in December, with a relatively superficial and thin aerosol layer gathering in the lowest layer (about 0.19 km$^{-1}$). The evolution of the monthly mean EXT$_{532}$ profile is also consistent with the previous observations (Zhang et al., 2023b). We further analyzed the top height of the aerosol layer; the top height of the aerosol layer is determined at points where the EXT$_{532}$ values were below 0.01 km$^{-1}$. A similar trend was found between the top height of the aerosol layer and the MLH. An extremely high aerosol layer is found in April, May, August, and September (more than 5 km) and the lowest in January (about 3.4 km), implying that the vertical diffusion capacity of aerosols in Beijing is mainly influenced by the MLH.

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**Figure 6.** Density distribution of PRL-derived aerosol optical parameters: (a) ML LR$_{355}$ and FT LR$_{355}$ and (b) ML PDR$_{532}$ and FT PDR$_{532}$. Seasonal density distribution of (c) LR$_{355}$ and (d) PDR$_{532}$.

**Table 1.** Annual and seasonal means and 1 standard deviation (calculated with ML estimation) of the PDR$_{532}$ and LR$_{355}$.

<table>
<thead>
<tr>
<th></th>
<th>PDR$_{355}$</th>
<th>LR$_{355}$</th>
<th>PDR$_{355}$</th>
<th>LR$_{355}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year</td>
<td>(%) (sr)</td>
<td>(%) (sr)</td>
<td>(%) (sr)</td>
<td>(%) (sr)</td>
</tr>
<tr>
<td>Spring</td>
<td>6.1 ± 5.0</td>
<td>51.6 ± 11.3</td>
<td>5.2 ± 3.6</td>
<td>6.5 ± 5.6</td>
</tr>
<tr>
<td>Summer</td>
<td>5.0 ± 3.4</td>
<td>50.1 ± 10.8</td>
<td>5.0 ± 3.6</td>
<td>6.1 ± 5.8</td>
</tr>
<tr>
<td>Autumn</td>
<td>5.9 ± 4.8</td>
<td>50.2 ± 11.6</td>
<td>5.1 ± 3.1</td>
<td>6.2 ± 5.4</td>
</tr>
<tr>
<td>Winter</td>
<td>4.5 ± 3.4</td>
<td>55.0 ± 11.4</td>
<td>3.8 ± 2.2</td>
<td>4.9 ± 3.8</td>
</tr>
</tbody>
</table>

https://doi.org/10.5194/acp-23-14271-2023
The PDR$_{532}$ profile exhibits considerably different seasonal variations than the EXT$_{532}$ profile (Fig. 7b). In January, February, July, August, and September, the monthly mean PDR$_{532}$ in the bottom layer (0.25 km) was extremely low (less than 5 %), indicating that the lower atmosphere of Beijing was almost unaffected by dust during this period. The monthly mean PDR$_{532}$ is highest in April and May, with values around 0.10 at 0.25 km, and the value of PDR$_{532}$ is between 8 %–12 % in the range of 1.6 to 4.1 km. The high PDR$_{532}$ values in spring are primarily impacted by the long-range transport of Asian dust. Under the influence of the spring cold front, dust from East Asia (particularly over the Taklimakan Desert and the Gobi Desert) is easily lifted into the FT and transported eastward, affecting the air quality in Beijing (Gui et al., 2022; Miao et al., 2015).

The LR$_{355}$ provides the possibility to estimate the properties and sources of the measured aerosols (Fig. 7e). Monthly mean LR$_{355}$ is highest in January; this is because Beijing is rarely affected by dust aerosol in January. Moreover, building heating leads to a high emission of absorbing air pollutants in January in Beijing (Zheng et al., 2019). The MERRA-2 global reanalysis data also show a higher concentration of black carbon in Beijing in winter (Fig. 8), which is consistent with the PRL observations. The monthly mean LR$_{355}$ profile of the whole year reveals that the monthly mean LR$_{355}$ is high below 2 km and has little change with a value of 52–64 sr. However, the monthly mean LR$_{355}$ profile above 2 km varies dramatically from 35 sr (more scattering) to 67 sr (moderate absorption), indicating the complexity of airborne particles transport in Beijing, the presence of more species, and relatively more aging aerosol layers.

### 3.3.2 Anthropogenic aerosols

The annual cycle of anthropogenic aerosols’ mass concentration retrieved by the POLIPHON method is analogous to that of EXT$_{532}$ (Fig. 7c), with the lowest anthropogenic aerosols’ mass concentration in December, less than 15 µg m$^{-3}$ in the range of 0.25 to 5 km. The monthly mean anthropogenic aerosols’ mass concentration in October, November, and January was comparable; there was a thin aerosol layer below 1.4 km (about 23 µg m$^{-3}$ at 0.25 km), which is due to the weak solar radiation and the low MLH in autumn and winter (Fig. 5b), inhibiting the upward uplift of particles. From February to May, the MLH gradually rises due to the gradual increase in surface wind speed and solar radiation, and the distribution of aerosol layer also rises significantly, while the monthly mean anthropogenic aerosols’ mass concentration at 0.25 km gradually decreases, from 40 µg m$^{-3}$ in February to 22 µg m$^{-3}$ in May. In June and July, the largest monthly mean anthropogenic aerosols’ mass concentration and the widest aerosol distribution altitude were found, especially in July, up to 57 µg m$^{-3}$ in the range of 0.4 to 0.9 km. This was observed by PRL measurements and could not be identified by ground observations.

Tang et al. (2015, 2016) found that regional transport contributes considerably to light pollution, while local contribution is dominant in heavy pollution, and low RH and high RH typically correspond to light pollution and heavy pollu-
Figure 8. Cluster analysis of seasonal 48 h air mass backward trajectories in Beijing from May 2019 to February 2022: initialized at (a) 100 m, (b) 2 km, and (c) 4 km in spring; initialized at (d) 100 m, (e) 2 km, and (f) 4 km in summer; initialized at (g) 100 m, (h) 2 km, and (i) 4 km in autumn; and initialized at (j) 100 m, (k) 2 km, and (l) 4 km in winter. We calculated the hourly air mass backward trajectories during each season. Then, cluster analysis was carried out in two to four category directions. The percentages at the bottom right of each panel indicate the percentage of each backward trajectory. The color in each panel indicates the AOD, dust AOD, and black carbon AOD for each season obtained from the MERRA-2 global reanalysis data.

Firstly, there are frequent southern transmission scenarios in the upper air over Beijing in summer. As can be seen from Fig. 8, the air mass backward trajectories at 2 and 4 km in summer are considerably different from the air mass backward trajectories in other seasons. The clustering analysis of the backward trajectories shows that in spring, autumn, and winter, the air masses at 2 and 4 km are primarily from the northwest; the northwest of Beijing is mainly mountainous with a relatively clean air mass; and in the further northwest, dozens of deserts are distributed. However, in summer, 40% and 21% of the air mass at 2 and 4 km is transmitted from the south, respectively. The densely populated, industrial NCP is the most polluted area in China (Zhang et al., 2019a). Thus, southerly transport brings anthropogenic aerosols with high RH from the NCP into the upper atmosphere of Beijing. Secondly, the hygroscopic growth of atmospheric particles further strengthens the air pollution in the upper air over Beijing. The atmospheric particle hygroscopicity in Beijing is significantly influenced by the source of air masses (Tang et al., 2016). When southerly air masses dominate, the polluted southerly air masses (heavy polluted and high RH) will enhance the hygroscopicity of atmospheric particles in Beijing, and as the mass fraction of inorganic salts (especially nitrate) increases, the particles undergo sufficient aging and mixing.
processes, and the particle hygroscopicity can be further increased (Cheng et al., 2016; Zhang et al., 2023a).

We further analyzed the relationship between the relative humidity and EXT$_{532}$ and PDR$_{532}$ at the top of the mixing layer in summer (Fig. 9). When the RH was greater than 40%, the EXT$_{532}$ increased with the increase of RH; that is, the air pollution increased with the increase of humidity. More importantly, with the increase of RH, the PDR$_{532}$ gradually decreases, indicating that there is hygroscopic growth of particles at the top of the mixing layer. Because the PDR$_{532}$ is related to hygroscopicity (Dawson et al., 2020), it is inversely proportional to the sphericity of atmospheric particles. Assuming a uniform refractive index, moistened aerosol particles are thought to be more spherical due to condensation of water vapor and surface tension, resulting in a lower PDR$_{532}$. Thus, there is hygroscopic growth of particles at the top of the mixing layer, but the contribution of hygroscopic growth to anthropogenic aerosols at the top of the mixing layer is still uncertain.

The monthly mean wind speed and RH obtained from the ERA5 reanalysis data further confirm the above scenario, with higher RH and weak southerly wind in Beijing during summer (Fig. 9). Low wind speed is a prerequisite for haze episodes, and high RH is favorable for air pollution formation. In addition, Beijing is cloudy in summer (cloud fraction $>50\%$) due to the intrusion of the western Pacific subtropical high pressure (Tang et al., 2016). When the cloud fraction increases, solar radiation and the MLH will drop sharply (Fig. 5b). Such unfavorable meteorological conditions (southerly transport, high RH, low MLH, and wind speed) lead to a rapid formation of atmospheric particles in the upper air over Beijing in summer. Therefore, the high anthropogenic aerosols’ mass concentration in the upper air (0.4–0.9 km) over Beijing in summer is mainly caused by the southward transport in the upper air, where the atmosphere is relatively stable and moist, favoring hygroscopic growth of particles.

In fact, similar upper-air pollution transport was also found in PRL observations during the rest of the seasons in Beijing. Tao et al. (2014) and our previous studies (Wang et al., 2020) have analyzed the persistent haze episodes of Beijing’s upper air in spring, autumn, and winter in detail. Different from summer, the air pollution in the upper air of Beijing is dominated by dust mixed with anthropogenic aerosols in spring, autumn, and winter, accompanied by the hygroscopic growth of atmospheric particles. In contrast, the upper atmosphere of Beijing is most polluted in summer because the southern transport is more frequent (Fig. 8), and the upper air of Beijing is more stable and humid (Fig. 10), which is more favorable for the hygroscopic growth of atmospheric particulates.

### 3.3.3 Asian dust

Meteorological conditions in Beijing vary significantly between the seasons. Air mass transport patterns are under the influence of strong westerlies over Beijing in spring (Kok et al., 2021; Gui et al., 2022). As early as February, there was substantial dust transport within 1.6–5 km, although it was not visible at the ground (Fig. 7d). From March to June, Beijing was frequently subjected to dust pollution, and the distribution of dust mass concentration at different heights was quite different. The dust mass concentration is 27, 59, 61, and 41 µg m$^{-3}$ at 0.25 km from March to June, respectively. With the increase in altitude, the dust mass concentration gradually decreased, and then it increased significantly after reaching a certain altitude. Dust aerosol is mainly distributed in 2.1–2.7 km (39–46 µg m$^{-3}$) in March and more widely distributed in April (1.1–3.0 km), with higher dust mass concentration at different heights was quite different. The dust mass concentration is 27, 59, 61, and 41 µg m$^{-3}$ at 0.25 km from March to June, respectively. With the increase in altitude, the dust mass concentration gradually decreased, and then it increased significantly after reaching a certain altitude. Dust aerosol is mainly distributed in 2.1–2.7 km (39–46 µg m$^{-3}$) in March and more widely distributed in April (1.1–3.0 km), with higher dust mass concentration between 40 and 60 µg m$^{-3}$. Compared with April, the distribution height of high dust mass concentration in FT increased further in May and June, ranging from 2.3–3.5 and 2.5–3.9 km, respectively. The difference in dust vertical distribution is mainly affected by meteorological factors and
Table 2. Means and 1 standard deviation of the PDR\textsubscript{352} and LR\textsubscript{355} for the three aerosol types.

<table>
<thead>
<tr>
<th></th>
<th>Anth. aero.</th>
<th>Polluted dust</th>
<th>Asian dust</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDR\textsubscript{352} (%)</td>
<td>4.3 ± 2.1</td>
<td>13.2 ± 3.5</td>
<td>28.7 ± 4.3</td>
</tr>
<tr>
<td>LR\textsubscript{355} (sr)</td>
<td>52.1 ± 11.3</td>
<td>47.8 ± 11.0</td>
<td>45.7 ± 5.1</td>
</tr>
</tbody>
</table>

3.3.4 Occurrence frequency of aerosol types

Three aerosol types in Beijing have been identified according to PDR\textsubscript{352}, Asian dust, anthropogenic aerosols, and their mixtures (polluted dust). The means and 1 standard deviations of the PDR\textsubscript{352} and LR\textsubscript{355} for the three aerosol types are listed in Table 2, and the vertical occurrence frequency of the three aerosol types is shown in Fig. 11. In general, anthropogenic aerosols occurred more frequently than dust and polluted dust below 5 km. In January, February, July, August, and September, the occurrence frequency of anthropogenic aerosols below 1.1 km is close to 100 %, and the occurrence frequency of anthropogenic aerosols above 1.1 km decreases more steeply in January and February than in July, August, and September, which is consistent with the vertical distribution characteristics of PDR\textsubscript{352}. From March to June and October to December, the maximum occurrence frequency of anthropogenic aerosols occurred at heights of 0.39 km (95 %), 0.77 km (86 %), 0.9 km (76 %), 0.75 km (91 %), 0.65 km (91 %), 0.62 km (91 %), and 0.69 km (99 %), respectively, none of which was at the bottom (0.25 km), indicating that a large number of elevated aerosols gather over Beijing. It is also mentioned in Sect. 3.3.2 that the southerly winds bring the high polluted and humid air masses from the NCP over Beijing, and under stable meteorological conditions, further hygroscopic growth of atmospheric particles leads to severe air pollution in the upper atmosphere of Beijing (Tao et al., 2014; Tang et al., 2015, 2016). It is worth noting that these upper-layer aerosols may trigger aerosol-radiation feedback effects, which will strengthen the surface air pollution in Beijing (Zhong et al., 2018).

The maximum occurrence frequency of dust typically occurs at 2–4 km in March to June, with a frequency of 0 %–9 % (Fig. 11), and a small amount of dust aerosols also appears in other months, usually above 1 km, with a frequency of less than 3 %. The vertical occurrence frequency of polluted dust also shows considerably seasonal variation. From March to June, polluted dust occurred more frequently (5 %–39 %) below 4 km, and the height of maximum occurrence frequency was about 2.5 km. In addition, the occurrence frequency of polluted dust at 0.25 km is between 10 % and 39 % from March to June, indicating the considerable contribution of dust aerosol to the bottom air pollution in Beijing. From July to September, a small amount of polluted dust occurs above 1.2 km, with a frequency of 0 %–19 %. In October and November, polluted dust is mainly distributed in 0–3 km and contributed more frequently to the bottom layer (0.25 km) and 1.2–2.4 km, up to 30 %. In December, January, and February, the polluted dust usually appeared in 1.1–2.4 km, while the anthropogenic aerosols were concentrated below 1 km, and aerosols tended to be vertically stratified in winter. The stratification of aerosols is mainly caused by the difference of meteorological conditions (dominated by northwest winds) and below (dominated by southerly winds).

dust sources. Firstly, with the increase in temperature and enhancement of convection from April to June, dust particles can be lifted higher into the air. Secondly, as two major dust sources in East Asia, the Taklimakan and Gobi Desert in northwest China have some differences in their eastward transmission paths (Sun et al., 2001). The Gobi Desert dust is transported eastward in the bottom troposphere and often affects the surface air quality in Beijing, while the Taklimakan Desert dust is readily lifted to the upper troposphere and transported eastward.

In July, August, and September, Beijing is rarely affected by dust aerosols (Fig. 7d), with monthly mean dust mass concentration below 5 µg m\textsuperscript{−3} in the range of 0.25 to 1.5 km and below 10 µg m\textsuperscript{−3} in the range of 1.5 to 5 km. In October and November, there was also a significant dust transport episode in Beijing, but its concentration and height distribution range were considerably smaller than those from March to June. The dust mass concentration in October and November at 0.25 km is 30 and 34 µg m\textsuperscript{−3}, respectively, and the dust in the lower FT was concentrated in the 1–2 km, with concentrations of around 25 µg m\textsuperscript{−3}. In December and January, the dust mass concentration was less than 5 µg m\textsuperscript{−3} below 1 km, while dust mass concentration increased slightly to about 10 µg m\textsuperscript{−3} within 1–2 km. The proportion of dust mass concentration in the total aerosol mass concentration at 0.25 km showed that (Fig. 7f) the highest share is in April (52 %), followed by May (51 %), October (38 %), June (37 %), November (34 %), and March (29 %). In the FT, the proportion of dust mass concentration of the total aerosol mass concentration is 10 %–60 % from March to June and 12 %–43 % from October to November.

In general, the PRL observations show that the dust is widely distributed in height, which can be consistently extended from the surface to more than 5 km; the dust concentration is discontinuous in the vertical direction; and there is stratification, not only on the ground but also in lofted layers that reach up to several kilometers. The heights of these lofted dust layers showed apparent seasonal dependence, with the height of the main dust layer gradually ascending from 1.1 km to about 2.5 km from April to June and below 3 km in autumn.

dust aerosols (Fig. 7d), with monthly mean dust mass concentration of around 25 µg m\textsuperscript{−3}. The Gobi Desert dust is transported eastward in the bottom troposphere and often affects the surface air quality in Beijing, while the Taklimakan Desert dust is readily lifted to the upper troposphere and transported eastward.

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the ML. This has been discussed in detail in our previous studies (Wang et al., 2020).

4 Discussion

The vertical structure of the ML is critical to surface air pollution because it affects the volume of air pollutants’ mixing (Miao et al., 2018). If the factor driving the variation of surface air pollution with the MLH is meteorological, then the discussion of their relationship can elucidate meteorological effects. We examine the MLH and the bottom (0.25 km) dust and anthropogenic aerosols’ mass concentration retrieved by PRL on a case-by-case basis. Scatter plots of the MLH with total, anthropogenic aerosols, and dust mass concentration are shown in Fig. 12a–c. Although total aerosol mass concentration is negatively correlated with the MLH, there are substantial differences and spreads in the correlations between dust and anthropogenic aerosols’ mass concentration and the MLH. Linear regression and inverse function fitting methods were applied to characterize the relationship between the MLH and dust/anthropogenic aerosols’ mass concentration. Anthropogenic aerosols’ mass concentration and the MLH were significantly negatively correlated, with a Pearson correlation of $-0.36$. In addition, the nonlinear inverse function was in high agreement with the mean of each bin and characterized the relationship between the MLH and anthropogenic aerosols’ mass concentration, with a higher correlation coefficient ($-0.49$), and this improvement was significant for total and anthropogenic aerosols’ mass concentration.

Anthropogenic aerosols’ mass concentration and the MLH are significantly negatively correlated, while dust mass concentration and the MLH are weakly correlated and not statistically significant (Fig. 12c). In fact, surface air pollution is affected by the combined effects of local emissions and meteorological conditions with different spatial and temporal distributions, and the MLH is only one of these influencing factors. Previous studies (Zhong et al., 2018) also pointed out that anthropogenic air pollution episodes in Beijing are typically divided into two processes, transmission and accumulation. In the transmission stage, air pollution is primarily induced by southerly transport, while in the accumulation stage, meteorological conditions are stagnant, and anthropogenic aerosols usually exhibit explosive growth. Subsequently, the anthropogenic aerosols within the ML scatter more solar radiation back to space, thus increasing the temperature inversion and lowering the MLH, further deteriorating the surface air quality. There is a positive feedback mechanism between the MLH and surface anthropogenic aerosols in this process. Therefore, the MLH shows a significant negative correlation with bottom anthropogenic aerosols’ mass concentration. However, dust aerosol in Beijing is governed by regional transport, and the MLH is not the key parameter controlling surface dust mass concentration. Our previous study (Wang et al., 2021c) shows that, for pure dust episodes, air masses are transported from the northwest, with stronger wind speed and a high MLH. For polluted dust episodes, air masses are transported from northwest to the NCP and then transmitted to Beijing by southerly winds, and the generation rate of new particles is further accelerated by the mixing of dust and anthropogenic aerosols (Nie et al., 2014), which is prone to persistent heavy haze events with a low MLH at this time. In addition, local dust sources (soil dust, construction dust, coal dust, and motor vehicle exhaust) in Beijing are usually concentrated on the ground (Wang et al., 2015), which is difficult to transmit to several hundred meters. Therefore, the bottom dust mass concentration is mainly...
Figure 12. The relationship between the PRL-derived MLH and total, anthropogenic aerosols, and dust mass concentration at 0.25 km in (a–c) the whole observation period, (d–f) spring, (g–i) summer, (j–l) autumn, and (m–o) winter in Beijing from May 2019 to February 2022. The black dots and vertical bars represent the mean values and 1 standard deviation for each bin. The interval of each bin is 0.2 km. The black and red lines represent the linear fit ($f(x) = Ax + B$) and inverse fit ($f(x) = A/x + B$), respectively. The fitting functions and correlation coefficients for linear fit (black) and inverse fit (red) are shown in the top right of each panel. The asterisks next to the correlation coefficient $R$ indicate statistical significance ($P < 0.01$). Color-shaded points represent normalized sample density.

Influenced by transport, and the correlation between the bottom dust mass concentration and the MLH is insignificant in long-term statistics.

The analysis in Sect. 3.1 and 3.2 shows that since both the MLH and air pollution in Beijing undergo the most pronounced seasonality, seasonal differences between the MLH and aerosol are analyzed further. Figure 12d–o focus on the seasonal dependence of the MLH and bottom dust and anthropogenic aerosols’ mass concentration. There is a significant negative correlation between anthropogenic aerosols
and the MLH in four seasons, while dust particles and the MLH are weakly correlated. The maximum magnitude of slope between $-1/\text{MLH}$ and anthropogenic aerosols’ mass concentration is about 45 km $\mu$g m$^{-3}$ in summer, followed by 40 km $\mu$g m$^{-3}$ in winter. The spring and autumn are comparable, with values of 25 and 28 km $\mu$g m$^{-3}$, respectively. This indicates that the sensitivity of anthropogenic aerosols to the MLH is strongest in summer, followed by winter, and is weakest in spring and autumn. The anthropogenic aerosols do not increase linearly with the decrease in the MLH. When the MLH was less than 1.2 km, anthropogenic aerosols increased rapidly with the decrease in the MLH, while anthropogenic aerosols changed more slowly when the MLH was greater than 1.2 km, and the relationship between the MLH and anthropogenic aerosols is virtually flat when the MLH is greater than 3 km.

In addition, Fig. 13a–c show the total, anthropogenic aerosols, and dust mass concentration profiles as a function of the MLH, as derived by PRL in Beijing from May 2019 to February 2022, with the POLIPHON method applied. Figure 13d–o show the seasonal relationship between the PRL-derived MLH and vertical profiles of total, anthropogenic aerosols, and dust mass concentration. The interval of the MLH is 0.2 km. We require that the number of averaged particle mass concentration profiles and MLH samples is larger than 20 for each interval of the MLH; otherwise, the data will be discarded, as shown in the white area of Fig. 13.

The anthropogenic aerosols’ mass concentration within the ML decreases continuously with an increasing ML (Fig. 13b). When the MLH is less than 0.4 km, high total aerosol mass concentration ($>50 \mu$g m$^{-3}$) extends up to 1.6 km, in which the anthropogenic aerosols (about 60 $\mu$g m$^{-3}$) are primarily in the ML, and the dust aerosol (about 50 $\mu$g m$^{-3}$) is mainly above the ML, and this aerosol stratification at a low MLH usually occurs in winter (Fig. 13m–o). When the MLH is between 0.4 and 2.0 km, the anthropogenic aerosols’ mass concentration increased from the bottom to the top of the ML and reaches a maximum at the top of the ML (35–60 $\mu$g m$^{-3}$). The accumulation of anthropogenic aerosols at the top of the ML is most significant in summer, with concentrations up to 57 $\mu$g m$^{-3}$, and a similar phenomenon also exists in spring, autumn, and winter, but the aerosol mass concentration is lower. We have discussed this phenomenon in detail in Sect. 3.3.2 because the southward transport is more frequent in summer (Fig. 8), and the air pollutants aloft in Beijing are more stable and moist (Fig. 10), which is favorable for the hygroscopic growth of atmospheric particles at the top of the ML. When the MLH is greater than 2 km, the anthropogenic aerosols’ mass concentration from the bottom to high altitude is less than 15 $\mu$g m$^{-3}$, and the atmosphere is relatively clean.

The relationship between the MLH and anthropogenic aerosols’ vertical profiles is quite different from the relationship between the MLH and dust vertical profiles. No matter how much the MLH is, there is always a shallow dust layer in the bottom layer, with mass concentration between 8 and 40 $\mu$g m$^{-3}$ (Fig. 13c), which further confirms that there is no significant correlation between the MLH and the bottom dust mass concentration. Moreover, high dust mass concentration is also distributed near the ML (1.4–3.4 km), with maximum mass concentration up to 104 $\mu$g m$^{-3}$. Seasonal variations show that (Fig. 13d–o), when the MLH ranges from 0.8 to 1.6 km in spring, high dust mass concentration extends up to 3.2 km, while dust aerosols are mainly concentrated near the ML (1.6–3.4 km) when the MLH is greater than 1.6 km. Different from spring, dust mass concentration is relatively low when the MLH is less than 1.6 km in summer, while high dust mass concentration is found around ML (1.6–3.4 km). The dust mass concentration in the ML is slightly lower than that above the ML in autumn and winter when the MLH is below 1.6 km.

In general, there is a significant negative correlation between bottom anthropogenic aerosols and the MLH. When the MLH is between 0.4 and 2 km, anthropogenic aerosols typically accumulate at the top of the ML, and the bottom anthropogenic aerosols have lower concentrations, especially in summer. Aerosols are prone to stratification when the MLH is less than 0.4 km. The relationship between dust and the MLH is complicated. There is no clear relationship between bottom dust mass concentration and the MLH, while the dust in the upper air tends to be distributed near the ML.

5 Conclusions

PRL was deployed to investigate the aerosol vertical distributions and their relationship with the MLH over Beijing from 22 May 2019 and 20 February 2022. The broad distributions of seasonal, ML, and FT PDR$_{532}$ and LR$_{355}$ were found over Beijing, indicating the occurrence of very diverse aerosol conditions. The PDR$_{532}$ ranges from values in pure anthropogenic aerosol pollution conditions (0 % to 9 %) to typical values in heavy dust outbreak episodes (up to 39 %). Also, the LR$_{355}$ is between 19–96 sr and clusters around 46–61 sr. The aerosol classification results also show that, besides anthropogenic aerosols, the maximum occurrence frequency of dust usually occurs at 2–4 km (0 %–9 %) in March to June, and the maximum occurrence frequency of polluted dust usually appears in 2.5 km (13 %–39 %) from February to June and 1.5 km (5 %–30 %) from November to January. These valuable datasets acquired through PRL can be used to improve lidar algorithms for current and/or future satellite lidar missions.

The annual variation of PRL-derived AOD showed a bimodal characteristic, with the first peak in March (0.61 ± 0.29) and the second peak in July (0.81 ± 0.37). In addition, the annual cycle of the ML AOD and FT AOD is virtually identical (Pearson correlation $R = 0.90$, $P < 0.01$); it may be regulated by the same mechanisms, such as regional transport of aerosols. Most importantly, the annual
The relationship between the PRL-derived MLH and vertical profiles of total, anthropogenic aerosols, and dust mass concentration in (a–c) the whole observation period, (d–f) spring, (g–i) summer, (j–l) autumn, and (m–o) winter for each bin in Beijing from May 2019 to February 2022. The interval of each bin is 0.2 km. The dashed black line represents the MLH.

cycle of AOD observed by PRL and the sun photometer was the highest in summer and the lowest in winter, while the surface observation showed diametrically contrary conclusions. This implies a high load of aerosols in the upper air of Beijing. Further retrieval of dust and anthropogenic aerosols’ mass concentration by the POLIPHON method revealed that, in June and July, the largest monthly mean anthropogenic aerosols’ mass concentration and the widest aerosol distribution altitude were found, especially in July, up to 57 µg m\(^{-3}\) in the range of 0.4 to 0.9 km. The high anthropogenic aerosols’ mass concentration in the upper air (0.4–0.9 km) over Beijing in summer is mainly caused by the southward transport in the upper air. The PRL observations show that the dust can be consistently extended from the surface to more than 5 km, and the dust concentration is discontinuous in the vertical direction, not only on the ground but also in lofted layers that reach up to several kilometers. The heights of these lofted dust layers exhibited apparent seasonal dependence, with the height of the main dust layer...
gradually rising from 1.1 km to about 2.5 km from April to June and below 3 km in autumn and winter.

There is a significant negative correlation between the bottom anthropogenic aerosols’ mass concentration and the MLH ($R = -0.36, P < 0.01$) due to the positive feedback mechanism between the MLH and surface anthropogenic aerosols, and an inverse function fit is more suitable to characterize this relationship ($R = -0.49, P < 0.01$). However, there is a weak correlation between the bottom dust mass concentration and the MLH in long-term statistics, which is not statistically significant because the bottom (0.25 km) dust mass concentration is mainly influenced by transport. The relationship between the MLH and anthropogenic aerosols’ mass concentration profiles shows that anthropogenic aerosols’ mass concentration increases from the bottom to the top of the ML and reaches a maximum at the top of the ML (35–60 µg m$^{-3}$). The accumulation of anthropogenic aerosols at the top of the ML is most pronounced in summer, with monthly mean mass concentration up to 68 µg m$^{-3}$, and similar phenomena also exist in spring, autumn, and winter but with lower mass concentration. This is because the upper-air southward transport is more frequent in Beijing in summer, and the atmosphere is more stable and moist, which is more conducive to the hygroscopic growth of atmospheric particles.

Although our results elucidate the long-term vertical distributions of dust (coarse) and anthropogenic aerosols (fine) and their relationships with the mixing layer height, our research also has two shortcomings. Firstly, PRL has an incomplete overlap region, about 0.25 km, which prevents us from capturing the evolution of air pollutants at the lowest level (0–0.25 km). Due to the incomplete overlap region, our inversion of the MLH also starts from 0.25 km, which may lead to overestimation of the MLH. Secondly, due to the limitations of the POLIPHON method, we excluded cases with RH greater than 85%, and the accumulation of particles at the top of the ML may undergo a significant hygroscopic growth, so the anthropogenic aerosols’ mass concentration at the top of the ML may be underestimated, and the hygroscopic growth of aerosols at the top of the ML is also worthy of further detailed exploration. The research on dust and anthropogenic aerosols’ vertical distributions and their correlation with the MLH contributes to further understanding of the contribution of dust–anthropogenic aerosols to the environment and climate and helps to supplement the dust–anthropogenic aerosol database with useful empirical information for upgrading atmospheric chemistry models and air pollution forecasting and warning.

**Code and data availability.** Measurement data in this study are available in the data repository maintained by Mendeley Data at https://doi.org/10.17632/tk7czvw7ty.1 (Wang et al., 2023). Additional code and data related to this paper may be requested from the authors.

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**Author contributions.** CL and CX conceived and supervised the study; ZW analyzed the Raman lidar data; ZW wrote the manuscript with input from CL, CX, and CS; and HZ reviewed and commented on the paper. All authors contributed to discussing the results and revising the manuscript.

**Competing interests.** The contact author has declared that none of the authors has any competing interests.

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