

Investigation of Aerosol Transport Flux Structure over Beijing Based on Lidar Observations and the Impact of Dust Transport on Air Quality

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Abstract. The origins, spatial distribution, and diffusion mechanisms of aerosols hold practical guiding significance for regional haze governance. The vertical and horizontal fluxes of aerosols serve as effective parameters for assessing the diffusion efficiency of aerosols, but they are less exploited due to insufficient observations. This study uses polarization lidar to differentiate between the aerosol sources of dust and non-dust and to estimate the mass concentration profiles of each. Combining the wind profiles acquired from Doppler wind lidar, the vertical and horizontal fluxes profiles of two type aerosols are calculated. This approach is designed to account for the influence of local aerosol transport mechanisms on air pollution, enabling a more precise reflection of the internal variations within a particular region. A winter haze event in Beijing from November 2nd to November 4th, 2023 was analyzed to distinguish the vertical distribution of and mass concentration and fluxes of aerosols brought by dust transported from the northwest monsoon and those from anthropogenic emissions within the North China plain area. Further analysis of different wind zones revealed that the aerosol concentration and fluxes from urban areas (regions with a higher density of anthropogenic sources) can be used to assess the local pollutant diffusion capacity, as well as the influence of vertical turbulence on ground PM₁₀ concentrations. Taking Beijing as an example, this study investigated the diffusion characteristics of urban aerosols, ultimately providing technical means and data references for early warning of urban air pollution and assessment of air pollution control measures.

1 Introduction

30 Atmospheric particulate matter or aerosols are fine solid or liquid particles that come from direct emissions or secondary conversion of gas - particulate matter and can suspend in the air. Atmospheric particulate matter pollution is one of the issues that the whole world pays attention to, and there is sufficient evidence showing that it has adverse impacts on human daily life

and health (Brunekreef et al., 2002). Anthropogenic aerosol emissions are one of the primary sources of air pollution. Klimont et al. (2017) estimated that the global anthropogenic total emissions of PM_{2.5} in 2010 were approximately 80 million tons, among which anthropogenic combustion aerosols (including transportation and industrial emissions) accounted for more than 80% (Klimont et al., 2017). China, with its vast territory, high population density and rapid economic development, has inevitably produced a large amount of anthropogenic aerosols during its development process, which has had a significant impact on air quality in East Asia. Since 2013, China has dedicated substantial efforts to air pollution governance and has promulgated numerous measures for managing and controlling aerosol emissions in cities (The State Council of the People's Republic of China, 2013; Geng et al., 2019). Nevertheless, these measures are primarily centered on the overall supervision of emission volumes. There exists an absence of a holistic approach to address various aspects of aerosol emissions, including the temporal dynamics, spatial patterns, dispersion processes, and the principal determinants that significantly contribute to air pollution. As a result, it remains challenging to establish an accurate and timely early warning system. Consequently, research on the emission and diffusion mechanisms of urban anthropogenic aerosols holds significant practical guiding value for the effective formulation and evaluation of emission policies.

Aerosols can be lifted from the Earth's surface into the atmosphere due to factors such as deep convection, frontal passages, and turbulence within the atmospheric boundary layer (Chen et al., 2012). This vertical transport mechanism is crucial for the initial step of regional-to-intercontinental transport. It is also of great significance for gaining an in-depth understanding of the frequent occurrence of long-duration and large-scale air pollution in China, as well as for improving urban air quality across the country. For example, it has revealed a new mechanism by which severe haze pollution in the Beijing-Tianjin-Hebei and Yangtze River Delta urban agglomerations during autumn and winter is intensified through mutual interaction and long-distance transport (Huang et al., 2020). Meanwhile, this can provide new insights for mitigating regional-scale mutual influences, thereby achieving scientific and precise haze control.

Atmospheric pollution in urban areas is mainly caused by anthropogenic emissions, including vehicle exhaust emissions, industrial emissions, and the combustion of carbon substances in daily life. Therefore, anthropogenic aerosols in urban areas can serve as an "emission indicator" for air pollution sources, which can be used to evaluate the effectiveness of air pollution reduction and air quality management measures. In the causes of air pollution, in addition to pollutant emission sources, the atmospheric planetary boundary layer height (PBLH) also plays an important role in controlling the temporal and spatial variations of pollutant concentrations (Ding et al., 2015). In most cases, anthropogenic air pollutants emitted from the surface are confined within the atmospheric planetary boundary layer, and their vertical mixing height directly determines the vertical mixing space of pollutants (Lou et al., 2019; Gerbig et al., 2008).

The interaction between vertical atmospheric pollution and the atmospheric boundary layer feedback process is an important factor in enhancing the transport of pollution between regions. The vertical flux of atmospheric aerosols is a key physical quantity used to quantitatively characterize the net transport of aerosols in the vertical direction per unit area and per unit time. It connects surface processes, boundary layer turbulence, aerosol cycling, and meteorological conditions. Its core significance lies in quantifying the intensity and direction of aerosol transport in the vertical direction, thereby revealing the

source and sink mechanisms of aerosols, their diffusion capabilities, and regional environmental impacts (Huang et al., 2024). Therefore, accurately assessing aerosol fluxes depends on the characteristics of the aerosol sources, the regional features, and the meteorological conditions within the atmospheric boundary layer. As Huang et al. (2020) demonstrated, aerosols themselves, especially light-absorbing components such as black carbon, can significantly influence the boundary layer structure through Aerosol-Boundary Layer Interaction (ABI), and amplify the accumulation and transport processes of regional pollutants (Huang et al., 2020). Due to the complexity and nonlinearity of the causes of atmospheric compound pollution under China's unique environmental conditions (Lv et al., 2017), more stringent requirements are placed on detection tools and methods in aerosol flux research. Typically, a comprehensive assessment of aerosol flux necessitates the employment of a combination of various methodologies and technologies, encompassing ground-based observations and vertical stereo-
70 detection techniques. Moreover, it is difficult to estimate the emission of particulate matter and the capacity of dry deposition at the surface to remove particulate matter from the air, which makes it impossible to parameterize these processes for application in atmospheric numerical simulation techniques.

Horizontal transport caused by wind speed and wind direction is also an important factor restricting the mixing space of pollutants (Weissmann et al., 2005; Wang et al., 2009; Tian et al., 2020; Zhang et al., 2020; Hou et al., 2022; Lin et al., 2023).
80 However, in the actual assessment and prediction of air pollution, it is difficult to consider the three factors of emission volume, atmospheric boundary layer, and dominant wind speed in a coordinated manner. Consequently, the horizontal flux can offer crucial insights into the horizontal transport of aerosol sources within cities and the net emissions resulting from external sources. For example, the lidar observations in Singapore (Huang et al., 2024) have shown that cross-border haze originating from biomass burning in Indonesia is transported via the nocturnal low-level jet. During the development of the daytime
85 boundary layer, it is entrained to the ground surface, leading to a 500% increase in the local PM_{2.5} concentration (Huang et al., 2024). Conversely, the findings of Zhang et al. (2015) suggest that the prevailing wind exerts a substantial influence on the pollution concentration in cities. For instance, the northerly wind is correlated with relatively lower pollutant concentrations in Beijing (Zhang et al., 2015).

Over the past two decades, many scholars have attempted to use different observation methods to conduct targeted studies
90 on the sources, diffusion mechanisms, spatial distribution, and other important influencing factors of particulate matter pollution (Tanaka et al., 2005; Weissmann et al., 2005; van Donkelaar et al., 2016; Huang et al., 2024). In the field campaigns of aerosol observation techniques, in-situ continuous monitoring remains one of the principal detection approaches. Its advantage is the ability to provide a high temporal resolution and straightforward aerosol mass concentration data. In-situ sampling monitoring can not only offer a relatively comprehensive understanding of aerosol characteristics but also be utilized
95 to analyze the composition of particulate matter. Additionally, a network of such sampling points can be established for research on regional near-surface particulate matter emissions (Liu & Daum, 2000; Drinovec et al. 2015). Currently, numerous methods for measuring the surface-atmosphere exchange processes of aerosols have been developed (Pryor et al., 2008). Among these, the most representative, direct, and productive systematic observational instruments and data analysis techniques for aerosol fluxes are associated with the European ACTRIS (Aerosol, Cloud and Trace Gases Research Infrastructure)

100 program. It is important to note that while eddy-covariance measurements for aerosols are not a standard component of the
ACTRIS core program, they are occasionally conducted as research activities at some ACTRIS sites. For example, such
measurements can be performed by combining ultrasonic anemometers and particle size spectrometers on meteorological
towers to obtain single-point aerosol fluxes. However, this approach can only represent relatively small areas. However, this
105 approach can only represent relatively small areas. As a result, when applying the aerosol flux results to mesoscale circulation
models, significant uncertainties may occur (Chen et al., 2008). Consequently, there is an urgent demand for vertical profile
observations of aerosol fluxes that span the entire atmospheric boundary layer.

In recent years, remote sensing technology, particularly lidar, has become a powerful tool for aerosol research due to its
high spatiotemporal resolution and detection sensitivity (Li et al., 2019; Lopatin et al., 2013; Che et al., 2019; Wang et al.,
2019; Weinzierl et al., 2011; Denjean et al., 2016; Córdoba-Jabonero et al., 2018). Unlike passive sensors, lidar systems
110 actively probe the atmosphere and can retrieve vertically resolved aerosol properties—such as depolarization ratio, extinction
coefficient, and mass concentration—along with wind field information within the planetary boundary layer (PBL) (Müller et
al., 2001; Böckmann et al., 2001; Mamouri & Ansmann, 2014). The combination of multi-wavelength polarization Raman
lidar and Doppler lidar offers a particularly advanced approach for quantifying aerosol fluxes. The pioneering work
successfully demonstrated the application of the eddy covariance technique by combining Doppler wind lidar and Raman
115 aerosol lidar to retrieve vertical aerosol flux profiles, validating the core methodology of synergistic lidar observations
(Wandinger et al., 2004). By leveraging covariance algorithms and co-located measurements of aerosol properties and 3D
wind vectors, this synergistic technique enables direct estimation of both vertical and horizontal aerosol transport with high
accuracy (Engelmann et al., 2008; Huang et al., 2024). Thus, the integration of these lidar technologies provides a feasible and
robust method for studying aerosol fluxes within the atmospheric boundary layer. However, most of the current detection
120 methods and contents have the following deficiencies: Insufficient detection methods and research on anthropogenic aerosol
emissions. Most of the current flux research methods use in-situ or passive remote sensing observations (e.g. Yuan et al., 2019;
Conte & Held, 2021; Casquero-Vera et al., 2022; Hu et al., 2022), lacking high-resolution vertical spatial scale variation results,
and rarely estimating natural and anthropogenic aerosols (dust and anthropogenic emissions) separately; Insufficient
consideration of regional meteorological conditions on the concentration of atmospheric pollutants, it is necessary to consider
125 the impact of both horizontal and vertical mixing on air pollution; Currently, there is a lack of effective quantitative assessment
of the sources and sinks (accumulation and removal rates) of atmospheric pollutants. Most importantly, urban aerosol fluxes
largely depend on site characteristics, so more urban particulate matter flux data from different cities are needed to fully
understand the role of urban pollution sources and their spatial variability.

This paper focuses on an air pollution event in Beijing during winter and conducts a detailed analysis of the impact of
130 wind direction shear on aerosol transport within a short period of time. Polarization lidar and Doppler wind profile lidar were
used to obtain the vertical and horizontal aerosol mass concentration flux profiles to consider the influence of local aerosol
transport mechanisms on air pollution. Although only one special case is selected in this study, it represents a
complete and typical pollution cycle driven by the complex interplay of meteorological factors and boundary layer

dynamics, which constitutes the core focus of our research. This case is possible to elucidate the fundamental mechanisms governing aerosol transport under the typical monsoon during winter over the North China Plain (NCP). The observation results can be used to accurately reflect the internal changes in a specific area, and ultimately provide technical means and data references for early warning of urban air pollution and assessment of air pollution control measures, such as in large cities like Beijing. The northwest of Beijing is mountainous, and the southeast is a plain. The plain slopes gently from northwest to southeast, with the height gradually decreasing. Due to the influence of topography and climate, a large amount of air pollutants is not only emitted in the urban area of Beijing, but also in the suburbs, rural areas, and surrounding cities. The mutual exchange of air pollutants among these cities makes the understanding of the formation of air pollution and the formulation of control strategies more complex. As a hot spot representative of the central area of the Beijing-Tianjin-Hebei region, the emission characteristics and meteorological conditions of the urban area of Beijing are different from those of other urban agglomerations. Therefore, the aerosol flux may differ from the observation results of other urban agglomerations.

2 Instrumentation and methodology

2.1 Doppler wind lidar

Wind lidar is the most effective method for global wind field information detection and can also obtain wind field information within the three-dimensional profile. In this study, a Doppler wind lidar system was used to monitor the vertical profile of the wind field. The detection range of this system is 3000 m, with a distance resolution of 50 m and a time resolution of 1 minute. During the long-term observation period, the lidar system was set to the velocity azimuth display (VAD) scanning mode, with a fixed elevation angle of 60° , an azimuth range of 0° to 300° , a scanning interval of 5° , and a total of 60 radial profiles were obtained for each scanning circle, lasting for 135 seconds. Then, based on the assumption of a horizontally uniform wind field, the horizontal wind speed, wind direction, and vertical wind speed were extracted from the measured radial velocities at different azimuth angles (Smalikho, 2003; Banakh et al., 2013). Due to physical obstructions at the instrument site, the laser transmission and reception path of the lidar was partially blocked by nearby structures, making effective detection impossible within the 300° to 360° sector. However, the absence of data from a 60° sector has a negligible impact on the accuracy of the wind profile retrieval—the uncertainty. Consequently, the calculated horizontal aerosol fluxes are not significantly affected.

2.2 Raman–Mie scattering aerosol LiDAR (REAL)

REAL is a single-wavelength polarized Raman-Mie scattering aerosol lidar with a wavelength of 532 nm, featuring a groundbreaking design that achieves a near-surface blind zone of less than 50 meters—significantly below the industry-typical blind zone of over 100 meters—which provides a crucial advantage for high-precision joint analysis with ground-based

165 observational data. The system's receiver is equipped with six channels, among which the core channels include the 532 nm
parallel polarization channel (P), the 532 nm perpendicular polarization channel (S), and the 607 nm nitrogen Raman channel.
This configuration, as detailed by Chen et al. (2024) regarding system parameters, enables the REAL to possess continuous
observation capabilities from the ground to the upper atmosphere. To ensure the dynamic range of the data from the three main
receiving channels, each main channel is divided into high and low sub-channels through a non-polarizing beam splitter and
170 data gluing is achieved through linear overlap (Zhang et al., 2014). To enhance the dynamic range of the lidar echo signal, in
the system design, optimization was made in terms of stray light, anti-electromagnetic interference in the detection circuit, and
polarization isolation. After the design was completed, it passed a series of tests of mutual comparison with the 532 nm
reference lidar MUSA (Wandinger et al., 2016) of the EARLINET lidar calibration center with high quality. In the mutual
comparison, it was found that the signal deviation of REAL and MUSA within the detection dynamic range was less than 2%,
175 and the dynamic detection range was larger.

In this research, the PM_{2.5} and PM₁₀ data employed were primarily sourced from the official website of the China National
Environmental Monitoring Centre (<http://www.cnemc.cn/>). This website offers real-time dissemination and historical data
query capabilities for air quality monitoring data across different regions in China. Furthermore, air quality-related information
(Beijing's PM concentration) published on the official website of the Beijing Municipal Ecology and Environment Bureau
180 (<http://sthj.beijing.gov.cn/>) was also consulted for supplementation and verification purposes. When analyzing the direction
and backward trajectories of long-range transported aerosols, we utilized the Hybrid Single-Particle Lagrangian Integrated
Trajectory (HYSPLIT) model (<https://ready.arl.noaa.gov/HYSPLIT.php>). The HYSPLIT model has been extensively applied
in calculating the trajectories of air quality, pollutants, and their transport processes (Kim et al., 2020).

2.3 Data processing

185 In this study, the aerosol optical properties were retrieved using a 532-nm single-wavelength Raman-Mie polarization
scattering lidar (REAL) system, and the wind profiles were obtained by means of a Doppler wind lidar. The raw dataset
measured by the REAL is sampled within 1 minute, with a vertical resolution of 15 m. Before retrieving aerosol optical
properties, these data undergo preprocessing steps, including dead time correction, background noise subtraction, signal gluing
and averaging, range correction, and polarization gain ratio calibration to ensure the quality of the data (Freudenthaler, 2016;
190 Wang et al., 2023; Mao et al., 2024). The aerosol backscattering coefficient is retrieved using the sum of the horizontally and
vertically polarized Mie scattering signals (D'Amico et al., 2015).

Since the presence of clouds can severely bias the retrieval of aerosol optical properties, cloud-affected range bins were
identified and masked before inversion using the Value Distribution Equalization (VDE) algorithm originally proposed for
ground-based lidar (Zhao et al., 2014). In VDE, semi-discretization is applied to suppress random noise while preserving sharp
195 layer gradients, followed by histogram-equalization to enhance far-range signal visibility; empirical gradient/attenuation tests
are then used to delineate cloud layers. The practical reliability of this screening under diverse atmospheric conditions has

been demonstrated in long-term, multi-instrument evaluations (Zou et al., 2024). Only cloud-free bins were used for the subsequent aerosol inversions and particle depolarization ratio (PDR)-based classification.

The separation of dust and non-dust aerosols was achieved following the Polarization Lidar Photometer Retrieval (POLIPHON) method (Freudenthaler et al., 2009; Mamouri & Ansmann, 2014), which utilizes the PDR as the key parameter.

$$\delta_p = \frac{(1+\delta_m)(\delta_v+1)\beta_a}{\beta_m(\delta_m-\delta_p)+(1+\delta_m)\beta_a} - 1, \quad (1)$$

In the text, δ_m represents the depolarization ratio of the molecules, δ_v denotes the volume depolarization ratio, and β_a and β_m are the backscattering coefficients of aerosols and molecules, respectively. Subsequently, classification thresholds are set: when $\delta_p \geq 0.31$, it is identified as pure dust aerosol (δ_{dust}); when $\delta_p \leq 0.05$, it is identified as non-dust aerosol ($\delta_{non-dust}$); the intermediate range is classified as a mixture of dust and other aerosols. Following the classification, the non-dust aerosol component is primarily attributed to anthropogenic emissions within the study context. Beijing is an inland megacity with no significant local sources of sea salt aerosols and is not influenced by volcanic emissions (Zhang et al., 2013; Huang et al., 2014; Liu et al., 2017). Therefore, for this study focusing on the Beijing region, the term "non-dust aerosol" serves as a reasonable proxy for anthropogenic aerosol, allowing us to investigate the transport characteristics of pollutants originating from local human activities separately from transported dust. The backscattering coefficient of dust is calculated following these rules: In a pure dust region, the total backscattering coefficient is solely contributed by dust. In a non-dust region, the contribution of dust is zero. In a mixed region, based on the linear mixing model, the contribution of dust is proportionally distributed according to the PDR. (Tesche et al., 2009; He et al., 2023, 2025):

$$\beta_{dust} = \beta_a \cdot \frac{\delta_p - \delta_{non-dust}}{\delta_{dust} - \delta_{non-dust}} \cdot \frac{1 + \delta_{dust}}{1 + \delta_p}, \quad (2)$$

The total backscattering coefficient minus the backscattering coefficient of dust yields the non-dust backscattering coefficient.

The aerosol mass concentration is derived via the joint calculation of the extinction coefficient and the mass concentration conversion factor (Mamouri & Ansmann, 2014).

$$m(z) = \beta_{non-dust/dust}(z) \cdot LR \cdot c, \quad (3)$$

Among them, the aerosol extinction coefficient which is calculated by dust and non-dust backscattering coefficient with assumed fixed lidar ratio (LR), and c represents the mass concentration conversion factor. In this study, the mass concentration conversion factor c for dust aerosols is adopted as $2.002 \times 10^{-3} \text{ kg} \cdot \text{m}^{-2}$, while that for non-dust aerosols is taken as $0.6355 \times 10^{-3} \text{ kg} \cdot \text{m}^{-2}$ (Mamouri & Ansmann, 2017).

The fluctuations of aerosol mass concentration (m') and vertical wind speed (w') for eddy covariance calculations were derived from datasets with a 1-minute temporal resolution. This resolution represents a balance between resolving the dominant turbulent eddies responsible for vertical transport and ensuring robust covariance statistics from the lidar measurements. The

Doppler wind lidar acquires radial wind velocities at a native frequency of 1 Hz, which are integrated over its 135-second VAD scanning cycle to retrieve wind profiles, resulting in a final 1-minute output. The aerosol optical properties from the REAL lidar are also provided at a 1-minute resolution. The spatial volume for each data point is defined by a 50-meter vertical range resolution and the horizontal averaging inherent to the VAD scan. This configuration (1-min, 50-m) is a standard and well-validated approach in lidar-based boundary layer flux studies. It effectively captures the energy-containing eddies that dominate vertical transport. While some attenuation of the flux may occur due to the smoothing of the smallest-scale turbulence by spatial and temporal averaging, the method robustly captures the principal patterns and vertical structure of the aerosol fluxes.

To ensure spatial alignment between the aerosol data and wind field data during flux calculations, we resampled the aerosol mass concentration data retrieved by the REAL to a vertical resolution of 50 m (consistent with the wind field data) before performing the covariance calculations. The vertical aerosol mass flux is calculated using the eddy covariance method (Swinbank, 1951). The eddy covariance method is a direct measurement method for vertical fluxes based on turbulence theory. Thus, the aerosol mass concentration vertical fluxes at different heights can be expressed as follows (Engelmann et al., 2008; Dipu et al., 2013):

$$F_m = \overline{m'(z)w'(z)}, \quad (4)$$

When calculating the vertical flux, the instantaneous fluctuation of the vertical wind speed at a given altitude is defined such that upward is positive and downward is negative. A positive vertical flux implies that the vertical motion is responsible for aerosol transport, while a negative vertical flux indicates that the vertical motion is involved in aerosol scavenging. While the aerosol mass concentration horizontal fluxes at different heights can be expressed as follows (Hu et al., 2022; Lin et al., 2023; Li et al., 2024):

$$F_h = \overline{m(z)} \times \overline{v(z)\cos(wd)} + \overline{m'(z)v'(z)}, \quad (5)$$

Among them, z represents the vertical range of the lidar. The time-averaged value (30 minutes) is calculated by computing the covariance between the specific aerosol mass concentration $m'(z)$ and the fluctuation $w'(z)$ of the vertical wind speed at different altitudes caused by turbulence; the time-averaged horizontal wind speed $\overline{v(z)}$ and aerosol concentration $\overline{m(z)}$ (30 minutes) are used for calculation. In addition, the covariance between the specific aerosol mass concentration $m'(z)$ and the fluctuation $v'(z)$ of the horizontal wind speed at different altitudes caused by turbulence is also considered in the horizontal fluxes in this study. When computing the horizontal flux, we decompose the horizontal wind speed into components in the north-south direction $\cos(wd)$, where the northwestward direction is defined as positive. When there is a positive correlation between the wind direction and the fluctuations in aerosol mass concentration, the horizontal flux is positive, signifying the northward transport of aerosols.

The column of both fluxes is calculated as the product of the transport flux per unit cross-sectional area and the sum of the heights of each layer (Hu et al., 2022):

$$F_c = \sum F_m \times z_i, \quad (6)$$

260 Among them, z_i represents the height interval of each layer. This study further calculates the emission rates of dust and non-dust aerosols, which are derived by dividing the aerosol flux by their respective mass concentrations, thereby directly reflecting the transport efficiency per unit concentration of aerosols.

3 Result Analysis

The observation site is located at the Beijing South Suburban Observatory (latitude: 39°54'20" N, longitude: 116°25'29" E). The observation period is selected from 19:30 China Standard Time (CST) on November 2, 2023 to 9:30 CST on November 4, 2023, with an interval of 38 hours. Throughout the observation, the sky was predominantly clear or had sparse cloud cover. Only the contributions of optical properties caused by aerosols are used so that ensured the continuous acquisition of aerosol vertical profiles and wind field information over the free troposphere.

270 Fig. 1 presents a comprehensive overview of the in-situ and lidar observation results over Beijing from 19:30 on November 2, 2023, to 09:30 on November 4, 2023 (CST). It integrates near-surface particulate matter concentrations with vertically resolved atmospheric profiles to illustrate the temporal evolution of a winter haze event. The figure is composed of six panels (a-f) that collectively depict the key parameters driving aerosol dynamics.

The event commenced with the arrival of a pronounced dust layer. Starting around 19:30 CST on November 2nd, the lidar observations revealed a high-aerosol-loading layer below 1.5 km (Fig. 1(b)), as indicated by the enhanced range-corrected signal at 532 nm (RCS532). This layer was characterized by a significantly elevated particle depolarization ratio (δ_p) of 0.3-0.4 (Fig. 1(c)), unequivocally identifying the presence of non-spherical dust aerosols (Peng et al., 2021; Floutsi et al., 2023; Tesche et al., 2009). This was synchronized with a drop in the near-surface $PM_{2.5}/PM_{10}$ ratio to below 0.1. The meteorological driver was a strong northwesterly flow (Fig. 1(d, f)) with wind speeds exceeding 10 m/s above 500 m, which transported dust from the northwest (confirmed by HYSPLIT back-trajectories in Fig. 4(a)). The boundary layer height (PBLH) was elevated 280 (~1.5 km), and strong vertical mixing was observed (Fig. 1(e)). Thus, it is inferred that the accumulation and subsequent dissipation of local pollution during this period were jointly influenced by long-range transported dust particles and local emissions.

A pivotal transition occurred around 02:30 CST on November 3rd. The low-level wind direction below 1 km abruptly shifted from northwesterly (NW) to northeasterly (NE) (Fig. 1(f)). The boundary layer height was compressed close to the ground (<0.5 km), consistent with observations of weaker vertical exchange rates below 500 meters (Fig. 1 (e)). This is likely 285 due to the change in near-surface temperature caused by the cold air from the north, which reduced vertical turbulence and thus inhibited the vertical development of the boundary layer (Seinfeld & Pandis, 2016). On the other hand, the northerly

airflow played a crucial role in the horizontal transport of local pollutants, leading to a rapid decrease in overall aerosol concentration. With the wind shear and the weakening of vertical turbulence, the boundary layer lost its dust source, and the aerosol type shifted from dust-dominated to locally emitted fine particles.

During the daytime of Nov 3, solar heating elevated the PBLH back to ~ 1.5 km. The wind field became complex: southerly winds prevailed near the surface, while a strong wind shear existed around 1-2 km altitude, with westerly/southwesterly winds aloft (Fig. 1(d, f)). This structure facilitated the vertical mixing of residual dust aloft and local anthropogenic emissions, resulting in a well-mixed aerosol layer with a moderate δ_p of ~ 0.2 below 1.5 km (Fig. 1(c)). Notably, a layer of dust ($\delta_p > 0.3$) persisted atop the boundary layer (1.5-2 km), indicating continuous advection.

Throughout the entire observation period, the core of the westerly jet stream, which existed at an altitude of approximately 1000 meters with an intensity of $15 \text{ m}\cdot\text{s}^{-1}$ on the night of November 2nd, significantly weakened and dissipated by the day on November 3rd. However, a westerly jet structure with adjusted intensity and position (around 2000-2500 meters, $12\text{-}15 \text{ m}\cdot\text{s}^{-1}$) was reestablished on the night of November 3rd. Regarding the near-surface wind field, the prevailing wind direction steadily shifted to southerly/southwesterly (S/SW) after 06:00 CST on November 3rd. This change in wind direction, coinciding with the typical timing following a cold front passage, indicates that the local influence of warm advection played a role after the cold front had passed. Lastly, a significant vertical discontinuity in the wind field consistently existed near 1000 meters, characterized by variable wind directions and strong vertical wind speed gradients at this altitude. The most intense shear event occurred around 18:00 CST on November 3rd, with a wind speed difference exceeding $10 \text{ m}\cdot\text{s}^{-1}$ observed within just 500 meters of vertical distance.

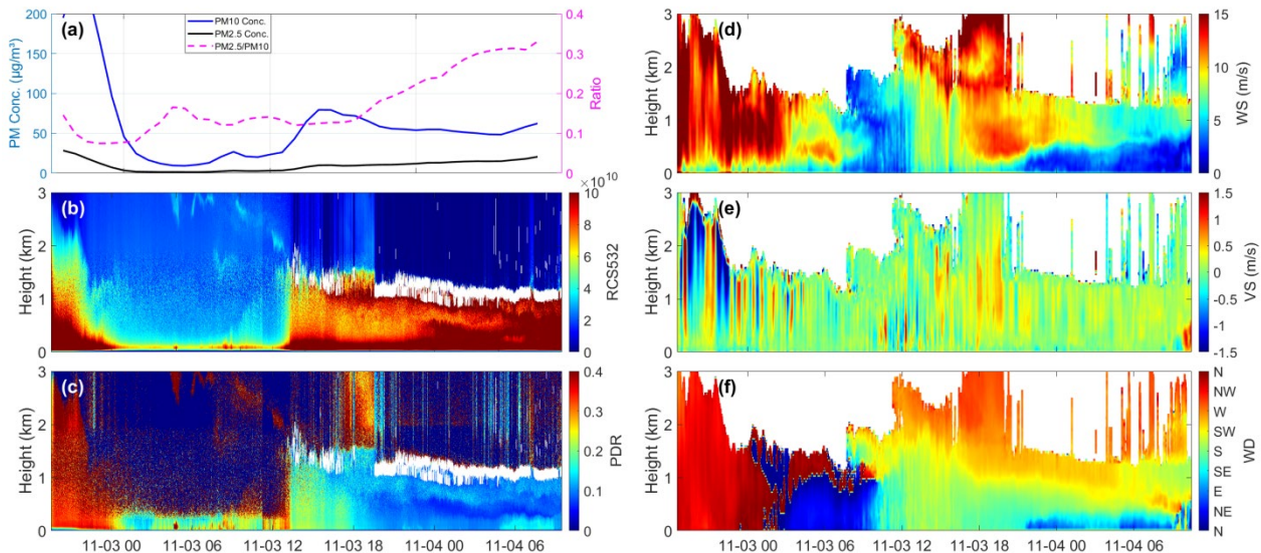
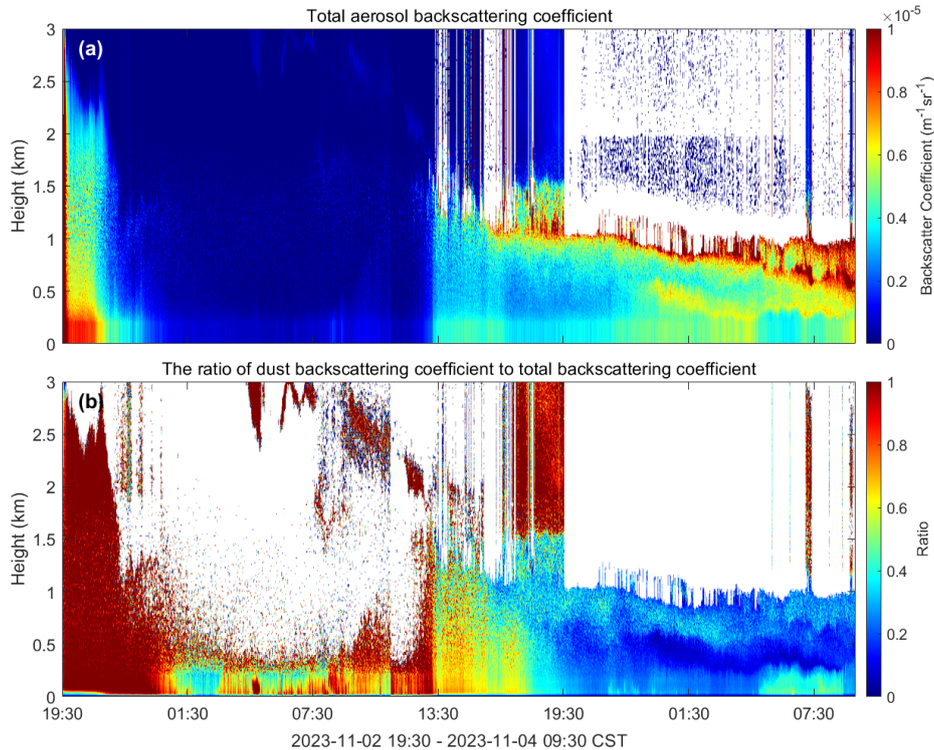


Figure 1: In-situ and lidar observation results over Beijing from 19:30 on November 2, 2023, to 09:30 on November 4, 2023 (CST). (a) Near-surface PM_{2.5} and PM₁₀ concentrations and their ratio from 19:30 on November 2, 2023, to 09:30 on November 4, 2023; (b) Observational results of the temporal and spatial distribution of the REAL and wind profiler lidar from 19:30 on November 2, 2023, to 09:30 on November 4, 2023, in the 0-3 km atmospheric layer over Beijing (b) 532 nm lidar range-corrected signal (RCS, representing vertical distribution information of relative aerosol loadings); (c) 532 nm particle depolarization ratio; (d) Temporal

and spatial distribution of horizontal wind speed; (e) Temporal and spatial distribution of vertical velocity; (f) Temporal and spatial distribution of wind direction. The white areas in (b) and (c) represent cloud layers identified by the Vertical Distribution Equalization (VDE) method, and the white areas in (d), (e) and (f) represent the absence of observation datasets.

315 During the period of rapid pollutant clearance (before 2:00 CST on November 3rd), both the atmospheric boundary layer height and wind direction were crucial for the efficient removal of pollutants. The decrease in boundary layer height prevented the mixing of dust aerosols from above into the surface layer, and the strong northwesterly winds facilitated the rapid dispersion of anthropogenic pollutants. Subsequently, the wind shear from northwest to southeast directions cut off the source of dust aerosols, while the elevated boundary layer height also contributed to the dilution of anthropogenically emitted aerosols.

320 Fig. 2(a) illustrates the spatio-temporal evolution of the total aerosol backscattering coefficient at 532 nm from 19:30 CST on November 2 to 09:30 CST on November 4, 2023. It reveals the complex vertical structural variations of aerosol loadings during this pollution event: at the initial stage of pollution (evening of November 2 to early morning of November 3), significant high-value (about $1 \times 10^{-5} \text{ m}^{-1} \cdot \text{sr}^{-1}$) appeared at low altitudes (0-1.5 km), indicating the accumulation of aerosols in the near-surface layer; subsequently, with changes in meteorological conditions, a strong aerosol layer (backscattering coefficient over $1 \times 10^{-5} \text{ m}^{-1} \cdot \text{sr}^{-1}$) emerged at higher altitudes (1.5-2.5 km), while the aerosol backscattering coefficient near-surface layer is lower to $0.1 \times 10^{-5} \text{ m}^{-1} \cdot \text{sr}^{-1}$.



330 **Figure 2: (a) the spatial-temporal evolution of the total aerosol backscattering coefficient at 532 nm wavelength from 19:30 on November 2, 2023, to 09:30 on November 4 (CST). (b) the ratio of dust backscattering coefficient to total backscattering coefficient. The white areas indicate the absence information.**

To gain an in-depth understanding of the transport behavior of aerosols from different sources, we have separated dust and non-dust aerosols following the POLIPHON method based on the depolarization signal (Fig. 2(b)). It indicates that dust aerosols were primarily concentrated in the elevated layer (approximately 1-3 km), although they were occasionally mixed down to near the surface (e.g., on November 2 and around 12:00 CST on November 3). In contrast, non-dust aerosols were
335 predominantly found in the lower atmospheric layer (below 1 km), which is mainly attributed to local anthropogenic emissions in the North China Plain. It noted that the spatial footprint of the measurement is height-dependent, meaning the influence of near-surface local sources diminishes with increasing altitude.

To investigate the regional transport of aerosols, the temporal variations in PM concentrations in Beijing and its surrounding cities during this period were analyzed (Fig. 3). On November 2nd, the five cities simultaneously reached peak
340 $PM_{2.5}$ concentrations, indicating a regional fine particulate matter pollution event. The $PM_{2.5}/PM_{10}$ ratio exceeded 0.6 during the peak period in all cities, confirming that the pollution was primarily composed of $PM_{2.5}$. Zhangjiakou first experienced an explosive increase in PM_{10} at 10:00 on November 2nd, followed by subsequent pollution peaks in Shijiazhuang, Tianjin, and Chengde, revealing a dust transport pathway from northwest to southeast. Subsequently, both PM_{10} and $PM_{2.5}$ concentrations in the five cities showed a rapid decline, and the $PM_{2.5}/PM_{10}$ ratio also decreased to varying degrees, reflecting the effective
345 clearance of local pollutants by strong meteorological processes.

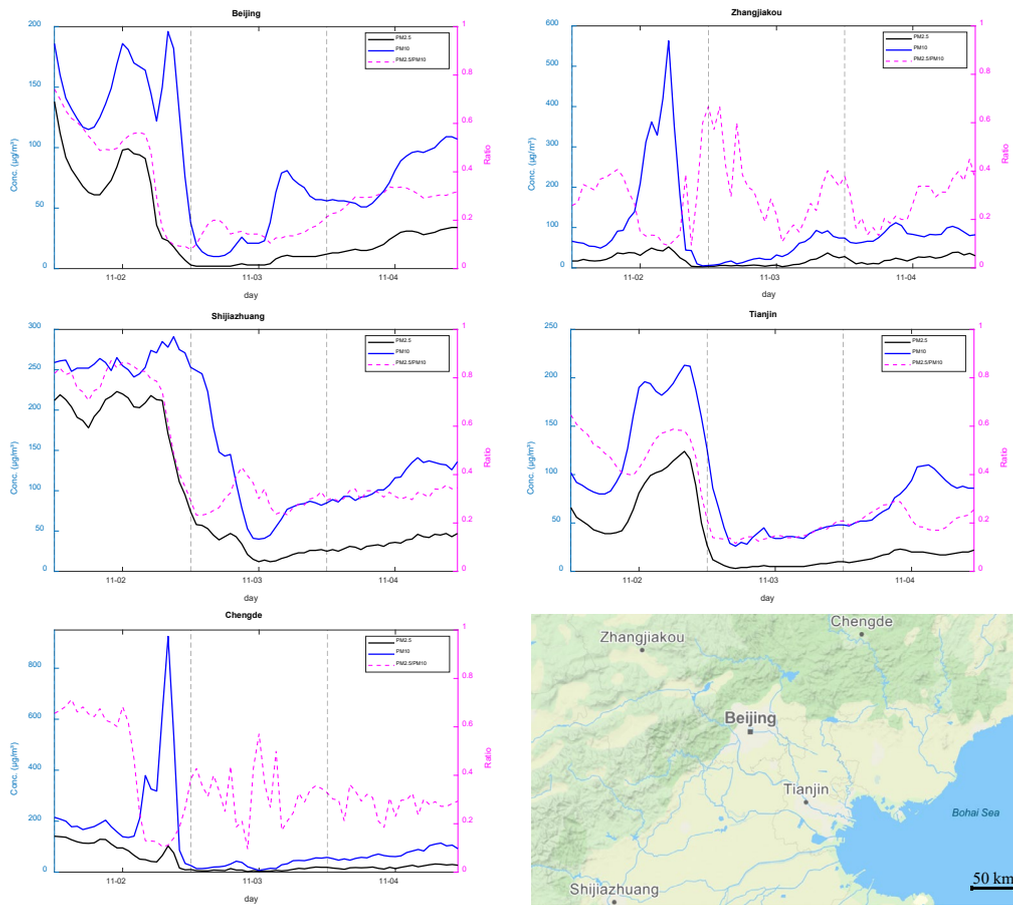
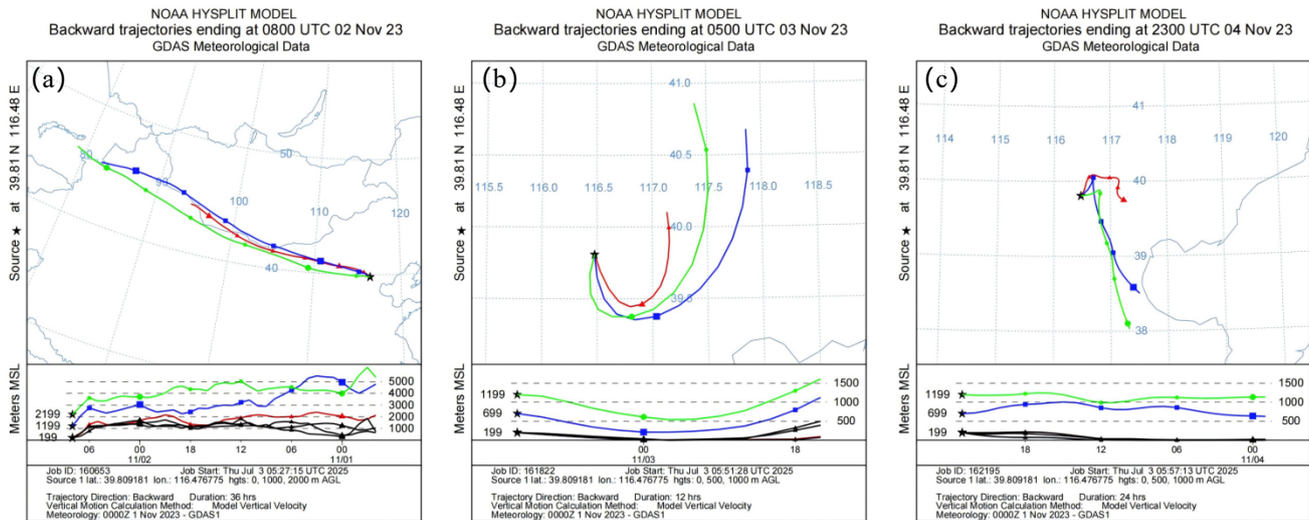


Figure 3: The hourly air quality trends of major cities in the Beijing-Tianjin-Hebei region (Beijing, Tianjin, Zhangjiakou, Chengde) from November 2 to 4, 2023 are presented. The map (© Mapbox, available at <https://www.mapbox.com/mapbox-studio>) illustrates the geographical distribution of the cities, with surrounding subplots showing the daily variation patterns of PM_{2.5} (black line), PM₁₀ (blue line) concentrations, and the PM_{2.5}/PM₁₀ ratio (pink dashed line) for each city. The concentration units are micrograms per cubic meter ($\mu\text{g}/\text{m}^3$), and the time span is from 0:00 CST to 23:00 CST daily.

350



355 **Figure 4: HYSPLIT modeled 48 h air mass backward trajectories for Beijing on 02-04 November 2023. (a) HYSPLIT modeled 36 h on 02 November 2023 at 00:00 China Standard Time (CST). All altitudes (green, blue, red and black), the trajectories mainly originated above the Taklimakan Desert. (b) HYSPLIT modeled 12 h on 02 November 2023 at 21:00 CST. All altitudes (green, blue, red and black), the trajectories mainly originated above the North China Plain. (c) HYSPLIT modeled 24 h on 04 November 2023 at 15:00 CST. At higher altitudes (green), the trajectories mainly originated above the North China Plain, while at lower altitudes (blue and red) they originated above the Taklimakan Desert.**

360 Fig. 4 shows the backward trajectories consistent with the wind direction results from the HYSPLIT model and wind profiler lidar observations, revealing the sources of aerosols within the aerosol layer during the air pollution event in Beijing, November 2023. From Fig. 4(a) (16:00 CST on November 2), it can be observed that the air mass trajectory follows a northwest-southeast path, originating from southern Mongolia and western Inner Mongolia, China (39°~41°N, 114°~116°E), indicating that the air mass during this period was primarily transported from the northwest, consistent with the contemporaneous wind field characterized by "predominantly northwesterly winds." In Fig. 4(b) (Beijing Time, 13:00 on 365 November 3), there is a dramatic change in the trajectory pattern, with all altitude levels turning southwestward. The air mass below 1000 meters originates from the southern North China Plain (35°N, 115°E), while the 1500-3000 meter layer recirculates inland from the Yellow Sea (34°N, 122°E), reflecting the warm and moist air mass along the coastal return flow. The trajectory at 500 meters altitude passed over the Bohai Bay (38°N, 120°E) 12 hours prior, suggesting a relatively clean air mass, although the possibility of sea salt particles being mixed in cannot be ruled out. In Fig. 4(c) (07:00 CST on November 5), the trajectories at all altitudes exhibit a southeast-northwest recirculation trend, with the lower layer (199~699 m) air mass originating from 370 the eastern North China Plain (38°~40°N, 117°~119°E), driven northwestward by southerly winds. The upper layer (1500~2199 m) trajectories extend southwestward, indicating the reverse transport after the recovery of the northwesterly winds at high altitudes. Combined with the observations of particle depolarization, it can be confirmed that dust aerosols primarily originate from the northwest (Taklamakan Desert), while non-dust aerosols mainly come from anthropogenic 375 emission sources in the North China region.

To quantitatively describe the aerosol transport flux during this event, we obtained the horizontal and vertical fluxes of aerosols by combining the estimated dust/non-dust aerosol mass concentrations with wind profile data (as shown in Fig. 5). The flux evolution clearly aligns with the three meteorological phases identified in Fig. 1.

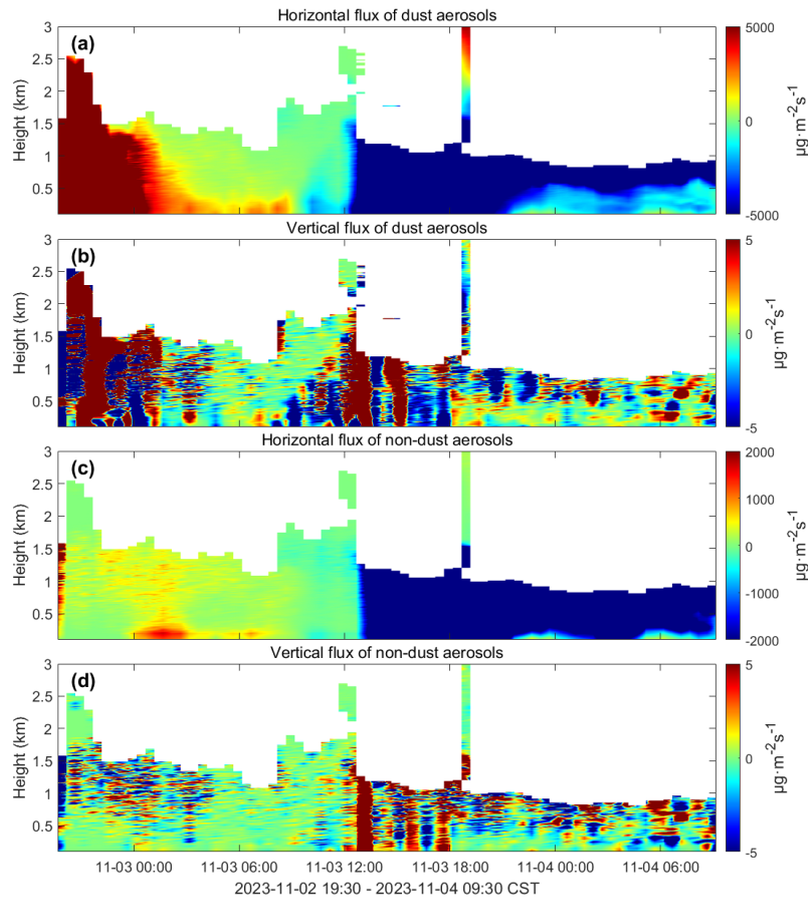
380 During Phase 1 (evening of Nov 2 to ~ 02:30 CST on Nov 3), dominated by strong northwesterly winds, both dust and non-dust aerosols were transported southeastward, as indicated by strong positive horizontal fluxes (Fig. 5(a, c)). The horizontal flux of dust aerosols peaked at approximately $5000 \mu\text{g}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ in the elevated layer. Concurrently, intense upward vertical fluxes of dust aerosols exceeded $5 \mu\text{g}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ (Fig. 5(b)), corresponding to the period of strong turbulent mixing within the deep boundary layer.

385 The pivotal transition in Phase 2 (~02:30 CST on Nov 3) is markedly reflected in the flux data. The abrupt shift in low-level wind direction to northeasterly directly triggered a sharp reversal of the dust horizontal flux from positive to near zero within an hour ((Fig. 5(a)), effectively cutting off the dust supply. This phenomenon confirms the modulating effect of wind shear on pollutant clearance proposed by Zhang et al. (2020), but this study further reveals its impact on the transport mechanism of aerosol types. This was accompanied by a shift of the dust vertical flux to significant deposition (approximately $-4 \mu\text{g}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$), reflecting scavenging due to boundary layer compression and weakened turbulence.

390 In Phase 3 (daytime on Nov 3), the horizontal flux of dust aerosols dramatically shifted to strong negative values (approximately $-5000 \mu\text{g}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$) below 2 km around noon (Fig. 5(a)), indicating northwestward transport of dust that was likely recirculated back as wind directions became complex. Meanwhile, the redevelopment of the boundary layer led to a second peak in the upward vertical flux of dust (over $5 \mu\text{g}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$, Fig. 5(b)), indicating secondary lifting. Throughout this period, the vertical flux of non-dust aerosols remained slightly positive, except for a brief deposition event around 14:00 CST
395 at 800 m altitude (Fig. 5(d)), underscoring that fine particles mainly undergo upward diffusion, contrasting sharply with the intermittent, strong deposition of dust.

The most notable dust/non-dust aerosol flux events in Fig. 5 occurred during the evening of November 2nd to the early morning of November 3rd, as well as from the morning to noon on November 3rd. These periods correspond to the strong horizontal wind speed events and wind direction abrupt changes depicted in Fig. 1. Strong horizontal winds and strong
400 boundary layer turbulence mixing are the primary driving forces behind the lifting of aerosol from the ground and the transport of mixture dust and anthropogenic emission particles.

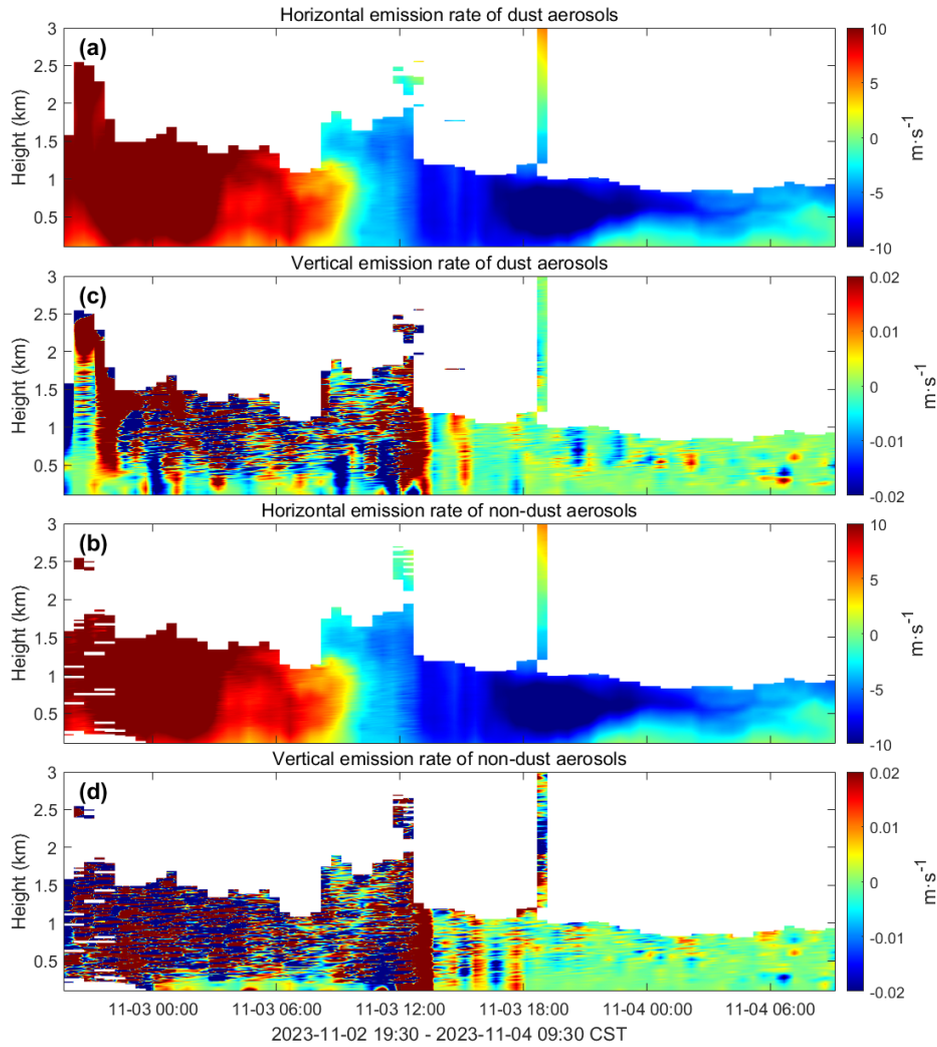
Compared to the study of near-surface aerosol fluxes (e.g., Casquero-Vera et al., 2022; Pryor et al., 2008; Yuan et al., 2019), the vertical cross-sectional aerosol fluxes in this research are particularly useful for understanding the inter-regional transport of aerosols. However, if similar lidar devices were deployed in multiple cities across the Beijing-Tianjin-Hebei region,
405 it would be possible to further quantitatively analyze the mutual transport of aerosols between cities. Regrettably, due to the limitations in observation duration, this study was unable to utilize long-term observational data to investigate pollution processes and pollutant transport characteristics under various weather patterns.



410 **Figure 5: Aerosol flux results from 19:30 on November 2, 2023, to 09:30 on November 4, 2023 (China Standard Time, CST), in the 0.1-3 km atmospheric layer over Beijing. (a) Horizontal flux of dust aerosols; (b) Vertical flux of dust aerosols; (c) Horizontal flux of non-dust aerosols; (d) Vertical flux of non-dust aerosols.**

Fig. 6 illustrates the spatial-temporal distribution of emission rates of dust and non-dust aerosols in the horizontal and vertical directions over the Beijing region from 19:30 CST on November 2 to 09:30 CST on November 4, 2023. The horizontal emission rate of dust aerosols (Fig. 6(a)) fluctuated significantly due to abrupt changes in the wind field. In the evening of November 2, driven by strong northwesterly winds, the horizontal emission rate showed a notable positive value (approximately $8-10 \text{ m}\cdot\text{s}^{-1}$), indicating transport of dust towards the southeast. Early on November 3, with the abrupt shift of the low-level wind direction to northeasterly, the horizontal emission rate sharply reversed to a strong negative value ($< -0.01 \text{ m}\cdot\text{s}^{-1}$), signifying a sudden increase in the efficiency of transporting dust northwestward. This drastic reversal in emission rates directly explains the rapid changes in the distribution of the high particle depolarization ratio layer observed in Fig. 1(c), as well as the abrupt decrease in near-surface PM_{10} concentrations—strong transport efficiency rapidly removed local dust. The horizontal emission rate of non-dust aerosols (Fig. 6(c)) exhibited a trend highly synchronized with that of dust aerosols, underscoring their co-variation in response to the same wind-field forcing.

425 In the evening of November 2, the boundary layer height was approximately 1.5 km, with strong convection, and the vertical emission rate of dust aerosols exceeded $0.01 \text{ m}\cdot\text{s}^{-1}$, indicating that strong convection efficiently lifted dust. Early in the morning of November 3, the boundary layer height compressed to less than 0.5 km, with weaker convection, and the vertical emission rate shifted to around $-0.01 \text{ m}\cdot\text{s}^{-1}$, reflecting a dominant deposition process. In the afternoon, as the boundary layer rose again, a peak in positive emission rates reemerged.

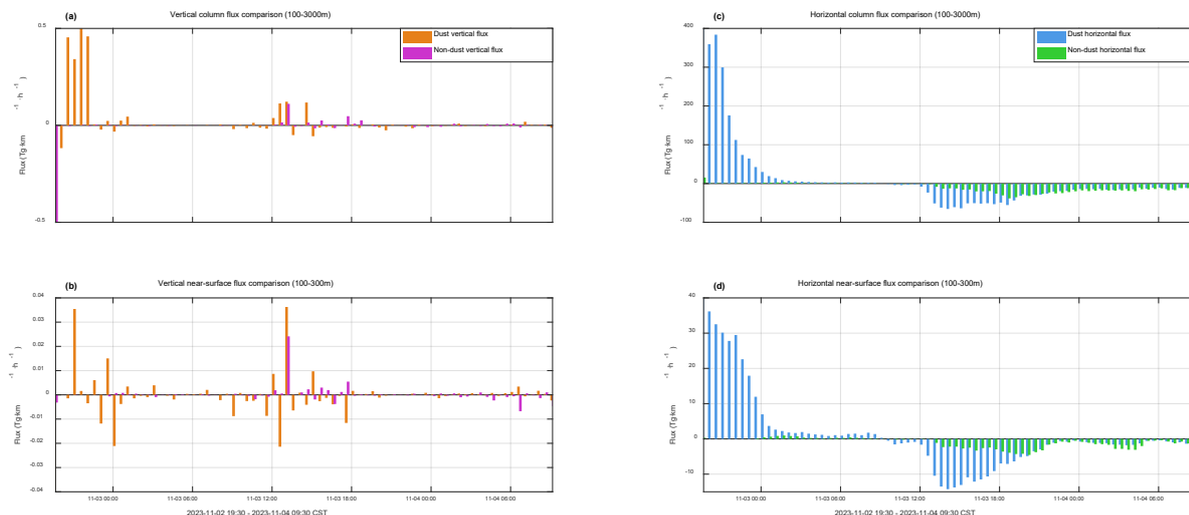


430 **Figure 6: Aerosol emission rate results from 19:30 on November 2, 2023, to 09:30 on November 4, 2023 (CST), in the 0.1-3 km atmospheric layer over Beijing. (a) Horizontal emission rate of dust aerosols; (b) Vertical emission rate of dust aerosols; (c) Horizontal emission rate of non-dust aerosols; (d) Vertical emission rate of non-dust aerosols.**

The most notable characteristic of the vertical emission rate of non-dust aerosols (Fig. 6(d)) was that it was mostly positive (0 to $0.02 \text{ m}\cdot\text{s}^{-1}$) throughout the observation period. This suggests that for non-dust aerosols, the vertical diffusion rate caused by convection was continuously greater than their minimal gravitational settling rate, resulting in a net upward transport. This

435 observation dynamically explains the slower decrease in $PM_{2.5}$ concentrations compared to PM_{10} in Fig. 1(a) and the phenomenon of pollution lingering, as fine particles, due to their extremely slow settling speed, are difficult to effectively remove through dry deposition and are more likely to be entrained and remain in the atmosphere for extended periods.

The results indicate a significant regulatory effect of meteorological conditions on aerosol emission rates: the diurnal variation in planetary boundary layer height drives the alternation between positive and negative vertical emission rates; 440 whereas abrupt changes in wind direction modulate the intensity of horizontal emission rates by altering the transport pathways.



445 **Figure 7: Distribution of aerosol column transport fluxes from November 2 to November 4, 2023. (a) Vertical column transport fluxes of aerosols within the atmospheric layer of 100-3000 meters; (b) Horizontal column transport fluxes of aerosols within the atmospheric layer of 100-3000 meters; (c) Vertical near-surface column transport fluxes of aerosols within the atmospheric layer of 100-300 meters; (d) Horizontal near-surface column transport fluxes of aerosols within the atmospheric layer of 100-300 meters.**

To analyze the changes in the near-surface and total fluxes of dust and non-dust aerosols during the diffusion process of this event, we obtained the column transport fluxes from 100-3000 meters (approximating the aerosol content throughout the entire atmospheric layer) and from 100-300 meters (representing the ground-level aerosol content) (Fig. 7). Negative non-dust 450 fluxes can be used to characterize the meteorological conditions' ability to clear locally emitted anthropogenic aerosols; positive non-dust fluxes indicate the meteorological conditions' capacity to accumulate locally emitted anthropogenic aerosols. Negative dust fluxes can be used to characterize the meteorological conditions' ability to clear already accumulated dust aerosols locally; positive dust fluxes indicate the meteorological conditions' capacity to accumulate dust aerosols transported to the local area.

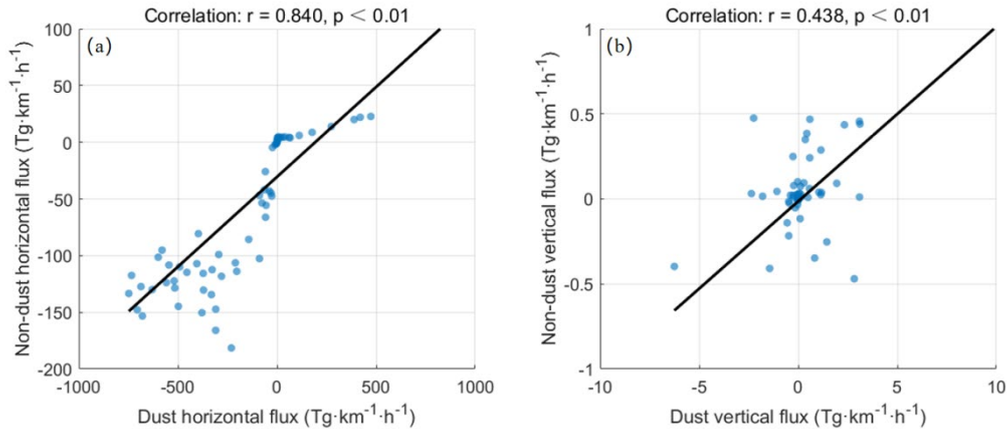
455 The comparison of near-surface vertical fluxes (100-300 m, Fig. 7 (b)) reveals distinct behaviors between dust and non-dust aerosols, indicative of their different sources. The vertical flux of dust exhibits strong intermittent deposition and occasional strong upward transport. In contrast, the vertical flux of non-dust aerosols remains consistently close to zero, with

much smaller amplitudes of both positive and negative fluctuations throughout the event. This indicates that in the near-surface layer, the net vertical exchange of locally emitted non-dust aerosols is minimal, whereas dust undergoes significant episodic deposition and lifting, driven by synoptic-scale processes. This pattern supports the interpretation that dust is primarily subject to long-range transport and dynamic meteorological forcing, while non-dust aerosols are dominated by local, near-surface emissions with limited net vertical exchange on the event scale.

Examining the column transport fluxes from 100-3000 meters (Fig. 7(a)), the highest value of positive vertical fluxes are about $0.5 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$ for dust aerosols and $0.15 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$ for non-dust aerosols, while the highest value of negative vertical fluxes are about $0.15 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$ for dust aerosols and $0.5 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$ for non-dust aerosols. It is observed that both dust and non-dust aerosol horizontal fluxes exceed their respective vertical fluxes, indicating that the horizontal dispersion of pollutants is more pronounced than their vertical dispersion. For the Southeastward horizontal transport (positive values shown in Fig. 7(c)), the maximal total column dust horizontal fluxes (100-3000 m) are estimated to be about $380 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$. In comparison, the maximal total column non-dust horizontal fluxes are found about 10 times smaller (about $30 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$). Regarding the near-surface level (Fig. 7(d)), the maximal column dust horizontal fluxes (100-300 m) are estimated to be about $36 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$. In comparison, the maximal column non-dust horizontal fluxes are found about 8 times smaller (about $3.5 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$). For the Northwestward horizontal transport (negative values shown in Fig. 7(c)), the maximal total column dust horizontal fluxes (100-3000 m) are estimated to be about $60 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$. In comparison, the maximal total column non-dust horizontal fluxes are found about 1-2 times smaller (about $40 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$). Compared with Southeastward horizontal transport, the Northwestward dust aerosols flux decreased significantly (approximately one-sixth of the Southeastward flux), while the non-dust aerosols flux slightly increased (about 1.3 times that of the southeastward flux). It may indicate that the long-range transported dust was mixed with local emissions in the downstream region of Beijing then impact the PM concentrations of Beijing. And also, it can indicate that the transportation cross-section in Southeastward is smaller than that in northwestward, which agrees with the terrain around Beijing region (Beijing has mountains in its northwest and plains in its southeast). In addition, to confirm with the near surface PM concentration over five cities (Fig. 4), it seems the aerosol transport cross-section probably reaches over 100 km levels. Regarding the near-surface level, the maximal column dust horizontal fluxes (100-300 m) are estimated to be about $14 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$. In comparison, the maximal column non-dust horizontal fluxes are found about 3 times smaller (about $4.5 \text{ Tg}\cdot\text{km}^{-1}\cdot\text{h}^{-1}$). Compared with Southeastward horizontal transport, it seems Northwestward horizontal transport brought 4 times less dust aerosols back but 1.2 times more non-dust aerosols. It indicates that the impact of local emissions is more than dust on the near surface. In general, a comparison between southeastward and northwestward air flows suggests that air pollutants over Beijing are more easily influenced by northwestward flows, which can bring about twice as much aerosol loading from the southeast region to Beijing.

The correlation analysis of aerosol fluxes reveals significant differences in the transport mechanisms of dust and non-dust components (Fig. 8). The horizontal fluxes exhibit a high positive correlation ($r = 0.856$, $p < 0.01$), indicating that dust and non-dust aerosols are controlled by the same atmospheric dynamical processes in the horizontal direction. In contrast, although the vertical fluxes show a positive correlation trend, the correlation is significantly weaker ($r = 0.438$, $p < 0.01$), suggesting a

big difference in their vertical transport processes. It is probably because one mainly comes from surface emission, while the other is mainly from sedimentation, and their emission rates are also comparatively different.



495 **Figure 8: Correlation between Dust and Non-Dust Aerosol Fluxes. (a) Scatter plot showing the correlation between dust horizontal flux and non-dust horizontal flux. (b) Scatter plot illustrating the correlation between dust vertical flux and non-dust vertical flux.**

4 Discussion and Conclusions

A strong dust transport event occurred from North-West China to the North China Plain in November 2023. During the transport periods, the Southeast-toward migration of cold air-masses carries not only the long-range transported dust but also local anthropogenic pollutants from the North China Plain to the Southeast region. This study quantitatively analyzed the vertical-horizontal flux structure of aerosols in the Beijing area using the polarization lidar and Doppler wind lidar observations on 2 November 19:00 CST to 4 November 9:30 CST, 2023. The main research findings are summarized as follows:

1. Meteorological factors exert a significant effect on aerosol concentrations

Temporal and spatial variations in wind direction and speed are the core drivers of horizontal aerosol transmission. From the evening of November 2 to the early morning of November 3, the low-altitude wind direction in Beijing shifted from Northwest to northeast, causing the horizontal transmission direction of dust aerosols to change from southeastward to northwestward. This confirms the path rule that "northwesterly winds push dust to spread southeastward, while northeasterly winds drive dust to transmit northwestward". Meanwhile, the dissipation and reconstruction of the upper-level westerly jet further affected the vertical transmission range of aerosols. In addition, the diurnal variation of the planetary boundary layer (PBL) height governs the vertical diffusion of aerosols by regulating turbulence intensity. When the boundary layer is stable at night, dust tends to settle and accumulate; during the day, the rising boundary layer, accompanied by strong turbulence, promotes the upward diffusion of aerosols, which is highly synchronized with the positive-negative transition of vertical emission rate (positive during uplift and negative during sedimentation). Aerosol concentrations control requires the synergy of "source reduction" and "meteorological factors. For a short period, meteorological factors are the dominant factors regulating aerosol concentration fluctuations, as they directly affect the transport and diffusion efficiency of

aerosols without changing emission intensity. Thus meteorological forecasting and early warning can be used to guide real regulation (e.g., enhanced emission reduction during stable boundary layer periods), in order to achieve effective control of aerosol concentrations.

2. There are fundamental differences in emission and diffusion characteristics between dust and non-dust aerosols.

520 The emission rate of dust aerosols fluctuates drastically, with their horizontal transmission significantly affected by sudden wind direction changes. Vertically, they exhibit an alternating "uplift-sedimentation" pattern and are mainly concentrated in the near-surface layer below 1000 meters, reflecting their close association with natural/anthropogenic activities such as surface dust emission. In contrast, non-dust aerosols (dominated by anthropogenically emitted fine particles) have a stable and low-intensity emission rate. Their horizontal transmission trend is consistent with that of dust but responds more gently.

525 Vertically, they primarily undergo weak upward diffusion, with rare large-scale sedimentation—characteristics that reflect the continuous emission from anthropogenic sources (e.g., industry and transportation). Moreover, due to their small particle size and slow sedimentation rate, fine particles are more likely to accumulate within the boundary layer.

3. The effectiveness of multi-source observation technologies and flux calculation methods is verified.

In this study, Real lidar was used to retrieve aerosol optical properties (e.g., particle depolarization ratio) for accurate

530 classification of dust and non-dust aerosols. Combined with Doppler wind lidar to obtain high-resolution wind profiles, the eddy covariance method was applied to calculate aerosol mass fluxes. This approach successfully captured the temporal and spatial distribution of emission rates of the two types of aerosols in horizontal and vertical directions. This technical system can effectively analyze the coupling mechanism of "meteorological conditions—pollution sources—diffusion processes" and provides a reliable method for quantifying anthropogenic aerosol emission rates in cities.

535 4. The synergistic impact of regional pollution transmission and local emissions is significant.

Observations of PM concentrations in cities around Beijing showed that on November 2, Zhangjiakou was the first to experience an explosive increase in PM₁₀ concentration, followed by pollution peaks in Shijiazhuang, Tianjin and other cities. This reveals the cross-regional transmission path of dust from northwest to southeast. Additionally, the peak PM_{2.5} concentration in Beijing corresponds to the stable emission of non-dust aerosols, indicating that the superposition of regional

540 transmission and local anthropogenic emissions is a key cause of this pollution event. Severe weather processes (e.g., cold air advection) have a significant scavenging effect on both types of pollutants, further verifying the potential of meteorological intervention in pollution control. The total amount of dust transmission rate is estimated to be about 1360 Tg·km⁻¹·h⁻¹ and the total amount of local pollutants transmission rate is estimated to be about 770 Tg·km⁻¹·h⁻¹ for the entire troposphere.

In conclusion, the established polarization-wind field joint inversion framework in this study overcomes the spatial

545 representativeness issues of traditional single-point flux observations. The aerosol type separation achieved by the depolarization ratio method enables the quantitative decomposition of the contributions to the aerosol flux. Combining both vertical and horizontal fluxes of aerosol can be used to estimate the total transport amount of dust along its trajectories and also the total transport amount between the cities. The aerosol type-specific flux observation technology developed in this study provides new insights for joint pollution prevention and control in urban agglomerations like the Beijing-Tianjin-Hebei

550 region and similar city clusters. This holds important reference value for the implementation of cross-regional coordinated air
pollution control in the context of the integrated development of the Beijing-Tianjin-Hebei and Yangtze River Delta regions.
It proves that using lidar remote sensing to study aerosol fluxes under different meteorological conditions (wind direction,
wind speed and boundary layer height) provides data suitable for a scientific basis for urban air pollution early warning and
555 the formulation of regional joint prevention and control strategies (e.g., northwest source control for dust and local emission
reduction for non-dust aerosols). Furthermore, the aerosol fluxes profiles can be useful to estimate the amount of pollutant
across-region transport. However, there are limitations: the observation duration is limited (38 hours), making it difficult to
cover fluxes characteristics under various weather patterns; the mass concentration conversion factor uses fixed values without
considering the hygroscopic growth effect during aerosol aging; and the non-dust contributions are assumed to be
560 anthropogenic emissions, of which the sources need to be further confirmed and a more accurate mass concentration conversion
factor is required. Moreover, the horizontal flux calculation does not incorporate multi-city network observations, limiting the
closure verification of regional transport fluxes. In the future, lidar observation networks can be deployed to track regional
transport fluxes in real-time and optimize emergency emission reduction strategies for heavy pollution weather. For instance,
when horizontal flux monitoring indicates an increase in dust input, dust suppression measures at construction sites can be
initiated in advance; whereas when the vertical flux of non-dust aerosols remains positive, it is necessary to strengthen the
565 control of industrial point sources and mobile sources.

Code and data availability. Code and data can be provided by the corresponding author on request.

Author contributions. All the authors made contributions to this research work and manuscript. In particular, LW, ZY, YH and
570 XW designed the whole strategy of this work. LW organized the observation campaign. ZT analyzed the data and wrote the
manuscript draft. SL, CY participated the data analysis. LW, ZY, YH, BZ, DC and XW participated in the scientific discussions,
reviewed and proofread the manuscript. TL participated in the observations and data collection. LL, BZ, DC, ZB, YC and XW
reviewed and proofread the manuscript. LW, ZY, YH, LL, XW and YC acquired the research funding and led the study. All
authors have read and agreed to the published version of the manuscript.

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580 *Competing interests.* The authors declare no conflict of interest.

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