

Reduced light absorption of black carbon (BC) and its influence on BC-boundary-layer interactions during “APEC Blue”

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Abstract. Light absorption and radiative forcing of black carbon (BC) is influenced by both BC itself and its interactions with other aerosol chemical compositions. Although the changes in BC concentrations in response to emission reduction measures have been well documented, the influence of emission reductions on the light absorption properties of BC and its influence on BC-boundary-layer interactions has been less explored. In this study, we used the online coupled WRF-Chem model to examine how emission control measures during APEC affect the mixing state/light absorption of BC, and the associated implications for BC-PBL interactions. We found that both the mass concentration of BC and the BC coating materials declined during the APEC week, which reduced the light absorption and light absorption enhancement (E_{ab}) of BC. The reduced absorption aerosol optical depth (AAOD) during APEC were caused by both the declines in mass concentration of BC itself (52.0%), and the lensing effect of BC (48.0%). The reductions in coating materials (39.4%) dominated the influence of lensing effect, and the reduced light absorption capability (E_{ab}) contributed 3.2% to the total reductions in AAOD. Reduced light absorption of BC due to emission control during APEC enhanced planetary boundary layer height (PBLH) by 8.2 m. Different responses of $PM_{2.5}$ and O_3 were found to the changes in light absorption of BC. Reduced light absorption of BC due to emission reductions decreased near surface $PM_{2.5}$ concentrations but enhanced near surface O_3 concentrations in the North China Plain. These results suggest that current measures to control SO_2 , NO_x , etc. would be efficient to reduce the absorption enhancement of BC, and to inhibit the feedback of BC on boundary layer. Yet

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enhanced ground O₃ might be a side effect of current emission control strategies. How to control emissions to offset this side effect of current emission control measures on O₃ should be an area of further focus.

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1 Introduction

Black carbon (BC) in the atmosphere is produced both naturally and by human activities, attributable to the incomplete combustion of hydrocarbons (*Bond et al., 2013; Ramanathan and Carmichael, 2008*). In addition to contributing to particulate matter and degraded air quality, it is the dominant absorber of visible solar radiation, playing a unique and pivotal role in the Earth's climate system (*Bond et al., 2013; Menon et al., 2002; Ramanathan and Carmichael, 2008; Yang et al., 2019*). The absorption of BC occurs not only in the atmosphere, but when it is deposited over snow or ice, it triggers positive feedbacks and exert a positive radiative forcing (*Flanner et al., 2007; Grieshop et al., 2009*). The direct radiative forcing of atmospheric black carbon was estimated to be 0.4 W m⁻² (0.05-0.8 W m⁻²) (*IPCC, 2014*), and BC has been targeted in emission control policies to mitigate both air pollution and global warming (*Grieshop et al., 2009*).

Before the 1950s, intense emissions of BC were concentrated in North America and Western Europe. In recent decades, South and East Asia have emerged to become major source regions (*Ramanathan and Carmichael, 2008*). BC emitted from China is responsible for a quarter of the total global emissions (*Bond et al., 2004*). Chemical transport model simulations suggest that the residential sector is the leading source for mass concentration of BC in China, followed by the industrial sector (*Li et al., 2016*). Mean BC direct radiative forcing in China is ~1.22 W m⁻², more than three times the global mean forcing (*Li et al., 2016*), with two-thirds to three fourths of which contributed by local emissions of BC in China, and the rest by emissions in other countries (*Li et al., 2016; Yang et al., 2017*).

Specific policies to address BC emissions have not been implemented in China, yet multiple measures targeting PM_{2.5} reduction have resulted in declines in BC (*Gao et al., 2018b; Yamineva and Liu, 2019*). A number of observational studies have revealed the declining trend of BC concentrations in China in recent years (*Ji et al., 2018, 2019a, 2019b; Qin et al., 2019*). From 2013 to 2018, the annual mean BC concentrations in Beijing declined from 4.0 µg m⁻³ to 2.6 µg m⁻³ (*Ji et al., 2019b*). Associated changes in BC radiative forcing can be expected from declines in mass concentration of BC in China, while the radiative forcing of BC is influenced also by the changes in other aerosol components.

BC absorption is closely connected with the aging process, which is defined as the interaction between BC and other aerosol chemical compositions (*Jacobson, 2001*). After being emitted from combustion processes, BC particles can coagulate and grow by condensation, during which both self-coagulation and hetero-coagulation happen (*Jacobson, 2001*). Although BC is mixed internally with other components, the system is impossible to be well-mixed due to the irregular shape of BC (*Jacobson, 2000*). A core-shell morphology is commonly established, with BC as the core and the coating materials (organics, sulfate, etc.) as the shell (*Jacobson, 2001; Zhang et al., 2018*). Numerous efforts have been made to explore the influence of aerosol components on internally mixed BC absorption (*Cappa et al., 2012; Chen et al., 2021; Bond*

70 *et al.*, 2006; Fuller *et al.*, 1999; Jacobson, 2001; Liu *et al.*, 2017; Peng *et al.*, 2016). It was proposed that the coating
components (shell) could act as a lens to focus more photons onto the core to enhance the light absorption of BC (Fuller *et al.*,
75 *et al.*, 1999). Bond *et al.* (2006) estimated that this lensing effect would increase the light absorption of BC by 50-100%.
Jacobson (2001) reported a global average BC absorption enhancement factor of 2, whereas other values, from negligible
(Cappa *et al.*, 2012) to as high as 2.4 (Peng *et al.*, 2016) were also found previously. This lensing effect has been recognized
also as an important factor affecting radiative forcing of BC (Jacobson, 2001). In the recent decade, efforts were made also
to develop models to predict the dynamic evolution of aerosol mixing states (Ching *et al.*, 2016; Curtis *et al.*, 2017; Matsui
80 *et al.*, 2013; Tian *et al.*, 2014). For example, Matsui *et al.* (2013) developed a 2-D aerosol bin scheme that can resolve BC
mixing state and BC aging processes. However, these approaches are far too computationally expensive for use in regional
3-D models (Barnard *et al.*, 2010).

80 Over the past several years, the State Council of China has issued a comprehensive Air Pollution Prevention and
Control Action Plan (APPCAP), covering major emission sectors (Liang *et al.*, 2017; Zhang *et al.*, 2019a). Long-term
observations of aerosol chemical composition indicate that both concentrations of BC and other coating components have
declined rapidly (Gao *et al.*, 2020b; Ji *et al.*, 2019b; Zhou *et al.*, 2019). Although the changes in BC concentrations in
response to emission reduction measures have been documented (Ji *et al.*, 2019b; Gao *et al.*, 2020b), the influence of
85 emission reductions on the aging processes and light absorption of BC has been less explored (Zhang *et al.*, 2019b). Zhang *et al.*
et al. (2018) observed that the declines in absorption of BC was mainly dominated by decreases in BC mass concentration
(86%), and the weakening of BC light absorption capability also played a role (14%). However, this finding was formulated
based on surface observations, little is known about the changes at upper layers. Given the importance of BC absorption in
the upper boundary layer as to buildup of pollution (Ding *et al.*, 2016), the impact of emission reductions on the light
90 absorption of BC, and its implications for the development of boundary layer and pollution episodes need further
investigations. On November 5-11, 2014, Beijing, China hosted the Asia-Pacific Economic Cooperation (APEC) meeting,
during which Beijing and surrounding regions cooperated to implement short-term emission control measures to ensure good
air quality. This event offers a great opportunity to study physical and chemical responses of atmospheric composition to
emission reductions.

95 In this study, we address the following questions using the APEC event as a study case: (1) how did emission
reductions affect the aging processes and light absorption of BC during APEC; (2) what were the relative contributions of
reduced mass concentrations of BC, aging processes of BC, and reshaped mixing state of BC to the changes in light
absorption of BC during APEC; and (3) how did these processes affect BC-PBL interactions and formation of air pollution?
In Sect. 2, we describe the WRF-Chem model configurations and observational datasets used in this study. Results are
100 presented in Sect. 3, and conclusions/discussions are provided in Sect. 4.

2 Methods and data

2.1 WRF-Chem model configuration

WRF-Chem model (*Grell et al., 2005*) version 3.8.1 was adopted in this study to simulate emission, chemical transformation and deposition of aerosols, as well as their interactions with radiation. We demonstrated in previous studies (*Gao et al., 2016a, 2016b, 2020b, 2020c*) that the spatio-temporal variations of air pollutants over China could be reproduced well by WRF-Chem. WRF-Chem enables multiple options for gas phase chemistry and aerosol modules (*Grell et al., 2005*). We employed the Carbon Bond Mechanism version Z (CBMZ) gas phase chemistry (*Zaveri and Peters, 1999*) coupled with the Model for Simulating Aerosol Interactions and Chemistry (MOSAIC) (*Zaveri et al., 2008*) aerosol module in this study. MOSAIC treats size resolved aerosol species, and we used 8 bins version in this study, corresponding to the particle diameter ranges of 0.039-0.078, 0.078-0.156, 0.156-0.312, 0.312-0.625, 0.625-1.25, 1.25-2.5, 2.5-5.0, 5.0-10.0 μm , respectively. Secondary organic aerosol (SOA) formation in MOSAIC was simulated with volatility basis set (VBS) (*Shrivastava et al., 2011*). We configured two nested domains with horizontal resolutions of 81km and 27km, and 31 vertical layers up to a pressure level of 50hPa. The configured domains cover most areas of East Asia and focus on the North China region (same as Figure 1 in *Gao et al., 2017*). Other chosen options for key physical parameterizations follow *Gao et al. (2016b)*. Meteorological initial and boundary conditions were provided by the NCEP $1^\circ \times 1^\circ$ degree final reanalysis dataset (FNL), and chemical initial and boundary conditions were obtained from the MOZART global chemistry simulations (*Emmons et al., 2010*). We used the YSU scheme in WRF-Chem to simulate the evolution of boundary layer, and the YSU scheme defines the top of the boundary layer using Richardson number (*Hong et al., 2010*). To allow the effects of aerosol on meteorological conditions in the model, we did not apply observational nudging or reanalysis nudging.

Anthropogenic emissions of particles and gases in China in the model were taken from the Multi-resolution Emission Inventory for China (MEIC) for year 2014 developed by Tsinghua University (*Zheng et al., 2018*). Anthropogenic emissions for areas outside China were obtained from the MIX Asian emission inventory developed for MICS-Asia and HTAP, which combines five emission inventories for Asia (*Li et al., 2017*). Both MEIC and MIX datasets provide monthly emissions of air pollutants at $0.25^\circ \times 0.25^\circ$ grids, which were interpolated to WRF-Chem modeling domains in this study. We adopted the MEGAN model version 2.04 to estimate biogenic emissions of gases and particles online (*Guenther et al., 2006*). The Global Fire Emissions Database version 4 (GFEDv4) (*Giglio et al., 2013*) were used as open fire emissions.

We simulated the period from October 16 to November 13, and discarded the first seven days as spin-up to avoid the influences of initial conditions. To explore the influences of coordinated emission control measures on BC absorption, we conducted multiple sets of simulations, as described in Table 1. For the NOCTL experiments, simulations were conducted with no perturbations in emissions. For the CTL experiments, emissions of SO_2 , NO_x , PM_{10} , $\text{PM}_{2.5}$, VOCs, and other species in Beijing were reduced by 39.2%, 49.6%, 66.6%, 61.6%, 33.6%, and 50%, respectively, over November 3-11 period. Emissions in Inner Mongolia, Shanxi, Hebei, Tianjin, and Shandong were reduced by 35%. These perturbation

factors were taken from the BMEPB reports (*Gao et al., 2017*). The locations of these provinces are marked in Figure 1 in
 135 *Gao et al. (2017)*.

The influences of BC absorption under different assumptions, including external/core-shell mixing and with/without emission reductions ($\Delta_{BC-Ext-NOCTL}$, $\Delta_{BC-Ext-CTL}$, $\Delta_{BC-CS-NOCTL}$, $\Delta_{BC-CS-CTL}$, and $\Delta_{BC-CYSN-CS}$), can be derived with equations (1-5) below. The description of each simulation is documented in Table 1.

$$\Delta_{BC-Ext-NOCTL} = NOCTL_{Ext} - NOCTL_{Ext-nobc} \quad (1)$$

$$\Delta_{BC-Ext-CTL} = CTL_{Ext} - CTL_{Ext-nobc} \quad (2)$$

$$\Delta_{BC-CS-NOCTL} = NOCTL_{CS} - NOCTL_{CS-nobc} \quad (3)$$

$$\Delta_{BC-CS-CTL} = CTL_{CS} - CTL_{CS-nobc} \quad (4)$$

$$\Delta_{BC-CS-CYSN} = CYSN_{CS} - CYSN_{CS-nobc} \quad (5)$$

The influences of emission reductions during APEC on changes in light absorption of BC and associated changes in
 145 meteorological/pollution conditions under external/core-shell mixing assumptions ($\Delta_{emission-Ext}$ and $\Delta_{emission-CS}$) can be inferred with equations (6-7) below. We use equation (8) to derive the impact of changed BC aging processes by comparing the differences between core-shell simulation and external mixing simulation. The influences of reduced coating due to emission control measures during APEC are calculated with equation (9). We use equation (10) to derive the influences of changes in light absorption enhancement (E_{ab}) of BC.

$$\Delta_{emission-Ext} = \Delta_{BC-Ext-CTL} - \Delta_{BC-Ext-NOCTL} \quad (6)$$

$$\Delta_{emission-CS} = \Delta_{BC-CS-CTL} - \Delta_{BC-CS-NOCTL} \quad (7)$$

$$\Delta_{aging} = \Delta_{BC-CS-CTL} - \Delta_{BC-CYSN-CS} \quad (8)$$

$$\Delta_{coating} = \Delta_{BC-CS-CYSN} - \Delta_{BC-CS-CTL} \quad (9)$$

$$\Delta_{E_{ab}} = \frac{\frac{\Delta_{emission-CS}}{\Delta_{BC-CS-NOCTL}} \frac{\Delta_{emission-Ext}}{\Delta_{BC-Ext-NOCTL}}}{\frac{\Delta_{emission-CS}}{\Delta_{BC-CS-NOCTL}}} \quad (10)$$

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2.2 Calculation of aerosol optical properties in WRF-Chem

WRF-Chem uses Mie theory to calculate layer aerosol optical depth (AOD), single scattering albedo (SSA), and asymmetry factor (g). First, the size parameter and spectral refractive index are used to calculate the Mie extinction efficiency Q_e . Then, the extinction coefficient $\sigma_e(\lambda)$ is provided by the integral of Q_e with consideration of the geometric
 160 size of the particle (πr^2) and the particle number size distribution $n(r)$ (equation (11)). $\sigma_e(\lambda)$ is a equation of wavelength λ . Similarly, absorption coefficient $\sigma_a(\lambda)$ and scattering coefficient $\sigma_s(\lambda)$ can be obtained with Mie absorption efficiency Q_a and Mie scattering efficiency Q_s . The value of SSA can be calculated with equation (12) using $\sigma_a(\lambda)$ and $\sigma_s(\lambda)$.

$$\sigma_e(\lambda) = \int_{r_{min}}^{r_{max}} Q_e \pi r^2 n(r) dr \quad (11)$$

$$SSA(\lambda) = \frac{\sigma_s(\lambda)}{\sigma_s(\lambda) + \sigma_a(\lambda)} \quad (12)$$

165 The calculated optical properties vary with the assumption of mixing state of aerosols. For external mixing, each particle is assumed to be a single chemical species. There are several models proposed for internal mixing, and the commonly used ones include the volume averaging model and core-shell model. In the volume averaging model, all species are assumed to be well mixed, while the core-shell model assumes that BC is coated by a well-mixed shell of other species (Jacobson, 2001). The volume-weighted refractive index m is obtained with the equation below:

$$170 \quad m = \frac{\sum_i V_i m_i}{\sum_i V_i} \quad (13)$$

In equation (13), V_i denotes the volume of species i and m_i represents the refractive index of species i . For core-shell internal mixing, similar averaging processes are applied to the core and shell separately. The scattering efficiency, absorption efficiency and asymmetry parameter are then obtained using the core-shell Mie theory documented in Ackerman and Toon (1981). Core-shell Mie calculation requests core radius, shell radius, refractive index of core and refractive index to shell as inputs (Ackerman and Toon, 1981; Toon and Ackerman, 1981). The official version of WRF-Chem does not calculate optical properties of aerosols with external mixing assumption. To assess the influence of mixing with coating particles on the light absorption of BC, estimated light absorption of pure BC is required. We modified the optical calculation module in WRF-Chem so that it does not mix BC with other chemical species in the calculation of optical properties. In the calculation of optical properties with internal mixing assumption, the volume-weighted refractive index will be inputs of the Mie code. In the calculation of optical properties of BC with external mixing assumption, we only allowed BC to go through the Mie code. The mass and particle number of BC, denoted in M_i and N_i in each bin i (1 through 8) are computed first, and the volume (V_i) are obtained then by dividing by the density of BC. During the calculation of physical diameter (equation 14), other chemical species are not considered. Accordingly, BC does not mix with other chemical species, and we named it as optical properties of BC with external mixing assumption here. It should be noted that this calculation is different from treatments of external mixing in other models that all particles are separated from each other. In our calculation, BC within each bin are internally mixed, although it does not mix with other chemical species. As the purpose of this study is to explore how coating particles on BC would affect the absorption of BC, this treatment would not be an issue.

$$185 \quad D_i = 2 \left(\frac{V_i}{\frac{4}{3}\pi} \right)^{\frac{1}{3}} \quad (14)$$

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Both observations of meteorological variables and air pollutants were used to evaluate the performance of model over the APEC study period in *Gao et al. (2017)* and in this study. The meteorological measurements were retrieved from the National Centers for Environmental Information website (<https://gis.ncdc.noaa.gov/maps/ncei#app=cdo>), which includes near surface temperature, relative humidity (RH), wind speed, and wind direction. The hourly surface concentrations of PM_{2.5} and daily PM_{2.5} chemical compositions were measured at the Institute of Atmospheric Physics (IAP), Chinese Academy of Sciences (CAS) site (*Liu et al., 2017; Yang et al., 2020*). We obtained also AAOD (absorption aerosol optical depth) from the AERONET network (*Dubovik and King, 2000; Holben et al., 1998*) to evaluate model performance. We derived AAOD at 440nm based on Angstrom exponent relation (*Schuster et al., 2006*) to make it consistent with observations. Data from more than 500 sites across the world are provided online at the AERONET website (<http://aeronet.gsfc.nasa.gov>).

3 Results

3.1 Model Evaluation

Model evaluation was conducted with surface observations of meteorological variables, PM_{2.5}, PM_{2.5} chemical components, and AAOD. Data at two meteorological sites in urban Beijing were averaged, and were compared against the model values for the domain grid cell containing the monitoring site. Figure 1 indicates that the daily mean temperature and relative humidity (RH) are captured well by the model. Observed strong wind conditions are slightly underestimated, which is a common issue due to inaccurate land use inputs or other problems in the model (*Gao et al., 2018a*). Our previous investigation (*Gao et al., 2017*) suggested that temperature and RH were lower and northerly winds became more frequent from before APEC to during APEC periods, contributing to pleasant air quality. Figure 2(a) displays the simulated and observed hourly PM_{2.5} concentrations in urban Beijing. Before APEC, observed high PM_{2.5} concentration is well captured by our model. During APEC, the NOCTL case overestimates PM_{2.5} concentrations, while the CTL case exhibits better agreement with observations. Implementing emission reductions in the model lowers the mean bias of the model from 30.8 to -4.0 $\mu\text{g m}^{-3}$. The performance of WRF-Chem in simulating wintertime PM_{2.5} chemical compositions was explored extensively in our previous investigations (*Gao et al., 2016b, 2018a*). Similarly, measured high concentrations of inorganic aerosols (sulfate, nitrate and ammonium) are underestimated, which could be partly due to missing sulfate formation pathways (*Cheng et al. 2016*). We used the updated version with heterogeneous sulfate formation (*Gao et al., 2016a*) to reduce the underestimation of sulfate in this study. Simulated BC concentration shows high degree of consistency with observations, while OC is slightly underestimated due to large uncertainties in current status of SOA modeling (Figure 2f). In general, the temporal variations and magnitudes of air pollutants are well represented in our model. Figure 1(d) compares

simulated AAOD with external mixing assumption and core-shell model against AERONET inferred AAOD during the APEC study period. AAOD simulated with external mixing assumption exhibits much lower values than observation. With the core-shell model, this underestimation is largely reduced. However, AAOD is still underestimated by the model, which might be caused by missing sources of absorbing particles in the model. Currently, the absorption of organics is not treated in the WRF-Chem model, which is likely to underestimate the light-absorbing capability of carbonaceous aerosols in the atmosphere (Andreae and Gelencser, 2006). Uncertainties in the aerosol size distribution in emissions may also contribute to this mismatch between the model and observations (Matsui, 2006).

3.2 Reductions in the concentrations of BC/coating pollutants and changes in BC aging degree

Previously, the reductions of air pollutants were estimated by comparing concentrations of air pollutants during the APEC period with those during other periods. Given the differences in meteorological conditions, such a comparison is not able to indicate the influence of emission control measures. As displayed in Figure 2(a), the concentrations during October 24-25 can be two times of those during October 26-27, although no emission reduction measures were implemented. Previously, we concluded that the meteorological conditions during the APEC week were generally favorable for good air quality compared to it during the week before the APEC week (Gao et al., 2017). Thus, we perturbed emissions in this study to examine how it would affect concentrations of air pollutants, including both BC and its coating pollutants. As displayed in Figure 3(a-b), mean concentrations of SO₂ and NO₂ in urban Beijing declined by 38.7% and 36.3%, respectively, in response to short-term emission control measures. Based on observations, Zhang et al. (2018) reported that SO₂ concentrations decreased by 35% (67%) and NO₂ concentrations decreased by 34% (45%) compared with that before (after) APEC. These declines in aerosol precursors would have modified secondary aerosol formation during the APEC week. Our model indicates that sulfate and nitrate declined by 40.0% and 28.2%, respectively. Given the slight underestimation of sulfate and nitrate, these values might have been moderately underestimated. Mass concentrations of BC declined by 34.6%, while the abundance of OC in the atmosphere exhibited a larger reduction (44.2%).

We used the ratio of the sum of pollutants (primary as well as secondary) to black carbon concentrations (r_{BC}) to track the relative abundance of BC and non-BC particles, which is essential in the C-S calculation:

$$r_{BC} = \frac{[sulfate]+[nitrate]+[ammonium]+[organics]+[dust]+[sodium]+[chloride]}{[BC]} \quad (14)$$

As shown in Figure 3(c), the impacts of emission reductions during APEC on r_{BC} behave differently at different sizes. For ultrafine particles, emission reductions generally lower the aging degree of BC. This is consistent with the observational evidence that smaller BC cores show larger reductions in aging degree as a result of emission control measures during APEC (Zhang et al., 2018). As most secondary aerosols are in smaller sizes, the effect of emission reduction on BC aging is more significant for smaller particles. Zhang et al. (2018) reported only the changes in sizes below 0.2 μm, our modeling results suggest, however, that the aging degree of BC might be enhanced under emission reductions for relatively

larger particles (Figure 3c). The impact of emission reductions on rBC behaves differently near the surface and at higher layers (Figure 3d). The aging degree is lowered in the CTL case near the surface, mainly due to reductions in coating materials. However, at layers higher than 200 meters, the aging degree of BC increases with emission reductions. In-situ near surface measurements indicate also that rBC was reduced during APEC, and the reduction was most likely caused by lower photochemical production (Zhang *et al.*, 2018).

3.3 Changes in AAOD and the light-absorption enhancement (E_{ab}) of BC during APEC

rBC values describe the aging degree of BC, while the exploration of how emission reductions affect light absorption of BC requires a sophisticated calculation of optical properties of BC. Mie theory is commonly used to calculate the light absorption enhancement of BC (E_{ab}) from lensing effect with a core-shell model. Zhang *et al.* (2018) estimated E_{ab} by dividing the light-absorption cross section of the whole BC-containing particle by that of BC core at a certain wavelength. Here we follow the method in Curci *et al.* (2019), and calculate E_{ab} as the ratio of BC AAOD estimated assuming core-shell internal mixing to that calculated with external mixing assumption (enhancement due to mixing with coating non-BC particles):

$$E_{ab} = \frac{BC_AAOD(550nm,core-shell\ mixing)}{BC_AAOD(550nm,external\ mixing)} \quad (\text{AAOD can be either layer or column}) \quad (15)$$

Inferred vertical profiles of layer E_{ab} values in the CTL and NOCTL scenarios are displayed in Figure 3e. At the layers below 5km, mean E_{ab} values are 1.96 and 1.95 for CTL and NOCTL scenarios, respectively. Below 500 m in the troposphere, emission reductions during APEC lower E_{ab} from 2.11 to 2.06. Previous study by Jacobson (2001) suggests a global average BC absorption enhancement factor of 2, which is consistent with current study. However, a wide range of enhancement factors have been reported, from negligible (Cappa *et al.*, 2012) to as high as 2.4 (Peng *et al.*, 2016). Liu *et al.* (2017) pointed that the enhancement factors depend on the particles' mass ratio of non-black carbon to black carbon. Our model results indicate also that the reductions in light absorption enhancement of BC are concentrated at lower layers, while enhancement could happen at higher layers (Figure 3e). This is consistent with the vertical profile of rBC where it decreases due to emission reductions near the surface while increases at higher layers (Figure 3d).

Figure 4a presents the daytime mean (defined as the mean BC AAOD over 10:00-17:00 time period) BC AAOD in Beijing inferred from simulations with different mixing assumptions and emission perturbations. In the NOCTL scenarios, BC AAOD simulated with core-shell model exhibits higher values than those with external mixing assumption (0.0220 for external and 0.0427 for core-shell). Due to reductions in emissions, these values decline to 0.0145 and 0.0283, respectively. Due to emission reductions (differences between CTL and NOCTL scenarios), mean daytime BC AAOD decrease by 0.0075 during the APEC week, as a result of declines in mass concentration of BC (52.0%, Table 2). However, the lensing effect of BC induces a further decline of 0.0069 (48.0%, Table 2). The influence of lensing effect is dominated by the reductions in

coating materials (39.4%, equation (9), Table 2). The BC absorption enhancement (E_{ab}) factor decreased by 0.003 due to
290 reductions in emissions (Figure 4b). We further quantified that the reduced light absorption capability (E_{ab}) resulting from
emission reductions during APEC contributed 3.2% to the total reductions in AAOD (equation (10), Table 2).

3.4 Influences on boundary layer process and air pollution

The vertical distribution of BC absorption plays an important role in modulating the temperature gradient and changing
295 boundary layer meteorology (Ding *et al.*, 2016). We conducted a series of numerical experiments to understand the
influences of reshaped BC absorption due to emission reductions during APEC on boundary layer process and the formation
of air pollution. Figure 4c illustrates the vertical profiles of BC absorption induced changes in equivalent potential
temperature (EPT), which is commonly used to indicate the stability of air in the atmosphere (Obremski *et al.*, 1989). When
EPT decreases with height, the atmosphere is unstable and vertical motion/convection is likely to occur. In all experiments,
300 BC absorption induces a positive impact on EPT in the air above ground acting to enhance the stability of the atmosphere
(Figure 4c). The maximum enhancement occurs at layers close to 1-2km (Figure 4c). At ~2.6km, the maximum ratio of
changes with core-shell model to those with external mixing reach above 2.5, indicating the important effects of mixing state
of BC in the upper boundary layer (Figure 4c).

In urban Beijing, BC absorption induced mean changes of daytime planetary boundary layer height (PBLH) during the
305 APEC week are -11.6 and -24.0 m for external mixing and core-shell model, respectively (Figure 4d). Under a relatively
clean condition (CTL scenarios), these values change to -8.8 and -15.6 m for external mixing and core-shell model for
NOCTL emissions (Figure 4d). Due to emission reductions, the impacts of BC absorption on PBL inhibition decrease by 8.2
m on average during the APEC week (reduced emissions enhance PBLH by 8.2 m). The influences of reduced mass
concentration of BC itself account for 35% of the total changes, while the lensing effect of BC explain the rest (65%, Table
310 2). The decreased coating due to emission reductions dominate the lensing effect of BC (47.4%, Table 2).

The corresponding changes in daytime mean near surface concentrations of O_3 and $PM_{2.5}$ in Beijing are displayed in Figure
4e and Figure 4f, respectively. The inhibited development of PBL due to BC absorption results in higher abundance of $PM_{2.5}$
within the PBL (Ding *et al.*, 2016; Gao *et al.*, 2016b). Previously, we quantified that the co-benefits of reduced aerosol
feedbacks could explain ~11% of the total decreases in $PM_{2.5}$ in Beijing during APEC. Here we focus on light absorption of
315 BC, and find that the lensing effect of BC decreases $PM_{2.5}$ concentration by $0.8 \mu\text{g m}^{-3}$ on average (Figure 4f). On average,
declines in BC mass concentration itself account for 64.3% of the total impact of reduced light absorption of BC on $PM_{2.5}$,
while 35.7% is attributed to the lensing effect of BC. However, inhibited PBL development does not necessarily lead to
enhanced levels of near surface O_3 , as the formation of O_3 is also affected by changes in aerosols and photolysis reactions
above the ground (Chen *et al.*, 2021; Lu *et al.*, 2019). As displayed in Figure 4e, near surface O_3 concentrations in urban
320 Beijing decrease in response to BC absorption.

The spatial distribution of $\Delta_{emission-Ext}$ and $\Delta_{emission-CS}$ in Figure 5 reveal that external mixing and core-shell models estimate similar patterns of changes in AAOD, PBLH, near surface O_3 and near surface $PM_{2.5}$. However, the responses of these variables are larger in the core-shell model due to lensing effects of coating materials. Reduced emissions of BC and its coating materials during APEC led to declined AAOD, less stabilized PBLH, decreased near surface $PM_{2.5}$ concentrations and enhanced near surface O_3 concentrations in the North China Plain (Figure 5). It was noted that $PM_{2.5}$ concentrations was enhanced in northwest China, particularly when we used C-S model. This is related absorption-modulated natural emissions of windblown dust.

Figure 6a,6b illustrates the cross sections in the northeast direction of changes in BC absorption coefficient due to emission reductions, as Beijing and polluted cities in South Hebei are covered. Pronounced declines are concentrated below 2km, and the core-shell model estimates stronger reductions due to lensing effects (Figure 6c). Emission control measures also reshaped the light absorption enhancement factor of BC, as indicated in Figure 6f. Within the lower boundary layer, E_{ab} values were reduced with emission reductions during APEC. Light absorption of BC stabilizes boundary layer to accumulate $PM_{2.5}$, yet this effect is inhibited at lower emission levels during APEC. These relationships are reflected in Figure 6g and 6h with negative changes in $PM_{2.5}$ near the ground.

The responses of O_3 to reduced light absorption of BC during APEC are in the opposite direction (*Gao et al., 2018c*), compared to those for $PM_{2.5}$. Strong absorption of BC tends to enhance photolysis above the aerosol layer, but to reduce photolysis near the ground. Figure 7d, 7g illustrate the changes in O_3 1D and NO_2 photolysis rates with emission reductions inferred from an external mixing assumption. With emission control implemented, photolysis rates near the ground are enhanced due to lower light absorption of BC, while the photolysis rates above the aerosol layer are reduced. Similar patterns but with larger values are found using the core-shell model (Figure 7e, 7h). The responses of O_3 are generally in line with the responses of O_3 1D and NO_2 photolysis rates (Figure 7a, 7b).

4 Summary and Discussions

In this study, we used the online coupled WRF-Chem model to understand how emission control measures during the APEC event would affect the mixing state/light absorption of BC, and the implications for BC-PBL interactions. Multiple observations, including surface observations of meteorological variables, $PM_{2.5}$, $PM_{2.5}$ chemical composition, and AAOD were used to evaluate model performance. A series of numerical experiments were conducted to address three questions: (1) how did emission reductions affect the aging processes and light absorption of BC during APEC; (2) what were the relative contributions of reduced mass concentrations of BC, aging processes of BC, and reshaped mixing state of BC to the changes in light absorption of BC during APEC; and (3) how did these processes affect BC-PBL interactions and formation of air pollution?

We found that both the mass concentration of BC and the BC coating materials declined during the APEC week, which reduced the light absorption and light absorption enhancement (E_{ab}) of BC. Below 500 m in the troposphere, emission reductions during APEC lowered the absorption enhancement factor E_{ab} from 2.11 to 2.06. The column absorption
355 enhancement was reduced also. The reduced AAOD during APEC is caused by both the declines in mass concentration of BC itself (52.0%) and the lensing effect of BC (48.0%). The reductions in coating materials (39.4%) dominated the influence of lensing effect, and the reduced light absorption capability (E_{ab}) contributed 3.2% to the total reductions in AAOD. Our estimate of the contribution of reduced light absorption capability (E_{ab}) exhibit lower values than *Zhang et al. (2016)*, which could be caused by the uncertainties in the assumption of the mixing state of BC in the core-shell model.

360 The diminished light absorption of BC during APEC promotes the development of PBL, as indicated in the changes in vertical profiles of EPT. Different responses of $PM_{2.5}$ and O_3 were found to the changes in light absorption of BC. The responses of $PM_{2.5}$ follow the enhanced PBLH to decrease, while O_3 concentrations increase near the ground. The enhanced levels of O_3 were mainly caused by the influences of BC absorption on photolysis rates. As displayed in the conceptual scheme plot in Figure 8, reduced emissions of BC and its coating materials during APEC led to declined AAOD, less
365 stabilized PBLH, decreased near surface $PM_{2.5}$ concentrations and enhanced near surface O_3 concentrations in the North China Plain.

This study with perturbations of emissions during APEC offer important implications on the potential effects of China's Clean Air Act. As discussed in our previous investigation (*Gao et al., 2017*), emission control measures have the co-
benefits of reducing aerosol feedbacks to accelerate the cleaning of air, which accounts for ~11% of the decreased $PM_{2.5}$
370 concentrations during APEC. In this study, we further clarified that the ongoing measures to control SO_2 , NO_x , etc. would be efficient to reduce the absorption capability of BC to inhibit the feedback of BC on the boundary layer. Our results also show that near ground O_3 responds differently from the changes in $PM_{2.5}$, which might be a side effect of current emission control strategies. *Ma et al. (2021)* reported that aerosol radiative effect could explain 23% of the total change in surface summertime O_3 in China. How to control emissions to offset this side effect of current emission control measures on O_3
375 should be an area of further focus. In addition to the influences on air quality and weather, a sudden reduction in aerosol emissions may potentially affect climate (*Ren et al., 2020; Yang et al., 2020*), which warrants further investigation.

Although careful validation was conducted in this study, uncertainties still remain in the current study. We concluded that the core-shell model captures the variation of AAOD better than external mixing assumption. However, the core-shell model is an ideal scenario that assumes all non-BC materials are internally mixed and coated on BC (*Miyakawa et al., 2014*). *Zhang et al. (2016)* observed that BC particles are heavily coated and are in a near-spherical shape in the North
380 China Plain. The usage of core-shell model seems to be reasonable in this study, whereas the assumption that all non-BC materials are coated on BC might not be true in real atmosphere. The observed ratio of coatings to PM_1 was ~25-70% in summer in Beijing (*Xu et al., 2019*), and the observed ratio of coatings to $PM_{2.5}$ was ~10-40% in winter in Beijing (*Wang et al., 2019*). We conducted additional simulations assuming 10%, 20%, 40%, 60%, and 80% mixing fractions of coating
385 aerosols, and explored how these assumptions would affect estimated BC AAOD, and its feedbacks on radiation, boundary

layer and air pollutants. As suggested in Figure 9, modelled BC AAOD increases gradually when mixing fraction rises from 0.1 to 0.4, but it keeps relatively stable when mixing fraction is between 0.4 and 1. The responses of near surface shortwave radiation and PBLH are in line with it, exhibiting relatively constant reductions when mixing fraction is higher than 0.4 (Figure 9). However, the no significant relationships are found for BC absorption induced changes in O_3 (<10% difference between fractions of 0.1 and 1) and $PM_{2.5}$ (<22% difference between fractions of 0.1 and 1) in Beijing. These results suggest that the findings demonstrated in this study might not be largely affected by the assumptions in mixing fractions of coating particles. Additionally, the simulated feedbacks of BC absorption on boundary layer processes are not well constrained. We used multiple coupled models to examine how these processes are represented, and we calculated ensemble mean to obtain the best current understanding (*Gao et al., 2018a, 2020a*). In the future, further efforts are needed also to constrain the uncertainties of these processes in the model.

Data availability. The measurements and model simulations data can be accessed through contacting the corresponding authors.

Author contributions. MG and JH designed the study. MG performed model simulations and analyzed the data with help from YY, HL, BZ, YZ, XL, CW, QZ and GRC. QZ provided the emission inventory. YW and ZL provided measurements. MG and JH wrote the paper with inputs from all the other authors.

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References

- Ackerman, T. P. and Toon, O. B.: Absorption of visible radiation in atmosphere containing mixtures of absorbing and non-absorbing particles, *Appl. Optics*, 20(20), 3661–3662, 1981.
- Andreae, M. O. and Gelencsér, A.: Black carbon or brown carbon? The nature of light-absorbing carbonaceous aerosols, *Atmos. Chem. Phys.*, 6(10), 3131–3148, doi:10.5194/acp-6-3131-2006, 2006.

- Barnard, J. C., Fast, J. D., Paredes-Miranda, G., Arnott, W. P., and Laskin, A.: Technical Note: Evaluation of the WRF-Chem “Aerosol Chemical to Aerosol Optical Properties” Module using data from the MILAGRO campaign, *Atmos. Chem. Phys.*, 420 10, 7325–7340, doi:10.5194/acp-10-7325-2010, 2010.
- Bond, T. C., Streets, D. G., Yarber, K. F., Nelson, S. M., Woo, J. and Klimont, Z.: A technology-based global inventory of black and organic carbon emissions from combustion, *J. Geophys. Res. Atmos.*, 109(D14203), doi:10.1029/2003JD003697, 2004.
- Bond, T. C., Habib, G. and Bergstrom, R. W.: Limitations in the enhancement of visible light absorption due to mixing state, *J. Geophys. Res.*, 111(D20211), doi:10.1029/2006JD007315, 2006.
- Bond, T. C., Doherty, S. J., Fahey, D. W., Forster, P. M., Berntsen, T., Deangelo, B. J., Flanner, M. G., Ghan, S., Kärcher, B., Koch, D., Kinne, S., Kondo, Y. and Quinn, P. K.: Bounding the role of black carbon in the climate system: A scientific assessment, *J. Geophys. Res. Atmos.*, 118(11), 5380–5552, doi:10.1002/jgrd.50171, 2013.
- Chen, D., Liao, H., Yang, Y., Chen, L. and Wang, H.: Simulated aging processes of black carbon and its impact during a severe winter haze event in the Beijing-Tianjin-Hebei region, *Sci. Tot. Env.*, 755, p.142712, 2021.
- Chen, Y., Beig, G., Archer-Nicholls, S., Drysdale, W., Acton, J., Lowe, D., Nelson, B. S., Lee, J. D., Ran, L., Wang, Y., Wu, Z., Sahu, S. K., Sokhi, R. S., Singh, V., Gadi, R., Hewitt, C. N., Nemitz, E., Archibald, A., McFiggins, G., and Wild, O.: Avoiding high ozone pollution in Delhi, India, *Faraday Discussions*, 10.1039/D0FD00079E, 2021.
- Ching, J., Riemer, N., and West, M.: Impacts of black carbon particles mixing state on cloud microphysical properties: sensitivity to environmental conditions, *J. Geophys. Res.-Atmos.*, 121, 5990–6013, <https://doi.org/10.1002/2016JD024851>, 2016.
- Curtis, G., Alyuz, U., Barò, R., Bianconi, R., Bieser, J., Christensen, J. H., Colette, A., Farrow, A., Francis, X., Jiménez-guerrero, P., Im, U. and Liu, P.: Modelling black carbon absorption of solar radiation: combining external and internal mixing assumptions, *Atmos. Chem. Phys.*, 19(1), 181–204, doi:10.5194/acp-19-181-2019, 2019.
- Curtis, J. H., Riemer, N., and West, M.: A single-column particle-resolved model for simulating the vertical distribution of aerosol mixing state: WRF-PartMC-MOSAIC-SCM v1.0, *Geosci. Model Dev.*, 10, 4057–4079, <https://doi.org/10.5194/gmd-10-4057-2017>, 204470
- Ding, A. J., Huang, X., Nie, W., Sun, J. N., Kerminen, V., Petäjä, T., Su, H., Cheng, Y. F., Yang, X., Wang, M. H., Chi, X. G., Wang, J. P., Virkkula, A., Guo, W. D., Yuan, J., Wang, S. Y., Zhang, R. J., Wu, Y. F., Song, Y., Zhu, T., Zilitinkevich, S., Kulmala, M. and Fu, C. B.: Enhanced haze pollution by black carbon in megacities in China, *Geophys. Res. Lett.*, 43(6), 2873–2879, 445 doi:10.1002/2016GL067745, 2016.
- Dubovik, O. and King, M. D.: A flexible inversion algorithm for retrieval of aerosol optical properties from Sun and sky radiance measurements A flexible inversion algorithm for retrieval of aerosol optical properties from Sun and sky radiance measurements, *J. Geophys. Res. Atmos.*, 105(D16), 20673–20696, doi:10.1029/2000JD900282, 2000.
- Emmons, L. K., Walters1, S., Hess1, P. G., Lamarque, J.-F., Pfister1, G. G., Fillmore1, D., Granier, C., Guenther, A., Kinnison, D., 450 Laepple1, T., Orlando, J., Tie, X., Tyndall, G., Wiedinmyer, C., Baughcum, S. L. and Kloster, S.: Description and evaluation

- of the Model for Ozone and Related chemical Tracers, version 4 (MOZART-4), *Geosci. Model Dev.*, 3(1), 43–67, doi:10.5194/gmd-3-43-2010, 2010.
- Flanner, M. G., Zender, C. S., Randerson, J. T. and Rasch, P. J.: Present-day climate forcing and response from black carbon in snow
Present-day climate forcing and response from black carbon in snow, *J. Geophys. Res. Atmos.*, 112(D11202), 455 doi:10.1029/2006JD008003, 2007.
- Fuller, K. A., Malm, W. C. and Kreidenweis, S. M.: Effects of mixing on extinction by carbonaceous particles Effects of mixing on extinction by carbonaceous particles, *J. Geophys. Res. Atmos.*, 104(D13), 15941–15954, doi:10.1029/1998JD100069, 1999.
- Gao, M., Carmichael, G. R., Wang, Y., Ji, D., Liu, Z. and Wang, Z.: Improving simulations of sulfate aerosols during winter haze over Northern China: the impacts of heterogeneous oxidation by NO₂, *Front. Environ. Sci. Technol.*, 10(5), 1–11, 460 doi:10.1007/s11783-016-0878-2, 2016a.
- Gao, M., Carmichael, G. R., Wang, Y., Saide, P. E., M. Yu, J. Xin, Z. Liu and Z. Wang: Modeling study of the 2010 regional haze event in the North China Plain, *Atmos. Chem. Phys.*, 16(3), 1673–1691, doi:10.5194/acp-16-1673-2016, 2016b.
- Gao, M., Liu, Z., Wang, Y., Lu, X., Ji, D. and Wang, L.: Distinguishing the roles of meteorology, emission control measures, regional transport, and co-benefits of reduced aerosol feedbacks in “APEC” Blue, *Atmos. Environ.*, 167, 476–486, 465 doi:10.1016/j.atmosenv.2017.08.054, 2017.
- Gao, M., Han, Z., Liu, Z., Li, M., Xin, J., Tao, Z. and Li, J.: Air Quality and Climate Change , Topic 3 of the Model Inter-Comparison Study for Asia Phase III (MICS-Asia III), Part I: overview and model evaluation, *Atmos. Chem. Phys.*, 18(7), 4859–4884, doi:10.5194/acp-18-4859-2018, 2018a.
- Gao, M., Ji, D., Liang, F. and Liu, Y.: Attribution of aerosol direct radiative forcing in China and India to emitting sectors, *Atmos. Environ.*, 190, 35–42, doi:10.1016/j.atmosenv.2018.07.011, 2018b.
- Gao, J., Zhu, B., Xiao, H., Kang, H., Pan, C., Wang, D., and Wang, H.: Effects of black carbon and boundary layer interaction on surface ozone in Nanjing, China, *Atmos. Chem. Phys.*, 18, 7081–7094, <https://doi.org/10.5194/acp-18-7081-2018>, 2018c.
- Gao, M., Han, Z., Tao, Z., Li, J., Kang, J., Huang, K. and Dong, X.: Air quality and climate change , Topic 3 of the Model Inter-Comparison Study for Asia Phase III (MICS-Asia III)–Part 2: aerosol radiative effects and aerosol feedbacks, *Atmos. Chem. Phys.*, 20(2), 1147–1161, doi:10.5194/acp-20-1147-2020, 2020a.
- Gao, M., Liu, Z., Zheng, B., Ji, D., Sherman, P., Song, S., Xin, J., Liu, C., Wang, Y., Zhang, Q., Xing, J., Jiang, J., Wang, Z. and Carmichael, G. R.: China’s emission control strategies have suppressed unfavorable influences of climate on wintertime PM 2.5 concentrations in Beijing since 2002, *Atmos. Chem. Phys.*, 20(3), 1497–1505, doi:10.5194/acp-20-1497-2020, 2020b.
- Gao, M., Gao, J., Zhu, B., Kumar, R., Lu, X., Song, S., Zhang, Y. and Jia, B.: Ozone pollution over China and India : seasonality and 480 sources, *Atmos. Chem. Phys.*, 20(7), 4399–4414, doi:10.5194/acp-20-4399-2020, 2020c.
- Giglio, L., Randerson, J. T. and Werf, G. R. Van Der: Analysis of daily , monthly , and annual burned area using the fourth-generation global fire emissions database (GFED4), *J. Geophys. Res. Biogeosciences*, 118(1), 317–328, doi:10.1002/jgrg.20042, 2013.

- Grell, G. A., Peckham, S. E., Schmitz, R., Mckeen, S. A., Frost, G., Skamarock, W. C. and Eder, B.: Fully coupled “online” chemistry within the WRF model, *Atmos. Environ.*, 39(37), 6957–6975, doi:10.1016/j.atmosenv.2005.04.027, 2005.
- Grieshop, A. P., Reynolds, C. C. O., Kandlikar, M. and Dowlatabadi, H.: A black-carbon mitigation wedge, *Nat. Publ. Gr.*, 2(8), 533–534, doi:10.1038/ngeo595, 2009.
- Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P. I. and Geron, C.: Estimates of global terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from Nature), *Atmos. Chem. Phys.*, 6(11), 3181–3210, 490 doi:10.5194/acp-6-3181-2006, 2006.
- Holben, B. N., Slutsker, T. I. E. I., Tar, D., Buis, J. P., Setxerj, I. I. A., Reagan, A., J, Y., Nakajima, T., Lavenu, F., Vemte, E., Jankowiak, I. and Smirnozt, A.: AERONET-A Federated Instrument Network and Data Archive for Aerosol Characterization, *Remote Sens. Environ.*, 66(1), 1–16, 1998.
- IPCC: Climate Change 2014: Synthesis Report, Contribution of Working Groups I, II and III to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, edited by: Core Writing Team, R.K. Pachauri and L.A. Meyer, 151pp., IPCC, Geneva, Switzerland, 2014.
- Hong, S.-Y.: A new stable boundary-layer mixing scheme and its impact on the simulated East Asia summer monsoon, *Q. J. Roy. Meteorol. Soc.*, 136, 1481–1496, 2010
- Jacobson, M. Z.: Strong radiative heating due to the mixing state of black carbon in atmospheric aerosols, *Nature*, 409(6821), 695–500 697, doi:10.1038/35055518, 2001.
- Ji, D., He, J., Sun, Y. and Gao, M.: Two-year continuous measurements of carbonaceous aerosols in urban Beijing, China: Temporal variations, characteristics and source analyses, *Chemosphere*, 200, 191–200, doi:10.1016/j.chemosphere.2018.02.067, 2018.
- Ji, D., Gao, W., Maenhaut, W., He, J., Wang, Z. and Li, J.: Impact of air pollution control measures and regional transport on carbonaceous aerosols in fine particulate matter in urban Beijing, China: Insights gained from long-term measurement, *Atmos. Chem. Phys.*, 19(13), 8569–8590, doi:10.5194/acp-2019-42, 2019a.
- Ji, D., Gao, M., Maenhaut, W., He, J., Wu, C., Cheng, L., Gao, W., Sun, Y., Sun, J., Xin, J., Wang, L. and Wang, Y.: The carbonaceous aerosol levels still remain a challenge in the Beijing-Tianjin-Hebei region of China: Insights from continuous high temporal resolution measurements in multiple cities, *Environ. Int.*, 126, 171–183, doi:10.1016/j.envint.2019.02.034, 2019b.
- Li, L., Liao, H., Mao, Y. and Ridley, D. A.: Source sector and region contributions to concentration and direct radiative forcing of black carbon in China, *Atmos. Environ.*, 124, 351–366, doi:10.1016/j.atmosenv.2015.06.014, 2016.
- Li, M., Zhang, Q., Kurokawa, J., Woo, J., He, K., Lu, Z. and Ohara, T.: MIX: a mosaic Asian anthropogenic emission inventory under the international collaboration framework of the MICS-Asia and HTAP, *Atmos. Chem. Phys.*, 17(2), 935–963, doi:10.5194/acp-17-935-2017, 2017.
- Li, F., Gao, M., Xiao, Q., Carmichael, G.R., Pan, X. and Liu, Y.: Evaluation of a data fusion approach to estimate daily PM_{2.5} levels in North China. *Environmental research*, 158, pp.54-60, 2017.

- Liu, D., Whitehead, J., Alfarra, M. R., Reyes-villegas, E., Spracklen, D. V., Reddington, C. L., Kong, S., Williams, P. I., Ting, Y., Haslett, S., Taylor, J. W., Flynn, M. J., Morgan, W. T., Mcfiggans, G., Coe, H. and Allan, J. D.: Black-carbon absorption enhancement in the atmosphere determined by particle mixing state, *Nat. Geosci.*, 10(3), 184–188, doi:10.1038/NGEO2901, 520 2017.
- Liu, Z., Hu, B., Zhang, J., Xin, J., Wu, F., Gao, W., Wang, M. and Wang, Y.: Characterization of fine particles during the 2014 Asia-Pacific economic cooperation summit: Number concentration, size distribution and sources, *Tellus B: Chem. Phys. Met.*, 69(1), p.1303228, 2017.
- Lu, X., Zhang, L., Zhao, Y., Jacob, D. J., Hu, Y., Hu, L., Gao, M., Liu, X., Petropavlovskikh, I., McClure-Begley, A., and Querel, R.: 525 Surface and tropospheric ozone trends in the Southern Hemisphere since 1990: possible linkages to poleward expansion of the Hadley circulation, *Sci. Bull.*, 64, 400–409, <https://doi.org/10.1016/j.scib.2018.12.021>, 2019Ma, X., Huang, J., Zhao, T., Liu, C., Zhao, K., Xing, J., and Xiao, W.: Rapid increase in summer surface ozone over the North China Plain during 2013–2019: a side effect of particulate matter reduction control?, *Atmos. Chem. Phys.*, 21, 1–16, <https://doi.org/10.5194/acp-21-1-2021>, 2021.
- Matsui, H., Black carbon simulations using a size-and mixing-state-resolved three-dimensional model: 1. Radiative effects and their 530 uncertainties, *J. Geophys. Res. Atmos.*, 121, 1793–1807, doi:10.1002/2015JD023998, 2016.
- Matsui, H., Koike, M., Kondo, Y., Moteki, N., Fast, J. D., and Zaveri, R. A.: Development and validation of a black carbon mixing state resolved three-dimensional model: Aging processes and radiative impact, *J. Geophys. Res.-Atmos.*, 118, 2304–2326, doi:10.1029/2012JD018446, 2013.
- 535
- Menon, S., Hansen, J., Nazarenko, L. and Luo, Y.: Climate Effects of Black Carbon Aerosols in China and India, *Science*, 297(5590), 2250–2253, doi:10.1126/science.1075159, 2002.
- Miyakawa, T., Takeda, N., Koizumi, K., Tabaru, M., Ozawa, Y., Hirayama, N., and Takegawa, N.: A new laser induced incandescence – mass spectrometric analyzer (LII-MS) for online measurement of aerosol composition classified by black 540 carbon mixing state. *Aerosol Sci. Tech.*, 48, 853-863, 2014.
- OBREMSKI, J., SAMSON, J., DUTKIEWICZ, V. and HUSAIN, L.: On the use of surface equivalent potential temperature in isolating the influence of local as opposed to transported sources on aerosol concentrations, *J. Geophys. Res. Atmos.*, 94(D8), 11117–11127, doi:10.1029/JD094iD08p11117, 1989.
- Onasch, C. D. C. T. B., Massoli, P., Worsnop, D. R., Bates, T. S., Cross, E. S., Davidovits, P., Hakala, J., Hayden, K. L., Jobson, B. 545 T., Kathryn R. Kolesar, D. A. L., Lerner, B. M., Li, S.-M., Mellon, D., Nuaaman, I., Olfert, J. S., Petäjä, T., Quinn, P. K., Song, C., Subramanian, R., Williams, E. J. and Zaveri, R. A.: Radiative Absorption Enhancements Due to the Mixing State of Atmospheric Black Carbon, *Science*, 337(6098), 1078–1081, doi:10.1126/science.1223447, 2012.
- Pan, Y., Tian, S., Liu, D., Fang, Y., Zhu, X., Gao, M., Wentworth, G.R., Michalski, G., Huang, X. and Wang, Y.: Source Apportionment of Aerosol Ammonium in an Ammonia-Rich Atmosphere: An Isotopic Study of Summer Clean and Hazy 550 Days in Urban Beijing. *Journal of Geophysical Research: Atmospheres*, 123(10), pp.5681-5689, 2018.

- Peng, J., Hu, M., Guo, S., Du, Z., Zheng, J., Shang, D., Levy, M. and Zeng, L.: Markedly enhanced absorption and direct radiative forcing of black carbon under polluted urban environments, *Proc. Natl. Acad. Sci.*, 113(16), 4266–4271, doi:10.1073/pnas.1602310113, 2016.
- Qin, W., Zhang, Y., Chen, J., Yu, Q., Cheng, S., Li, W., Liu, X. and Tian, H.: Variation, sources and historical trend of black carbon 555 in Beijing, China based on ground observation and MERRA-2 reanalysis data, *Environ. Pollut.*, 245, 853–863, doi:10.1016/j.envpol.2018.11.063, 2019.
- Ramanathan, V. and Carmichael, G. R.: Global and regional climate changes due to black carbon, *Nat. Geosci.*, 1(4), 221–227, doi:10.1038/ngeo156, 2008.
- Ren, L., Yang, Y., Wang, H., Zhang, R., Wang, P., and Liao, H.: Source attribution of Arctic black carbon and sulfate aerosols and 560 associated Arctic surface warming during 1980–2018, *Atmos. Chem. Phys.*, 20, 9067–9085, <https://doi.org/10.5194/acp-20-9067-2020>, 2020.
- Schuster, G. L., Dubovik, O., and Holben, B. N.: Angstrom exponent and bimodal aerosol size distributions, *J. Geophys. Res.*, 111, D07207, doi:10.1029/2005JD006328, 2006.
- Shrivastava, M., Easter, R. C., Northwest, P., Zaveri, R. A., Northwest, P., Saide, P. E. and Angeles, L.: Modeling organic aerosols in 565 a megacity : comparison of simple and complex representations of the volatility basis set approach, *Atmos. Chem. Phys.*, 11(13), 6639–6662, doi:10.5194/acp-11-6639-2011, 2011.
- Tian, J., Riemer, N., West, M., Pfaffenberger, L., Schlager, H., and Petzold, A.: Modeling the evolution of aerosol particles in a ship plume using PartMC-MOSAIC, *Atmos. Chem. Phys.*, 14, 5327–5347, <https://doi.org/10.5194/acp-14-5327-2014>, 2014.
- Toon, O. B. and Ackerman, T.P.: Algorithms for the calculation of scattering by stratified spheres, *Appl. Opt.*, 20, 3657–3660, 1981.
- Wang, J., Liu, D., Ge, X., Wu, Y., Shen, F., Chen, M., Zhao, J., Xie, C., Wang, Q., Xu, W. and Zhang, J., 2019. Characterization of 570 black carbon-containing fine particles in Beijing during wintertime. *Atmospheric Chemistry and Physics*, 19(1), pp.447–458.
- Xu, W., Xie, C., Karnezi, E., Zhang, Q., Wang, J., Pandis, S. N., Ge, X., Zhang, J., An, J., Wang, Q., Zhao, J., Du, W., Qiu, Y., Zhou, W., He, Y., Li, Y., Li, J., Fu, P., Wang, Z., Worsnop, D. R., and Sun, Y.: Summertime aerosol volatility measurements in Beijing, China, *Atmos. Chem. Phys.*, 19, 10205–10216, <https://doi.org/10.5194/acp-19-10205-2019>, 2019.
- Yan, Y., and Liu, Z.: Cleaning the air , protecting the climate : Policy , legal and institutional nexus to reduce black carbon 575 emissions in China, *Environ. Sci. Policy*, 95, 1–10, doi:10.1016/j.envsci.2019.01.016, 2019.
- Yang, Y., Wang, H., Smith, S. J., Ma, P.-L., and Rasch, P. J.: Source attribution of black carbon and its direct radiative forcing in China, *Atmos. Chem. Phys.*, 17, 4319–4336, <https://doi.org/10.5194/acp-17-4319-2017>, 2017.
- Yang, Y., Smith, S. J., Wang, H., Mills, C. M., and Rasch, P. J.: Variability, timescales, and nonlinearity in climate responses to black 580 carbon emissions, *Atmos. Chem. Phys.*, 19, 2405–2420, <https://doi.org/10.5194/acp-19-2405-2019>, 2019.
- Yang, Y., Ren, L., Li, H., Wang, H., Wang, P., Chen, L., Yue, X., and Hong, L.: Fast climate responses to aerosol emission reductions during the COVID-19 pandemic, *Geophys. Res. Lett.*, 47, e2020GL089788, <https://doi.org/10.1029/2020GL089788>, 2020.

- Zhang, Q., Zheng, Y., Tong, D., Shao, M., Wang, S., Zhang, Y. and Xu, X.: Drivers of improved PM 2.5 air quality in China from 585 2013 to 2017, Proc. Natl. Acad. Sci., 116(49), 24463–24469, doi:10.1073/pnas.1907956116, 2019a.
- Zhang, Y., Zhang, Q., Cheng, Y., Su, H., Kecorius, S., Wang, Z., Wu, Z., Hu, M., Zhu, T., Wiedensohler, A., and He, K.: Measuring the morphology and density of internally mixed black carbon with SP2 and VTDMA: new insight into the absorption enhancement of black carbon in the atmosphere, Atmos. Meas. Tech., 9,1833–1843, https://doi.org/10.5194/amt-9-1833-2016, 2016.
- Zhang, Y., Li, X., Li, M., Zheng, Y., Geng, G., Hong, C. and Li, H.: Reduction in black carbon light absorption due to multi-pollutant 590 emission control during APEC China 2014, Atmos. Chem. Phys., 18(14), 10275–10287, doi:10.5194/acp-2018-274, 2018.
- Zhang, Y., Li, M., Cheng, Y., Geng, G., Hong, C., Li, H., Li, X. and Tong, D.: Modeling the aging process of black carbon during atmospheric transport using a new approach: a case study in Beijing, Atmos. Chem. Phys., 19(14), 9663–9680, doi:10.5194/acp-19-9663-2019, 2019b.
- Zhang, B., Tong, D., Li, M., Liu, F., Hong, C., Geng, G., Li, H., Li, X. and Peng, L.: Trends in China’s anthropogenic emissions since 600 2010 as the consequence of clean air actions, Atmos. Chem. Phys., 18(19), 14095–14111, doi:10.5194/acp-18-14095-2018, 2018.
- Zhou, W., Gao, M., He, Y., Wang, Q., Xie, C., Xu, W., Zhao, J., Du, W., Qiu, Y., Lei, L. and Fu, P.: Response of aerosol chemistry to clean air action in Beijing, China: Insights from two-year ACSM measurements and model simulations, Environ. Pollut., 600 255, 113345pp., doi:10.1016/j.envpol.2019.113345, 2019.
- Zaveri, R. A. and K.Peters, L.: A new lumped structure photochemical mechanism for long-scale applications, J. Geophys. Res. Atmos., 104(D23), 30387–30415, doi:10.1029/1999JD900876, 1999.
- Zaveri, R. A., Richard C, E., D.Fast, J. and K.Peters, L.: Model for Simulating Aerosol Interactions and Chemistry (MOSAIC), J. Geophys. Res., 113(D13204), doi:10.1029/2007JD008782, 2008.

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Table 1: Descriptions of model simulations

Experiments	Descriptions
<i>NOCTL_{Ext}</i>	No perturbations in emissions; assuming external mixing of BC.
<i>NOCTL_{Ext-nobc}</i>	No perturbations in emissions; assuming external mixing of BC; assuming no absorption of BC.
<i>NOCTL_{CS}</i>	No perturbations in emissions; calculating optical properties using core-shell assumption.
<i>NOCTL_{CS-nobc}</i>	No perturbations in emissions; calculation of optical properties using core-shell assumption; assuming no absorption of BC.

CTL_{Ext}	emissions are reduced during APEC; assuming external mixing of BC.
$CTL_{Ext-nobc}$	emissions are reduced during APEC; assuming external mixing of BC; assuming no absorption of BC.
CTL_{CS}	emissions are reduced during APEC; calculation of optical properties using core-shell assumption.
$CTL_{CS-nobc}$	emissions are reduced during APEC; calculation of optical properties using core-shell assumption; assuming no absorption of BC.
$CYSN_{CS}$	emissions of BC are reduced while emissions of other species are not during APEC; calculation of optical properties using core-shell assumption.
$CYSN_{CS-nobc}$	emissions of BC are reduced while emissions of other species are not during APEC; calculation of optical properties using core-shell assumption; assuming no absorption of BC.

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Table 2: The division of the impact of BC absorption into the impact of BC mass itself and BC mixing state

Influenced variables	BC mass itself	BC lensing effect	Reduced coating	Reduced E_{ab}
AAOD	52.0%	48.0%	39.4%	3.2%
PBLH	34.9%	65.1%	47.4%	-
PM _{2.5}	64.3%	35.7%	-	-
O ₃	49.1%	50.9%	-	-

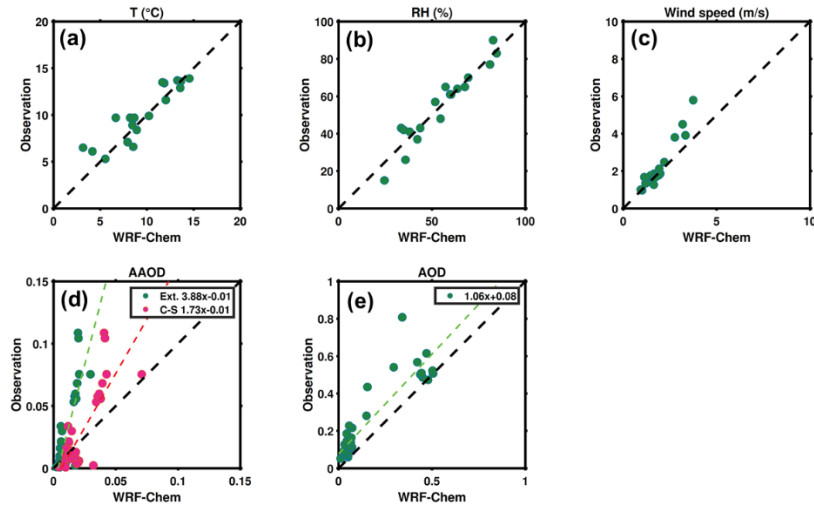


Figure 1: Scatter plots of modeled and observed near surface meteorological variables (a: T, b: RH, c: wind speed); modelled AOD with core-shell model/external mixing assumption, and the comparisons against observations (d); modelled and observed AOD (e).

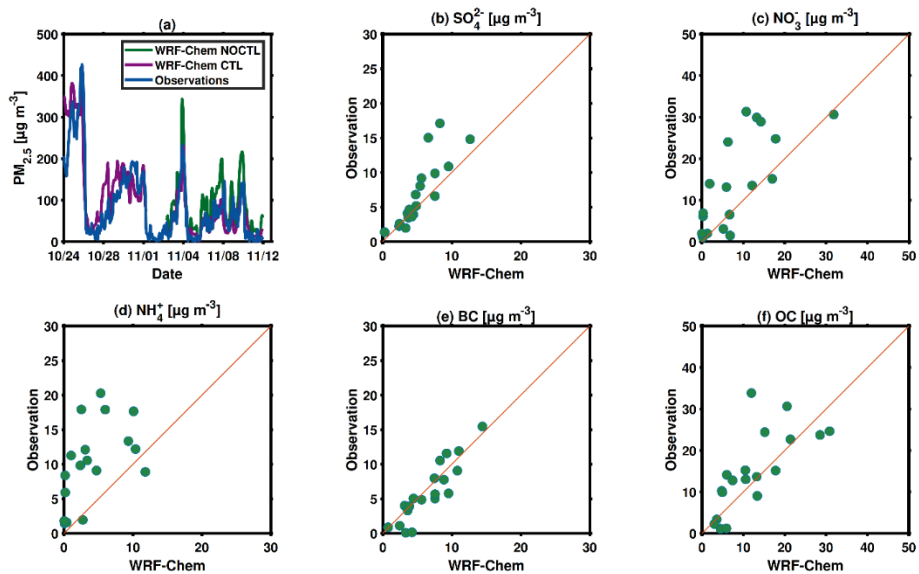


Figure 2: Modeled and observed time series of $PM_{2.5}$ concentrations in urban Beijing (a); Scatter plots of modeled and observed near surface daily mean concentrations of sulfate, nitrate, ammonium, BC (black carbon) and OC (organics) in Beijing (b-f).

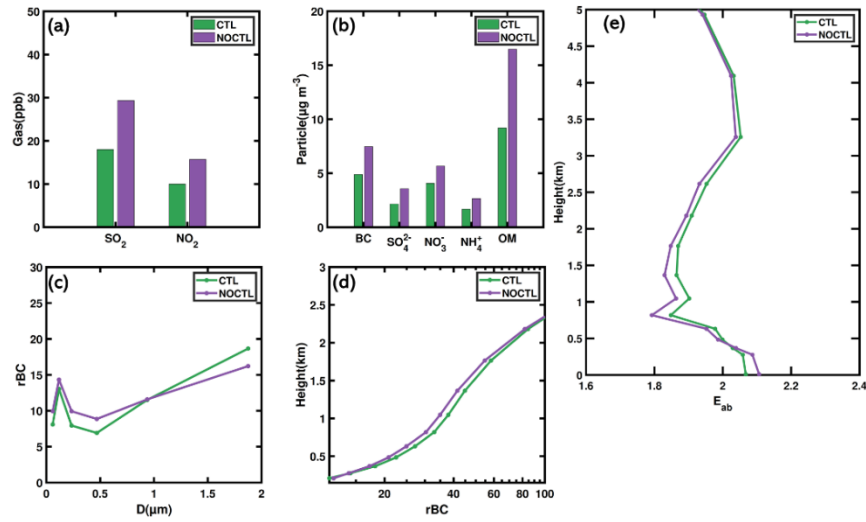


Figure 3: Mass concentrations of gaseous and condensed pollutants in the CTL and NOCTL cases (a-b); the distribution of rBC with sizes in the CTL and NOCTL cases (c); the distribution of rBC with height in the CTL and NOCTL cases (d) and the distribution of E_{ab} values with height in the CTL and NOCTL cases (e).

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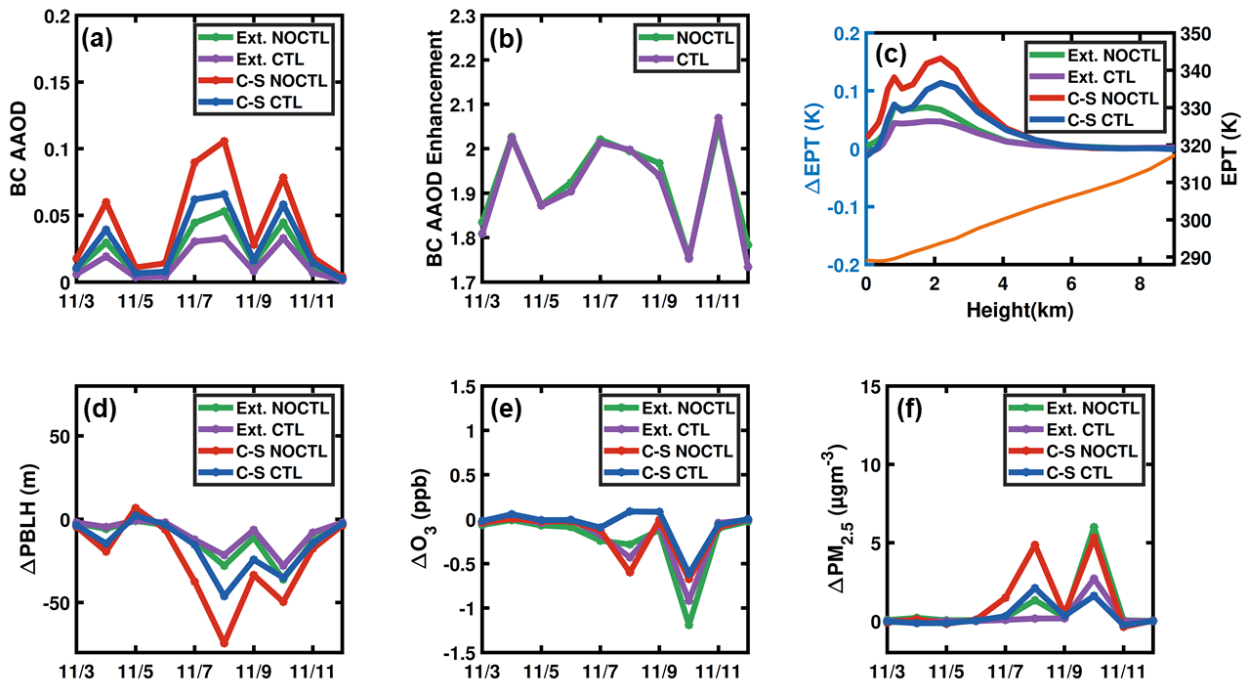


Figure 4: Daytime (10:00-17:00 local time) mean BC AAOD in Beijing inferred from different simulations (a) and the BC AAOD enhancement in the NOCTL and CTL scenarios (b); BC absorption induced changes in EPT (orange line indicates the vertical profile of EPT) (c), PBLH (d), O₃ (e), and PM_{2.5} (f).

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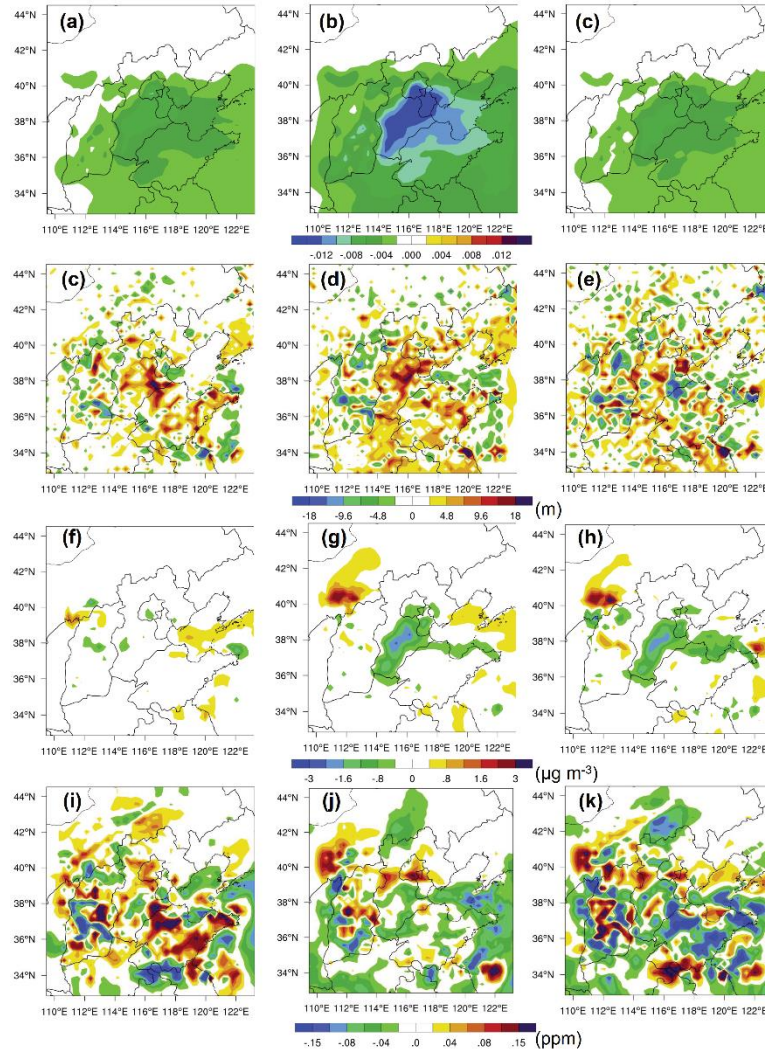


Figure 5: Spatial distribution of daytime (10:00-17:00 local time) mean BC AAOD (first row), and mean BC absorption induced changes after emission reductions (CTL minus NOCTL) in PBLH (second row), PM_{2.5} (third row), and O₃ (fourth row); first, second and third columns represent $\Delta_{emission-Ext}$, $\Delta_{emission-CS}$, and Δ_{aging} .

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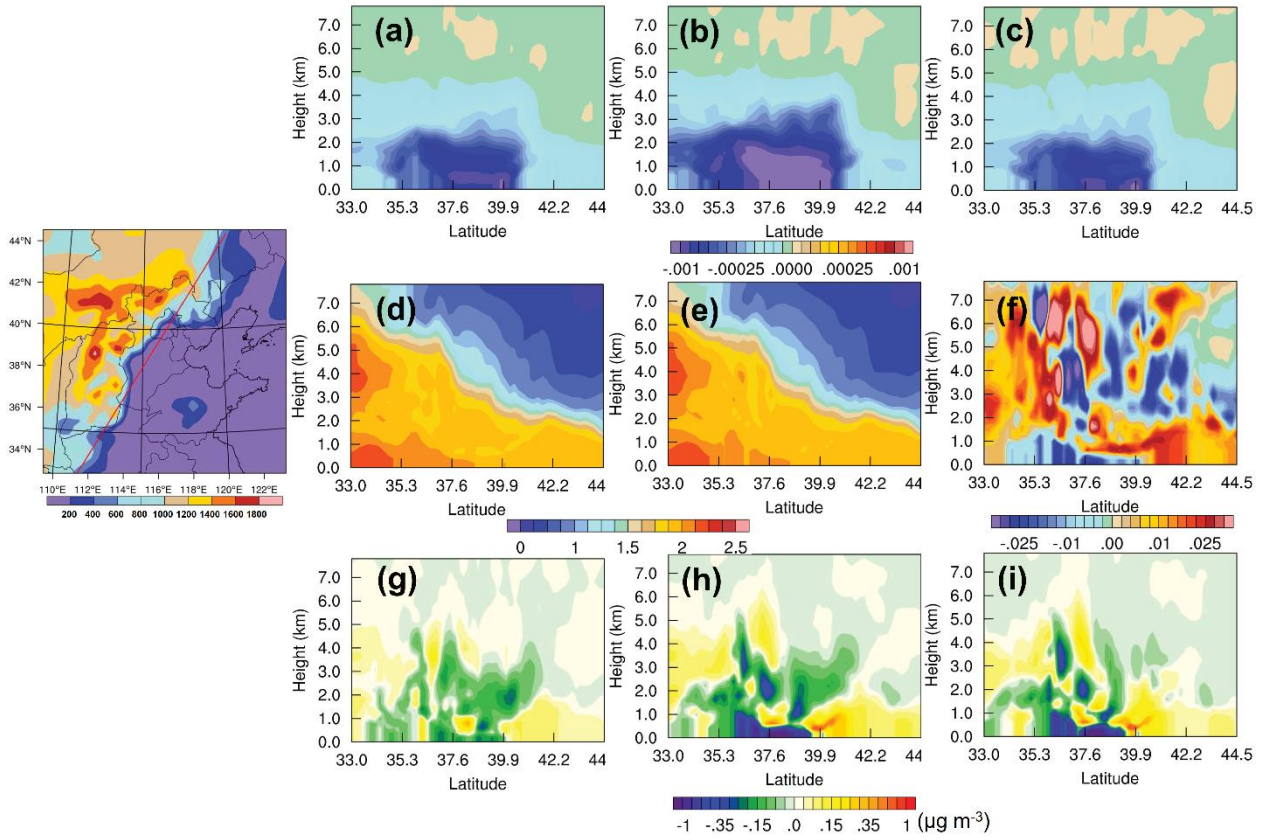
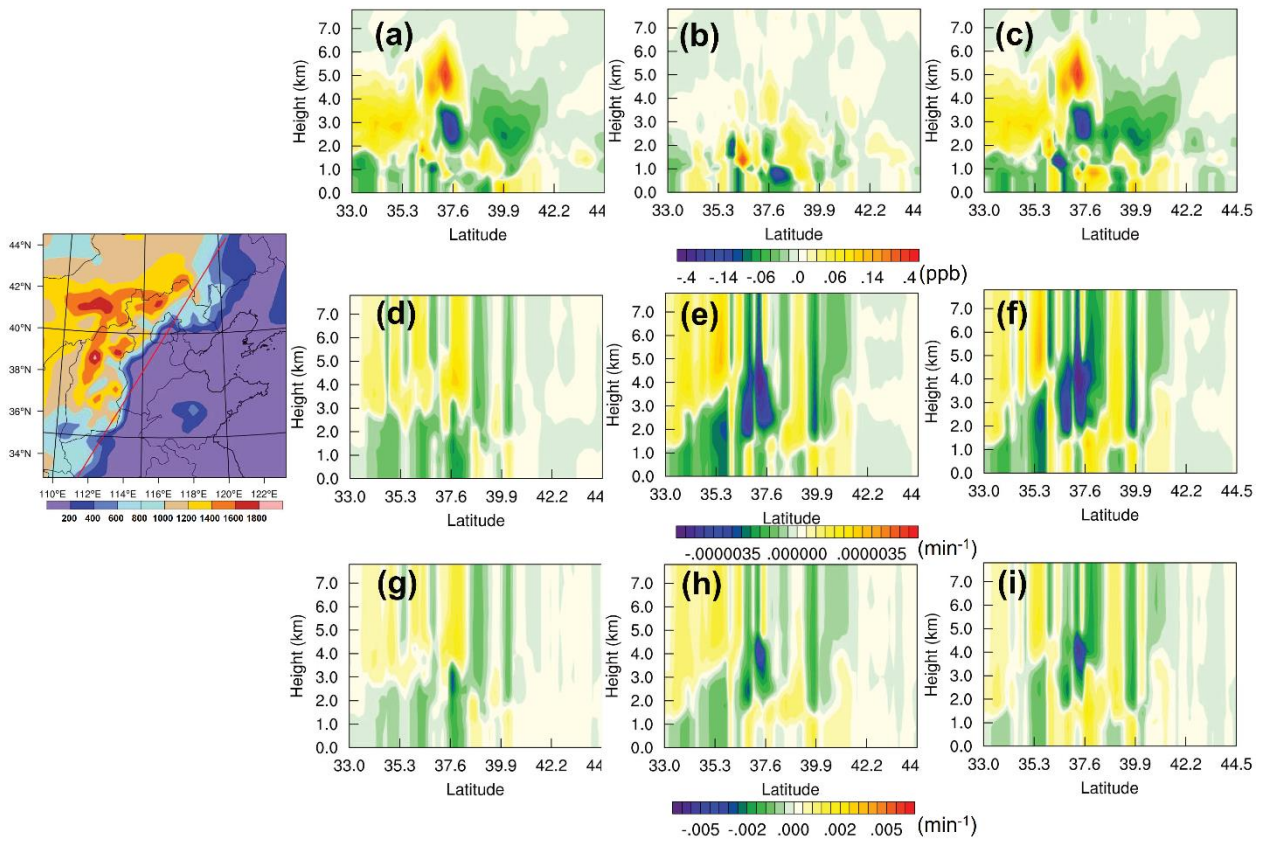


Figure 6: Cross sections of daytime (10:00-17:00 local time) mean changes in BC absorption coefficient (first row, CTL minus NOCTL), E_{ab} (second row), BC absorption induced changes after emission reductions (CTL minus NOCTL) in $PM_{2.5}$ (third row); first, second and third columns represent $\Delta_{emission-Ext}$, $\Delta_{emission-CS}$, and Δ_{aging} .

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655 **Figure 7: Cross sections of daytime (10:00-17:00 local time) mean changes in BC absorption induced changes after emission reductions (CTL minus NOCTL) in O₃ (first row), O₃1D photolysis rate (second row), and NO₂ photolysis rate (third row); first, second and third columns represent $\Delta_{emission-Ext}$, $\Delta_{emission-CS}$, and Δ_{aging} .**

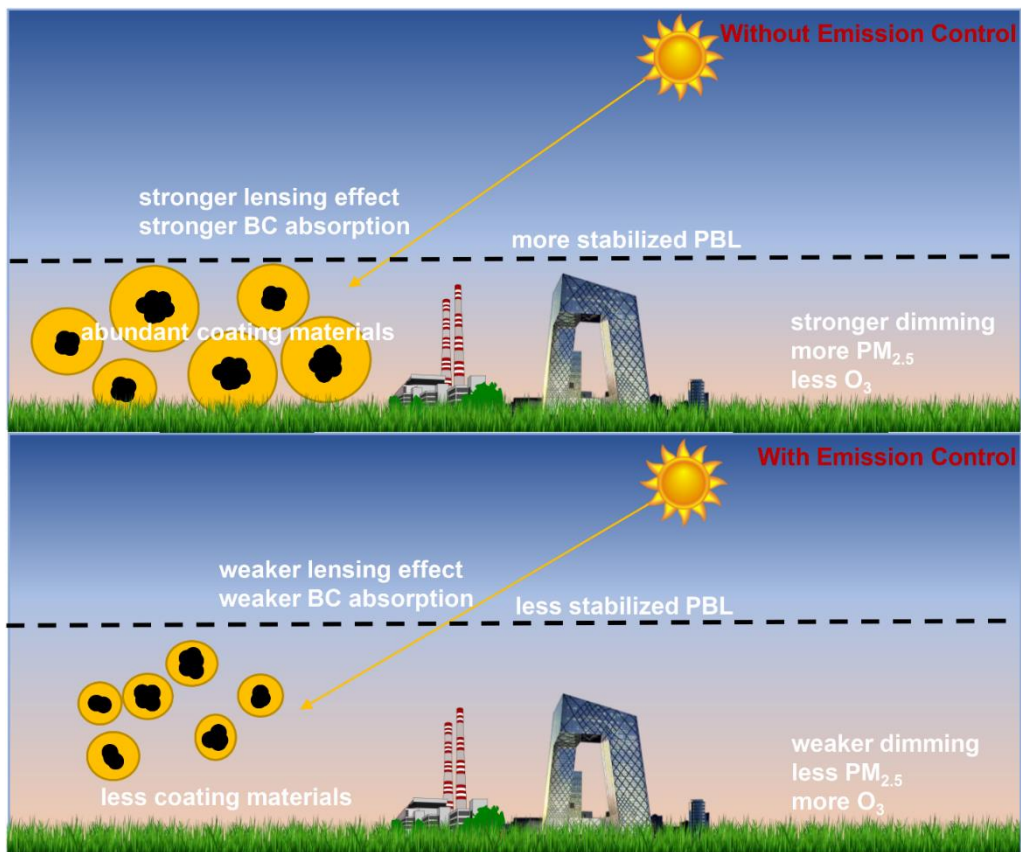


Figure 8: Conceptual scheme of the effects of emission control during APEC on light absorption capability of BC, PBL and air quality.

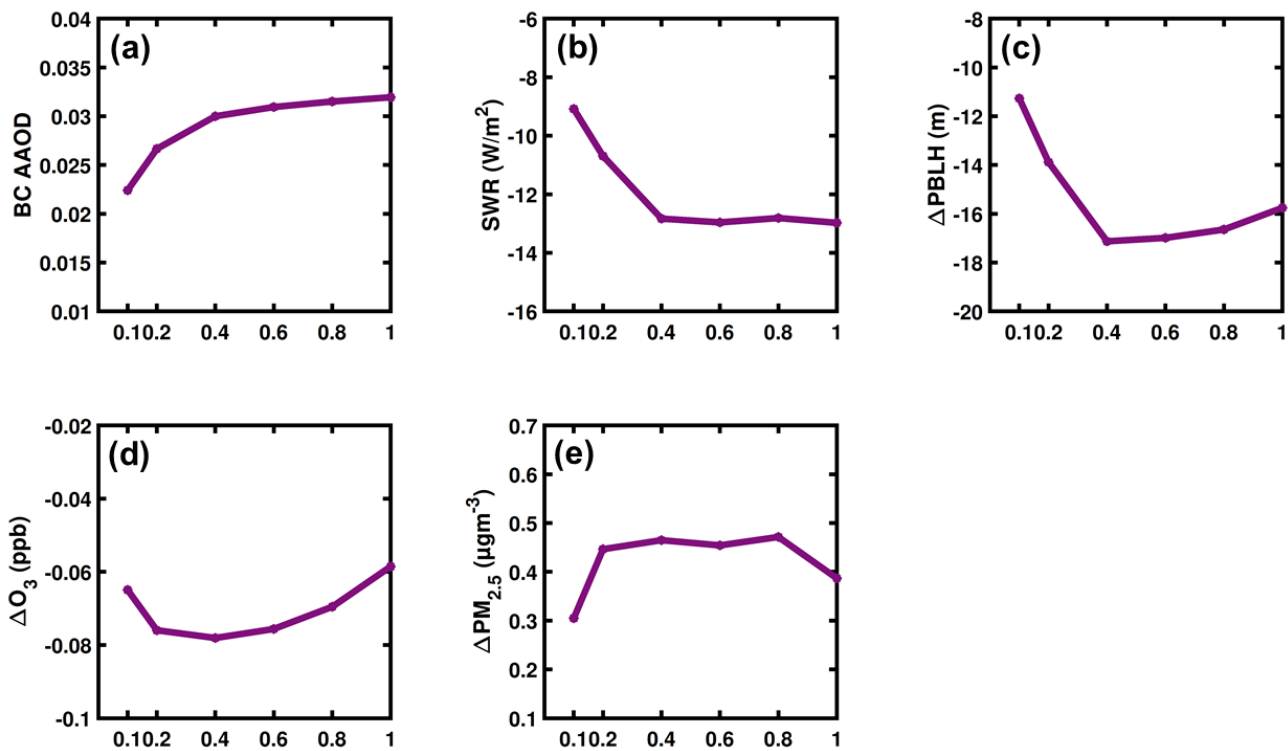


Figure 9: Daytime (10:00-17:00 local time) mean BC AAOD in Beijing (a) BC absorption induced changes in shortwave radiation (b), PBLH (c), O₃ (d), and PM_{2.5} (e) as a function of fraction of non-BC particles mixed with BC.