



Modeling simulation of aerosol light absorption over the Beijing-Tianjin-Hebei region: the impact of mixing state and aging processes

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Abstract. The mixing state and aging characteristics of black carbon (BC) aerosols are the key factors in calculating their

- optical properties and quantifying their impacts on radiation balance and global climate change. Considerable uncertainty still exists in the absorption properties of BC-containing aerosols and the absorption enhancement (E_{abs}) due to the lensing effect. It is crucial to reasonably represent the mixing of BC with other aerosol components to reduce the uncertainty. In this study, the absorption properties of PM_{2.5} were investigated based on the nested air quality prediction model system (NAQPMS) with different assumptions of the aerosol mixing state. The absorption coefficient (b_{abs}) is highest under uniform
- 20 internal mixing, lower under core-shell mixing, and lowest under the assumption of external mixing. The result under coreshell mixing is closest to the observation. The aging process and coating thickness were well produced by the advanced particle microphysical module (APM) in NAQPMS. Then the fraction of embedded BC and secondary components coating aerosols was used to constrain the mixing state. The E_{abs} at 880 nm over the Beijing-Tianjin-Hebei region was 2.0~2.5 under core-shell mixing. When the fraction of coated BC and the coating layer are resolved, the E_{abs_880} caused by the lensing effect
- 25 can decrease by 30~43% to 1.2~1.7, which is close to the range reported in previous studies. This study highlights the importance of representing the microphysical processes governing the mixing state and aging of BC and provides a reference for quantifying its radiative effect.

1 Introduction

Aerosols have important environmental and climatic effects, affecting not only transportation and public health but also the 30 global radiation balance (Xu et al., 2013; IPCC, 2021). Mainly originating from incomplete burning, black carbon (BC) is an important component of aerosols. The light absorption of BC particles is enhanced by the "lensing effect" of coating, which affects the heating of the atmosphere by BC (Fuller et al., 1999). When BC is coated by hydrophilic components, it can act



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as cloud condensation nuclei affecting cloud and rainfall. Previous studies have demonstrated the important radiative forcing effect of BC (Jacobson, 2013; Bond et al., 2013). However, uncertainty in calculating the optical parameters of BCcontaining aerosols still exists. It is challenging to quantify the radiative effect of BC.

The mixing state describes the distribution of properties across a population of particles (Riemer et al., 2019). Externally mixed means that each particle in a population is composed of a single species. Internally mixed means that each particle in the population consists of the same mixture of all chemical species (Stevens and Dastoor, 2019). The aerosol mixing state is dynamic and changes due to several processes, such as emission, new particle formation, transport, condensation, and coagulation processes. Purely internal and external mixing states are rare in the real atmosphere (Bondy et al., 2018). In addition, observations have shown that not all BC particles are coated, and not all secondary aerosols are coated on BC cores (Li et al., 2016). Transmission electron microscope has shown that only a proportion of BC aerosols is embedded (Wang et al., 2021b). The fraction of thickly coated BC in Beijing winter reduced from 48% to 29% from 2012 to 2018 (Wu et al., 2021). The presentation of aerosol mixing states in atmospheric models at different scales was highlighted

- 45 by Riemer et al. (2019). In many models, aerosols are presented in a few modes with assumed size distribution and the same mixing state and compositions in a mode (Liu et al., 2016; Mann et al., 2010). The sectional approach is used in limited models to represent the mixing state (Yu et al., 2012; Matsui et al., 2014; Matsui, 2016; Yu et al., 2015). Furthermore, particle-resolved models can accurately simulate BC mixing states, but it is applied only to the box and single-column models due to the high computational cost (Riemer et al., 2009; Zaveri et al., 2010; Curtis et al., 2017). Yao et al. (2022)
- 50 verified the important yet complicated role of mixing state in calculating aerosol optical properties using an ensemble of 1800 aerosol populations from particle-resolved simulations. Trade off the detail presentation and computational cost, most chemical transport models (CTMs) typically simplify aerosol representation by tracking separate aerosol populations rather than individual particle components (Riemer et al., 2019). Mie theory based on a simple fixed mixing state (external, fully internal, or core-shell mixing) assumption is often used in chemical transport models (Li et al., 2020; Gao et al., 2020).
- 55 Comparison between different mixing states conducted in previous studies showed that the BC absorbing properties are sensitive to the mixing state assumptions. Curci et al. (2019) found that aerosol optical depth (AOD) is mainly determined by the aerosol mass and only secondarily affected by the mixing state, however, the absorption enhancement (E_{abs}) values depended on the mixing assumption made in the model. The underestimation in modeled absorption AOD decreased from 66% in the external mixing case to 43% in the core-shell mixing case (Tuccella et al. 2020). Partial internal mixing is the most
- 60 likely mixing state of aerosols. The fraction of internally mixed particles can be calculated using the parameterization of Cheng et al. (2012) and Curci et al. (2019). The fraction of core-shell can be parametrized as a function of the bulk volume ratio (Hu et al., 2022). The mixing state index and mass ratio of coating to BC were used to improve the BC mixing state presentation and aerosol particles in the accumulation mode were partitioned into BC-free and BC-containing particles (Shen et al., 2024). E_{abs} calculated in a partial internal mixing state were approximately 10% lower than those from core-shell
- 65 mixing simulations (Tuccella et al., 2020). However, quantitative investigations considering the evolution of mixing states



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based on microphysical properties are limited (Li et al., 2024). Therefore, the fraction of coated BC and the coating fraction of other components based on microphysical processes should be considered in the optical calculation.

Coating thickness and the heterogeneity of the mixing state are proposed to be the main reasons explaining the gap between field observation, lab investigation, and model simulation in light absorption enhancement (Zhao et al., 2021; Fierce et al., 2020; Wang et al., 2021a). The coating fraction significantly influences the absorption of BC particles. The coating thickness of BC particles can be detected by a single-particle soot photometer (SP2), although the lower measurement limits of SP2 result in an overestimation of the concentrations of pure BC (Zhao et al. 2020), the mass ratio of coating to BC core (MR) can highly impact absorption enhancement (Liu et al. 2017; Zhao et al., 2021). The black carbon aging process is an

- important source of uncertainty in the assessment of its contribution to global warming (Wang et al., 2022). Black carbon can be coated by other aerosols, which increases the complexity of optical properties. Kang et al. (2023) showed that the aging of BC at night in the residual layer can be higher than in the daytime and enhance its light absorption. The aging degree and mixing state can change very quickly in a polluted environment (Peng et al., 2016). An inadequate understanding of the mixing state of BC greatly hinders the assessment of its climate effects (Huang et al., 2023). However, due to this complexity, few three-dimensional (3-D) models sufficiently resolve BC aging processes (Xie et al., 2023; Zhang et al.,
- 80 2018). A 3-D modeling study found that the aging time of BC varied greatly and showed significant spatial heterogeneity over the polluted areas in China (Chen et al., 2017b).

In this study, the nested air quality prediction model system (NAQPMS) coupled with an advanced particle microphysics module was used to investigate the absorption properties of aerosols over Northern China in November 2018. The simulation of aerosol components was conducted using the 3-D air quality model. Firstly, three ideal mixing states

85 (external, internal mixing, and core-shell mixing) are considered to study the effect of the mixing state by a Flexible Aerosol Optical Depth module (FlexAOD) based on observation and simulation of PM_{2.5} components. Then, the fraction of coated BC based on the aging degree of BC, the fraction of coating, and the detailed microphysics process were considered during the absorption calculation. Finally, the impact of aging degree on light absorption enhancement was investigated.

2 Data and methods

90 2.1 Observation data

The study period is from November 1, 2018, to November 30, 2018. Observations of PM_{2.5} components were obtained from China National Environmental Monitoring Centre. The observation site in Beijing (BJ), located at the China National Environmental Monitoring Centre (40°2'N, 116°41'E), is a typical urban site. Water soluble species (Na⁺, K⁺, sulfate, nitrate, ammonium, and Cl⁻) in PM_{2.5} were measured by Gas and Aerosol Collector (GAC), and particles were collected by wet

95 denuder and detected by Ion chromatography. OC and EC are detected by the Thermo optical transmittance method. OC could further be classified into primary organic carbon (POC) and secondary organic carbon (SOC) using a revised



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(3)

elemental carbon (EC) tracer method (Zhao et al., 2013). Then secondary organic aerosols (SOA) and primary organic aerosols (POA) were calculated with the following equations (Castro et al., 1999):

$SOC = OC - EC \times (OC/EC)_{min}$	(1)
$SOA = 1.6 \times SOC$	(2)

$$POA = 1.6 \times OC - SOA$$

In this study, (OC/EC)_{min} calculated from the observed data was 1.16. Organic matter (OM) was assumed to be 1.6 times of OC.

- The light extinction parameters were measured at the tower site of the Institute of Atmospheric Physics (IAP), Chinese
 Academy of Sciences (39°58′28″N, 116°22′16″E) in Beijing. The absorption coefficient at a wavelength of 880nm was directly measured by a seven-wavelength Aethalometer (AE33, Magee Scientific Corp.) (Sun et al., 2021). The extinction coefficient (*b_{ext}*, *λ*=630 nm) of PM_{2.5} was measured by a cavity-attenuated phase shift extinction monitor (CAPS PMext Aerodyne Research Inc.). The absorption coefficient at 630 nm is derived using a fitted power law relationship at seven wavelengths (Ran et al., 2016). The absorption coefficient at 880 nm was used to analyze the performance of the model as
 BC is the major contributor to aerosol absorption at 880nm. PM_{2.5}, component, and absorption data were used to evaluate the
 - model performance.

2.2 Air quality model

NAQPMS is a 3-D Eulerian terrain-following chemical transport model developed by the Institute of Atmospheric Physics, Chinese Academy of Sciences (Wang et al., 2001). The NAQPMS model coupled with an advanced particle microphysics

- 115 module (APM, Yu and Luo, 2009) was used in this study (Chen et al., 2014). NAQPMS includes physical processes such as advection, convection, diffusion, and deposition, and chemistry processes such as gas-phase chemistry, aqueous chemistry, and aerosol processes. A volatility basis set (VBS) framework for secondary organic aerosols has been coupled to NAQPMS to improve the performance of SOA (Yang et al., 2019; Chen et al., 2021). The APM module includes microphysical processes like nucleation, condensation, evaporation, and coagulation (Yu and Luo, 2009; Chen et al., 2021). Particles are
- 120 represented by the sectional bin scheme in the APM. Secondary inorganic particles are distributed by 40 bins covering 0.0012–12 μm. BC and OC are represented using 28 bins, and other primary particles such as dust and sea salt are represented by 4 bins. The evolution of particle size distributions and aging processes of BC due to condensation and coagulation are well reproduced by the model (Chen et al., 2017a; Du et al., 2019).
- A two-nested model domain was set up in this study. The parent domain covers Northern China at a resolution of 27 km and the second domain covers Beijing-Tianjin-Hebei and surrounding regions with a resolution of 9 km, see Figure 1. To represent fine vertical structures, 30 vertical levels were adopted, including 17 levels below 2 km. The mesoscale model WRF was used to provide the meteorological field for NAQPMS. The initial and boundary conditions for meteorology were provided by NCEP final reanalysis data (FNL) every six hours (https://rda.ucar.edu/datasets/ds083.2/), and MOZART model provided initial chemical fields. The Multi-resolution Emission Inventory for China (MEIC) developed by Tsinghua





130 University with a resolution of 0.1×0.1° was used (http://meicmodel.org.cn). The base year of the emission inventory was 2018. MEIC covers 10 species including BC, OC, PM_{2.5}, PM₁₀, CO, NH₃, SO₂, NOx, CH₄, and VOCs. The configuration of WRF and NAQPMS can be seen in Table S1. The simulation of WRF and NAQPMS started on October 24, 2018, and the first 7 days were set aside as spin-up time.



135 Figure 1. (a) The model domain. (b) Emission rate of primary PM_{2.5} and the location of observation sites. Black dots are pollution sites.

2.2.1 Flexible Aerosol Optical Depth (FlexAOD) module

- In this study, the particles are assumed to be spherical and Mie theory (Mie, 1908) is applied to study their optical properties. The FlexAOD (http://pumpkin.aquila.infn.it/flexaod/) (Curci et al., 2015) was employed to calculate extinction and single scatter albedo. The components of PM_{2.5} and relative humidity (RH) at ground level are used as input parameters. The average volume of particles is computed for each species by dividing mass by the species' density. Mixing states considered in the study include the external mixing assumption (EXT), internal homogeneous assumption (HOM), and the core-shell assumption (CS). For the external mixing assumption, extinction is the sum of each species under specific relative humidity. For internal mixing cases (HOM and CS), the particles conform to a lognormal size distribution. For the internal homogeneous assumption, the volume average refractive index is a function of particle size over all species. For the coreshell assumption, the refractive index for BC core and homogeneously mixed shell (secondary inorganic aerosols and secondary organic aerosols) are calculated separately. The Mie code based on Toon and Ackerman (1981) is used for the core-shell internal mixing, and the code based on Mishchenko et al. (1999) is adopted for external and homogeneous internal mixing. The size distribution of the different aerosols is taken from the OPAC (Optical Properties of Aerosols and Clouds)
- 150 database (Hess et al., 1998). The mean diameter of BC is assumed to be 30 nm based on Dentener et al. (2006). The density,





complex refractive index, particle hygroscopic growth factor, mean radius, and standard variation of log-normal size distribution are shown in Table S2.

2.2.2 Optical module based on APM

APM is a size-resolved, mixing-state-resolved advanced particle microphysics model coupled in NAQPMS. The mixing
state in APM is assumed to be semi-external mixing, which includes internal mixing, external mixing, and core-shell mixing. The seeding particles generated by emission and nucleation (including BC, OC, sulfate, dust, and sea salt) can be coated by secondary particles (including sulfate, nitrate, ammonium, and SOA) through condensation, coagulation, chemical reactions, equilibrium uptake, and hygroscopic growth processes. Sulfate coated by SIA or SOA is considered to be internal mixing. BC, OC, dust, and sea salt coated with SIA or SOA are considered to be core-shell mixing. These coated particles are
externally mixed. The mixing of BC particles with other aerosol components can be well resolved hourly. More details can be seen in Yu and Luo (2009) and Chen et al. (2017b).

When calculating the optical parameters of aerosols, the scheme by Yu et al. (2012) was used. The particles are assumed to be spherical and key particle optical properties including extinction efficiency, single scattering albedo, and asymmetry parameter at each wavelength are calculated by Mie theory based on the core diameter, shell diameter, real and imaginary components of the refractive index of core and shell. The core-shell code based on Toon and Ackerman (1981) is used in APM. To reduce computation cost, three lookup tables are used: one for particles without solid absorbing cores, the second for coated BC, and the third for coated dust. The volume-averaged refractive indices of species other than BC and dust are calculated based on the composition simulated by NAQPMS. Details can be seen in Yu et al. (2012) and references therein.

170 2.3 Sensitivity test design

In this study, the 3-D chemical transport model NAQPMS coupled with the advanced particle module (APM) was used to reproduce the evolution and spatial distribution of pollutants. The mass concentration, size distribution and mixing state of aerosols are calculated by NAQPMS+APM. FlexAOD is a module that calculates the extinction property of aerosols under different mixing state assumptions based on Mie theory and a fixed size distribution, using the input of aerosol components'

- 175 mass concentration and relative humidity as shown in Sect. 2.2.1. There are two approaches to calculating optical properties. The absorption property of aerosols can be investigated by FlexAOD with the input of component concentration simulated by NAQPMS+APM and assumed size distribution. The fraction of embedded BC and the traction of coating aerosols calculated by NAQPMS+APM can be used to constrain the mixing state in FlexAOD. In the other approach, the absorption property can be investigated by the optical module based on APM with core and shell information calculated by NAQPMS+APM as shown in Sect. 2.2.2. Then a series of sensitivity tests were designed to explore the impact of mixing
- state, components, aging process, and detailed microphysical processes (Table 1).





Firstly, to see the effect of mass concentration and mixing state on the optical properties, sensitivity tests with different mixing states (external, internally homogeneous, and core-shell) were conducted using FlexAOD. EXTO, HOMO, and CSo refer to cases calculated using FlexAOD with observed components as input under external, homogeneous internal, and coreshell mixing states, respectively. EXTs, HOMs, and CSs refer to cases calculated using FlexAOD with components simulated by NAQPMS+APM as input under external, homogeneous internal, and core-shell mixing states, respectively. Comparing EXTo, HOMo, and CSo can show the impact of the mixing state. Comparing EXTs, HOMs, and CSs can also show the impact of the mixing state. Comparing CS₀ with CS_s, the impact of mass concentration on optical properties can be obtained. Secondly, to see the impact of the aging process (fraction of embedded BC core and fraction of coating aerosols), simulations using partial core-shell mixing state in FlexAOD, CS-F_{in} and CS-F_{in}F_c, were designed. Additionally, components, size distribution, and mixing state simulated by NAQPMS+APM were used to calculate the optical properties (CS-APM). The impact of the microphysical process can be investigated by comparing CS-F_{in}F_c with CS-APM.

Case	Method	Input	Size distribution	Mixing state			
EXTo	FlexAOD	observed	fixed	external			
HOMo	FlexAOD	observed	fixed	internal homogeneous		internal homogeneous	
CSo	FlexAOD	observed	fixed	core-shell			
EXTs	FlexAOD	simulated	fixed	external			
HOM _s	FlexAOD	simulated	fixed	internal homogeneous			
CS_S	FlexAOD	simulated	fixed	core-shell			
CS-F _{in}	FlexAOD	simulated	fixed	partial core-shell and partial bare BC			
CS F. F	FloyAOD	simulated	simulated fixed partial core-shell, partial bare partial coating aerosol		fixed	partial core-shell, partial bare BC and	
CD-1 inf c	TRAGD	sinulated					
CS-APM	APM	simulated	simulated	semi-external (hourly)			
Impact			Des	cription			
EXT ₀ vs. HOMo vs. CSo Impact of mixing state when it			when inputting observed data				
EXT _s vs. HC	M _S vs. CS _S	Impact of mixing state when inputting simulated data					
CSo vs. CSs			Impact of aerosol mass concentration				
CSs vs.	CS-F _{in}	Impact of aging process (fraction of embedded BC)					

Table 1 Simulation test design





CSs vs. CS-F_{in}F_cImpact of the aging process (fraction of embedded BC and coating shell)CS-F_{in}F_c vs. CS-APMImpact of detailed microphysical process

2.4 Model evaluation

- 195 Statistical parameters such as the correlation coefficient (R), normalized mean bias (NMB), index of agreement (IOA), the fraction of the simulations within a factor of two of the observations (FAC2), mean fractional bias (MFB) and mean fractional error (MFE) were used in this study to evaluate the performance of NAQPMS (Table S3). NAQPMS reproduced the temporal distribution of PM_{2.5} in Beijing well (Fig. 2). As shown in Table S4, the R between the observed and simulated hourly PM_{2.5} concentrations of six sites over Beijing-Tianjin-Hebei and surrounding regions were within 0.55~0.76. The
- 200 NMB was within -0.22~0.13, which satisfied the model performance criteria proposed by Emery et al. (2016). There was only a small overestimation of 13% in the simulated $PM_{2.5}$ in Zhengzhou. And the IOA reached more than 0.72. The MFB and MFE of $PM_{2.5}$ in all six sites were within the benchmarks, which satisfied the model performance criteria proposed by Boylan and Russell (2006).







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Figure 2. Model evaluation of PM2.5 at eight sites in the Beijing-Tianjin-Hebei region in November 2018.

NAQPMS also exhibited good performance in representing the $PM_{2.5}$ components in Beijing (Fig. 3). The R values

between the observed and simulated nitrate, sulfate, ammonium, element carbon, primary organic carbon, and secondary organic carbon were 0.74, 0.83, 0.74, 0.55, 0.49, and 0.47 in Beijing, respectively. However, the simulation of secondary inorganic aerosols was underestimated by -62%~-8%. This is likely caused by insufficient heterogeneous formation of sulfate and nitrate (Li et al., 2018). Black carbon and primary organic aerosols were overestimated by 7.2% and 26.1%, which is probably related to the emission inventory. Monthly mean emissions were used in this study and there is substantial uncertainty in emission inventory (Li et al., 2017). However, the reasonable simulation of aerosol mass concentration lays a solid foundation for simulating the optical properties of BC-containing aerosols.







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Figure 3. Simulated and observed PM_{2.5} components in Beijing

3 Results

3.1 Absorption properties based on observed components

3.1.1 Description of observations

The time series and proportion of various chemical components of PM_{2.5} are shown in Fig. 4. During the study period, the average mass concentration of PM_{2.5} was 74.4 ± 68.7 µg m⁻³. Nitrate is the main component of PM_{2.5}, accounting for 36.4% on average, followed by organic matter, ammonium, and sulfate, accounting for 16.6%, 15.4% and 11.5%, respectively. EC, crustal elements, and chloride salt accounted for 3.9%, 8.8%, and 4.1%, respectively. The average RH during the period was $39 \pm 17.9\%$ and the temperature was 8.3 ± 3.2 °C. The average $b_{sca} (\pm 1\sigma)$ and $b_{abs} (\pm 1\sigma)$ at 630 nm during the study period were 169.1 ± 212.3 Mm⁻¹ and 46.5 ± 48.5 Mm⁻¹ in Beijing, respectively. The average $b_{abs} (\pm 1\sigma)$ at 880 nm during the study period were 30.7 ± 25.2 Mm⁻¹. The decrease in visibility is mainly caused by particle scattering extinction. The b_{sca} and b_{abs}

in this study were much lower than those observed in Beijing in the winter of 2016 (Xie et al., 2019), but the b_{ext} in this study





was higher than that observed in Beijing in the winter of 2019 (Sun et al., 2021), indicating that the decreases in $PM_{2.5}$ in recent years also caused similar reductions in extinction coefficients.



230 Figure 4. Evolution of observed PM_{2.5}, components, aerosol extinction coefficient, and meteorology parameters in November 2018 at Beijing

3.1.2 Absorption coefficient calculated by FlexAOD based on observation

The observed $PM_{2.5}$ components and RH were used to calculate the optical properties using FexAOD. It should be noted that the times missing component data were excluded when calculating optical properties. Comparison between the observed and simulated absorption coefficient showed that the simulations by FlexAOD under the three mixing state assumptions are highly correlated with the observation, and the correlated coefficient can reach 0.88. However, using different mixing state assumptions led to widely varying results, see Figure 5. On average, the b_{abs} of 880 nm calculated for the core-shell mixing state was 2.3 times higher than that of external mixing. For the external mixing state, the calculation was underestimated by 59%, while it was overestimated by 79% under uniform internal mixing. The simulation for core-shell mixing was closest to

240 observations, with an underestimation of 4.7%. The absorption coefficient under uniform internal mixing is the highest,





followed by core-shell mixing and the calculation for external mixing is the lowest. This is consistent with the findings of Curici et al. (2019). The assumption of internal mixing or external mixing is not realistic. Partial internal mixing with partial coating is closer to reality and it should be considered for absorption calculations as reported by Curci et al. (2019).



245 Figure 5. Observed absorption and calculated absorption at 880nm under external mixing (EXTo), homogeneous internal (HOMo), and core-shell mixing (CSo) by FlexAOD based on observed components.

3.2 Absorption property based on simulations by NAQPMS

Comparison of the absorption coefficients between observation and calculation with FlexAOD based on PM_{2.5} simulation from NAQPMS with different mixing states can be seen in Table 2. In this study, only times in which the simulated PM_{2.5} was within a factor of two of observation were considered in the optical calculation. The average of observed b_{abs} is 43 Mm⁻¹. 250 There were large variations in the absorption coefficient under different mixing states. In the EXTs case, the absorption coefficient at 880m was underestimated by 54%, and only 40% of the modeled values were within a factor of 2 of the observations. In the HOMs case, the absorption at 880m was overestimated by 95%, and 58% of the modeled values were within a factor of two of observations. In the CS_S case, FAC2 increased to 0.93 compared with EXTs simulations, and the model overestimated the absorption coefficient by 10%. This was related to the overestimation of black carbon. In this study, 255 black carbon and POA are assumed to have light absorption properties and only BC particles act as cores in FlexAOD. The core-shell mixing state produces the most accurate results compared to observations. Comparing the results of CS_s with CS_o (where calculations with FlexAOD were based on observed components under core-shell mixing state), the results show that the effect of simulated components on the absorption coefficient can be up to 15%, which is much smaller than the impact of 260 the mixing state.







Figure 6. Comparison of observed and simulated absorption coefficient at 880 nm under different mixing states (EXT_s, HOM_s, CS_s, CS-F_{in}, CS-F_{in}F_c, and CS-APM) at IAP in Beijing.

	Table 2. Intercomparison of the performance of absorption coefficient at 880 nm under different mixing states. b is the ratio of
265	simulation to observation.

Schemes	b	R	NMB	FAC2
EXTs	0.46	0.82	-0.54	0.40
HOMs	1.95	0.79	0.95	0.58
CSs	1.10	0.82	0.10	0.93
CS-F _{in}	0.83	0.77	-0.17	0.84
CS-F _{in} F _c	0.66	0.78	-0.34	0.66
CS-APM	0.58	0.79	-0.42	0.72





3.3 Constraint of the fraction of embedded BC and secondary components coating aerosols

In the real world, the mixing state of particles is complex. Wang et al. (2021b) using an electron microscope found that the embedded fraction of BC significantly influenced the absorption. In the extremely polluted winter period of January 2013, more than half of BC particles were thickly coated by non-refractory materials (Wu et al., 2016). Along with the implementation of the Air Pollution Prevention and Control Action Plan, the mass of BC and the fraction of thickly coated BC changed (Wu et al., 2021). Cheng et al. (2012) proposed that the fraction of internally mixed particles can be parameterized based on oxidized nitrogen oxides and total reactive nitrogen. Curci et al. (2019) used the mass ratio of secondary inorganic aerosols and organics to BC as the fraction of internally mixed particles.

- Emitted hydrophobic black carbon becomes hydrophilic due to aging processes. In this study, the aging of BC can be resolved by NAQPMS+APM. The detailed aging processes of aerosols are considered in a physical manner. The model represents the aging processes by simulating condensation and coagulation. The ratio of hydrophilic BC to total BC is used as a proxy for the fraction of embedded BC. The evolution of the ratio of particle diameter to BC-core size (Dp/Dc), the fraction of embedded BC (F_{in}), and the fraction of secondary components coating on BC (F_c) at Beijing is shown in Fig. 7. When F_{in} is equal to 0, it means the BC is externally mixed with other aerosols, and when F_{in} is equal to 1, it means all BC
- 280 particles are coated by other aerosols. The average ratio of F_{in} during the study period in Beijing is 34.1%, which is a bit lower than the ratio of 0.48 and 0.63 obtained by the method of Cheng et al. (2012) and Curci et al. (2019), respectively. Also, F_{in} in this study is closely related to the ratio of secondary inorganic aerosols to BC, with an R of 0.72. Zheng et al. (2022) also found that secondary inorganic aerosols dominated the light absorption enhancement with online observational datasets. We consider a separate case, CS-F_{in}, where the F_{in} fraction of BC particles is core-shell mixed with other aerosols,
- and a 1- F_{in} fraction of BC particles is bare and external mixing. The calculated BC absorption at 880 nm (b_{abs_880}) for the CS- F_{in} case was 35.5 Mm⁻¹, which was close to the measured mean value, and 84% of simulations of absorption coefficients were within a factor of two of the observations.







Figure 7. Evolution of ratio of particle diameter to BC-core size (Dp/Dc), the fraction of embedded BC (Fin), and the fraction of 290 secondary components coating on BC (Fc) at IAP in Beijing. "APM simulated" refers to parameters simulated by the advanced particle microphysics module in NAOPMS.

Aerosols in the atmosphere include BC-containing aerosols (coated BC and bare BC) and BC-free aerosols (Zhao et al., 2022). In this study, the fraction of secondary aerosol coating on BC is also considered in the optical calculation. As sulfate aerosols include sulfate coating on BC, OC, dust, and sea salts, then the fraction of sulfate coating on BC was used as the 295 fraction of secondary components coating on BC (F_c). When F_c is equal to 0, there is no coating over BC particles, and when F_c is equal to 1, it means all other aerosols are coated on BC. We thus consider a scenario CS- $F_{in}F_c$ where F_{in} fraction of BC particles are core-shell mixed and the F_c fraction of secondary components and other aerosols are externally mixed. The average F_c during the study period in Beijing is 34.3%. The calculated BC absorption at 880 nm ($b_{abs 880}$) for the CS- $F_{in}F_c$ case was 28.1 Mm⁻¹, and 66% simulations of absorption coefficients were within a factor of two of observations.

300 As described in section 2.2.2, optical properties were calculated based on Mie theory using the core and shell information calculated by APM considering microphysical processes. The observed absorption coefficient and that simulated by NAQPMS+APM under a semi-external mixing state, namely CS-APM, is shown in Table 2 and Fig. 6f. The results show that the simulated absorption at 880nm matches the observation reasonably well, with an R of 0.79, although there was an underestimation with NMB of 0.42. The CS- $F_{in}F_{c}$ case considers the fraction of embedded BC and the fraction of secondary components coating BC calculated with APM. Comparison between CS-F_{in}F_c and CS-APM showed the impact of 305





considering the detailed microphysics process on absorption property. The FAC2 of 0.72 in the CS-APM case is greater than 0.66 in the CS- $F_{in}F_c$ case. The underestimation of 42% in CS-APM is larger than the 34% in CS- $F_{in}F_c$. This underestimation can be attributed to the assumed morphology of BC-containing particles, the size distribution of primary particles input to APM, and the concentration of secondary components coated on BC. As shown in Fig. 3, there is an underestimation of 8-62% in the simulation of secondary inorganic aerosols. Even if the mode of the physical process is correct, the coating on BC can

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be underestimated, which affects the absorption characteristics of aerosols.

3.4 Light absorption enhancement due to mixing state

The light absorption enhancement is the ratio of the light absorption coefficient of coated BC and bare BC. E_{abs} is proposed to quantify the lensing effects, however, large uncertainty exists in E_{abs} and the radiative effect of black carbon.

315 3.4.1 The impact of the detailed microphysics process on absorption enhancement

We modified the APM module in NAQPMS so that BC does not mix with other chemical species in the calculation of the microphysics process and optical properties. This sensitivity test was conducted by turning off the coating process in APM. The radiative absorption enhancement was the ratio of the absorption coefficient in the base simulation to that in the sensitivity test.

- The mass ratio of the coating of BC to BC (MR) can be used to represent the aging degree (Du et al., 2019; Wang et al., 2019). To compare with previous studies, E_{abs} at 630nm is shown in Fig. 8. The evolution of E_{abs} and MR shows that E_{abs} is positively correlated to MR, and the R can reach 0.88. This is consistent with Liu et al. (2017) who showed that E_{abs} is closely related to MR. Under the same MR, E_{abs} can vary by 0.49. When MR equals 3, E_{abs} varied by 0.25. The E_{abs} in the CS-APM case in Beijing is much higher than the measurement in Taizhou (Zhao et al. 2021). However, the measurements in
- 325 Beijing by Xie et al. (2019) fall in the range of this study when MR is less than 5. E_{abs} in the CS-APM case is higher than that from the laboratory study in Peng et al (2016) when MR is less than 3, but it is lower when MR is bigger than 5.

The spatial distribution of E_{abs} at 880nm is shown in Fig. 9d. The E_{abs_880} over Beijing-Tianjin-Hebei and the surrounding region was about 1.3~1.8. The E_{abs_880} in the CS-APM case is a bit higher than that in the CS- $F_{in}F_{c}$ case. The spatial distribution of E_{abs_880} also showed lower values of 1.3~1.7 over the source region and higher values of 1.6~1.8 over

330 the outflow region. The average E_{abs} at Beijing at 630nm and 880nm from APM and Mie theory are 1.58 and 1.55.







Figure 8. The absorption enhancement in CS-APM case under different mass ratios (MR) of coating materials and BC core

3.4.2 Impact of the aging processes on light absorption enhancement

The spatial distribution of E_{abs} at 880nm is shown in Fig. 9. The spatial distribution of the absorption enhancement in CS-Fin
and CS-F_{in}F_c cases showed that E_{abs} was lower near the emission source and higher in the outflow region (Bohai and Yellow Sea, Taihang mountain). This is because BC was aged by condensation and coagulation processes during transport in the atmosphere. As shown, the values of E_{abs_880} over Beijing-Tianjin-Hebei and the surrounding region from FlexAOD under the core-shell mixing state are about 2.0~2.5. After considering the fraction of embedded BC, the E_{abs} decreased to 1.3~2.1, representing a decrease of 11%~34%. Considering the fraction of embedded BC and the fraction of coating, the E_{abs}
decreased to 1.2~1.7, representing a decrease of 30%~43%. The values of E_{abs} in the CS-F_{in}F_c case are 1.2~1.5 near the emission sources and 1.5~1.7 over the outflow region. These values are similar to the currently accepted range of 1.2–1.6 (Bond et al., 2013; Matsui et al., 2016; Liu et al., 2017; Curci et al., 2019). The distribution of average E_{abs} and SSA values with height in Beijing is shown in Fig. 10. E_{abs} increased with height while SSA decreased with height in CS-F_{in} and CS-F_{in}

 $F_{in}F_c$ cases. Relatively low E_{abs} values (1.3~1.6) are concentrated in layers below 500 m. This is related to the low-level 345 anthropogenic emission and the ability of BC in the upper layer to be transported over wider regions.







Figure 9. The absorption enhancement at 880 nm in (a) core-shell mixing (CS), (b) CS-F_{in}, (c) CS-F_{in}F_c, and (d) CS-APM. And changes in radiation absorption enhancement in (e) CS-F_{in} and (f) CS-F_{in}F_c cases compared with core-shell mixing.







350 Figure 10. The distribution of Eabs and SSA values with height

In Beijing, the absorption enhancement at 630 nm and 880 nm is 2.65 and 2.39 for the core-shell mixing state using FlexAOD (Fig. 11). When considering the fraction of embedded BC, the E_{abs} in the CS- F_{in} case at 630 nm and 880 nm are 1.94 and 1.80, decreasing by 26.7% and 24.5% compared to the CS case, respectively. If the fraction of secondary aerosol coating on BC is also considered at the same time, the E_{abs} in the CS- $F_{in}F_c$ case at 630 nm and 880 nm are 1.51 and 1.43, decreasing by 43% and 40.2% compared to the CS case, respectively. Therefore, considering the fraction of secondary aerosol is also considered at the same time, the E_{abs} in the CS- $F_{in}F_c$ case at 630 nm and 880 nm are 1.51 and 1.43, decreasing by 43% and 40.2% compared to the CS case, respectively. Therefore, considering the fraction of secondary aerosol is also considered at 0.2% compared to the CS case, respectively.

355 decreasing by 43% and 40.2% compared to the CS case, respectively. Therefore, considering the fraction of secondary aerosol coating on BC, E_{abs} at 630 nm and 880 nm can decrease by 16.2% and 15.7%, respectively, compared to the CS case. The ratios of E_{abs} at 630 and E_{abs} at 880 nm by FlexAOD in this study were both less than 1. This is consistent with the fact that E_{abs} are expected to decrease with increasing wavelength (Liu et al., 2018).







360 Figure 11. The radiation absorption enhancement of 630nm and 880nm under different mixing state assumptions in Beijing.

Comparing the E_{abs} obtained in this study under different mixing states with previous studies, the E_{abs} in CS-F_{in} and CS-F_{in}F_c considering the aging process were a bit higher than other laboratory and ambient measurement studies in Beijing (1.03~1.3) (Wang et al., 2019). Sun et al. (2021) using the thermodenuder (TD) method found that E_{abs} at 870nm at an urban site in Beijing was 1.24 ± 0.15. Zhang et al. (2021) using the mass absorption cross section (MAC) method by SP2 found that E_{abs} at 880nm at a rural site in Gucheng was 1.33 ± 0.57. Cui et al. (2016) found that the E_{abs} increased from 1.4 during fresh combustions to approximately 3 for aged BC at a rural site on the North China Plain. But the results for CS-F_{in}, CS-F_{in}F_c, and CS-APM were lower than other model simulations (Curci et al., 2019; Tuccella et al., 2020). The results show that considering the aging process of BC has a significant effect on the absorption enhancement, and should be considered during model E_{abs} calculation.

370 4 Summary and conclusions

Black carbon-containing aerosols have a significant impact on global warming. However, the extent of the impacts is highly uncertain. Component concentration, mixing state, and aging processes are important parameters. In this study, observed and simulated concentrations of PM_{2.5} components in November 2018 are used with Mie theory to investigate the impact of the mixing state and aging process on the light absorption properties of aerosols.

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5 Through a series of sensitivity tests, a systematic comparison was conducted to explore the impact of components, mixing state, aging process, and detailed microphysics on absorption property. Under the same mixing state with observed



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and simulated components, b_{abs} can be highly impacted by the simulated concentration of PM_{2.5} components. Sensitivity tests with different mixing states (external, internally homogeneous, and core-shell) using FlexAOD showed that different mixing state assumptions led to widely varying results. The absorption coefficient is the highest under uniform internal mixing, lower under core-shell mixing, which is closest to observation, and lowest under external mixing.

Partial internal mixing and partial coating are the closest to reality. The detailed microphysical processes can be resolved by an advanced particle microphysics module in NAQPMS. The ratio of hydrophilic BC and total BC is used as a proxy for the fraction of embedded BC and the fraction of sulfate coating on BC is used as a proxy for the fraction of secondary components coating on BC. Then the fraction of embedded BC and secondary components coating aerosols was used to constrain the mixing state. Considering the fraction of embedded BC and secondary components coating on BC, the mixing state is closer to reality and the simulation of absorption is also acceptable. The NMB of the simulated absorption coefficient has changed from 10% to -34% in Beijing, and the R changed from 0.82 to 0.78.

Accounting for the aging process of BC has a significant effect on radiative absorption enhancement. The E_{abs} at 880 nm over the Beijing-Tianjin-Hebei area reduced from 2.0~2.5 under core-shell mixing state to 1.3~2.1 when considering the fraction of embedded BC, and to 1.2~1.7, a decrease of 30%~42%, when considering the fraction of embedded BC and the fraction of coating. Considering the detailed microphysical processes, E_{abs} in the CS-APM case was positively correlated

- with MR with an R of 0.88. The E_{abs} values in CS- $F_{in}F_c$ and CS-APM cases were a bit higher than those from other laboratory and ambient measurement studies in Beijing, but were within the range of previous studies.
- The optical property can be affected by uncertainties in the size distribution of primary particle emission (Zhou et al. 2012; Matsui, 2016). Geometric radius and standard variation are two important parameters of size distribution. The optical depth of mineral dust and organic was sensitive to standard variation (Obiso and Jorba, 2018). There is a sector and spatial difference in the size distribution of primary emission (Paasonen et al., 2016). Sensitivity tests should be conducted to see the impact of size distribution on σ_{abs} and E_{abs} in future studies. More efforts considering the morphology and the absorption characteristics of coating can also help understand the radiative effect of BC-containing aerosols (Liu et al., 2020; Li et al., 2024).

Overall, this study underscores the importance of the representation of microphysical processes related to BC aerosols and their mixing state. Our results indicate that resolving the fraction of coated BC and the coating layer can significantly impact the calculated E_{abs} . Although modeling the mixing state and microphysics process is a challenge for the chemical transport model, the fraction of aged BC and coating aerosols can be used to constrain the mixing state. This study provides

405 a reference for simulating the radiative effect of black carbon aerosols using three-dimensional models.

Data availability.

The $PM_{2.5}$ observation data can be obtained from the China National Environmental Monitoring Centre (https://air.cnemc.cn:18007/). The simulated data of this study are available upon request to the corresponding author.





Author contributions.

410 HD, JL and XC designed the work. HD performed the simulation and analysis. Gabriele C and ZFW provided the software. YS, XD and SG processed the measurement data. ZW, WY and LW validated the simulated data. HD wrote the original draft with assistance from co-authors. JL, XC and FY reviewed and edited the manuscript.

Competing Interests

At least one of the (co-)authors is a member of the editorial board of Atmospheric Chemistry and Physics.

415 Acknowledgements

We thank the support of the Technological Infrastructure project "Earth System Science Numerical Simulator Facility" (EarthLab). The FlexAOD code can be provided upon request to gabriele.curci@aquila.infn.it.

Financial support.

This research was supported by the National Key R&D Program of the Ministry of Science and Technology, China (grant no.
2022YFC3700703, 2020YFA0607803), the Strategic Priority Research Program (B) of the Chinese Academy of Sciences (XDB0760300), the National Natural Science Foundation of China (NSFC) research project (42207133, 42377105).

References

- Bond, T. C., Doherty, S. J., Fahey, D. W., Forster, P. M., Berntsen, T., DeAngelo, B. J., Flanner, M. G., Ghan, S., Karcher,B., Koch, D., Kinne, S., Kondo, Y., Quinn, P. K., Sarofim, M. C., Schultz, M. G., Schulz, M., Venkataraman, C., Zhang,
- H., Zhang, S., Bellouin, N., Guttikunda, S. K., Hopke, P. K., Jacobson, M. Z., Kaiser, J. W., Klimont, Z., Lohmann, U., Schwarz, J. P., Shindell, D., Storelvmo, T., Warren, S. G., and Zender, C. S.: Bounding the role of black carbon in the climate system: A scientific assessment, Journal of Geophysical Research-Atmospheres, 118, 5380-5552, 10.1002/jgrd.50171, 2013.
 - Bondy, A., Bonanno, D., Moffet, R., Wang, B., Laskin, A., and Ault, A.: The diverse chemical mixing state of aerosol particles in the southeastern United States, Atmospheric Chemistry and Physics, 18, 12595-12612, 10.5194/acp-18-12595-2018, 2018.
 - Boylan, J. W., and Russell, A. G.: PM and light extinction model performance metrics, goals, and criteria for threedimensional air quality models, Atmospheric Environment, 40, 4946-4959, 10.1016/j.atmosenv.2005.09.087, 2006.





- Castro, L., Pio, C., Harrison, R., and Smith, D.: Carbonaceous Aerosol in Urban and Rural European Atmospheres:
 Estimation of Secondary Organic Carbon Concentrations, Atmospheric Environment, 33, 2771-2781, 10.1016/s1352-2310(98)00331-8, 1999.
 - Chen, X., Wang, Z., Li, J., Chen, H., Hu, M., Yang, W., Wang, Z., Ge, B., and Wang, D.: Explaining the spatiotemporal variation of fine particle number concentrations over Beijing and surrounding areas in an air quality model with aerosol microphysics, Environmental pollution, 231, 1302-1313, 10.1016/j.envpol.2017.08.103, 2017a.
- 440 Chen, X. S., Wang, Z. F., Li, J., and Yu, F. Q.: Development of a Regional Chemical Transport Model with Size-Resolved Aerosol Microphysics and Its Application on Aerosol Number Concentration Simulation over China, Sola, 10, 83-87, 10.2151/sola.2014-017, 2014.
 - Chen, X. S., Wang, Z. F., Yu, F. Q., Pan, X. L., Li, J., Ge, B. Z., Wang, Z., Hu, M., Yang, W. Y., and Chen, H. S.: Estimation of atmospheric aging time of black carbon particles in the polluted atmosphere over central-eastern China
- using microphysical process analysis in regional chemical transport model, Atmospheric Environment, 163, 44-56, 10.1016/j.atmosenv.2017.05.016, 2017b.
 - Chen, X., Yu, F., Yang, W., Sun, Y., Chen, H., Du, W., Zhao, J., Wei, Y., Wei, L., Du, H., Wang, Z., Wu, Q., Li, J., An, J., and Wang, Z.: Global–regional nested simulation of particle number concentration by combing microphysical processes with an evolving organic aerosol module, Atmos. Chem. Phys., 21, 9343-9366, 10.5194/acp-21-9343-2021, 2021.
- 450 Cheng, Y. F., Su, H., Rose, D., Gunthe, S. S., Berghof, M., Wehner, B., Achtert, P., Nowak, A., Takegawa, N., Kondo, Y., Shiraiwa, M., Gong, Y. G., Shao, M., Hu, M., Zhu, T., Zhang, Y. H., Carmichael, G. R., Wiedensohler, A., Andreae, M. O., and Poschl, U.: Size-resolved measurement of the mixing state of soot in the megacity Beijing, China: diurnal cycle, aging and parameterization, Atmospheric Chemistry and Physics, 12, 4477-4491, 10.5194/acp-12-4477-2012, 2012.
- Cui, X. J., Wang, X. F., Yang, L. X., Chen, B., Chen, J. M., Andersson, A., and Gustafsson, O.: Radiative absorption
 enhancement from coatings on black carbon aerosols, Science of the Total Environment, 551, 51-56, 10.1016/j.scitotenv.2016.02.026, 2016.
 - Curci, G., Hogrefe, C., Bianconi, R., Im, U., Balzarini, A., Baró, R., Brunner, D., Forkel, R., Giordano, L., Hirtl, M., Honzak, L., Jiménez-Guerrero, P., Knote, C., Langer, M., Makar, P. A., Pirovano, G., Pérez, J. L., San José, R., Syrakov, D., Tuccella, P., Werhahn, J., Wolke, R., Žabkar, R., Zhang, J., and Galmarini, S.: Uncertainties of simulated aerosol
- optical properties induced by assumptions on aerosol physical and chemical properties: An AQMEII-2 perspective,
 Atmospheric Environment, 115, 541-552, 10.1016/j.atmosenv.2014.09.009, 2015.
 - Curci, G., Alyuz, U., Baro, R., Bianconi, R., Bieser, J., Christensen, J. H., Colette, A., Farrow, A., Francis, X., Jimenez-Guerrero, P., Im, U., Liu, P., Manders, A., Palacios-Pena, L., Prank, M., Pozzoli, L., Sokhi, R., Solazzo, E., Tuccella, P., Unal, A., Vivanco, M. G., Hogrefe, C., and Galmarini, S.: Modelling black carbon absorption of solar radiation: combining external and internal mixing assumptions, Atmos Chem Phys, 19, 181-204, 10.5194/acp-19-181-2019, 2019.



470



- 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, 10.5194/gmd-10-4057-2017, 2017.
- Dentener, F., Kinne, S., Bond, T., Boucher, O., Cofala, J., Generoso, S., Ginoux, P., Gong, S., Hoelzemann, J. J., Ito, A., Marelli, L., Penner, J. E., Putaud, J. P., Textor, C., Schulz, M., van der Werf, G. R., and Wilson, J.: Emissions of
- primary aerosol and precursor gases in the years 2000 and 1750 prescribed data-sets for AeroCom, Atmospheric Chemistry and Physics, 6, 4321-4344, DOI 10.5194/acp-6-4321-2006, 2006.
 - Du, H., Li, J., Chen, X., Wang, Z., Sun, Y., Fu, P., Li, J., Gao, J., and Wei, Y.: Modeling of aerosol property evolution during winter haze episodes over a megacity cluster in northern China: roles of regional transport and heterogeneous reactions of SO2, Atmospheric Chemistry and Physics, 19, 9351-9370, 10.5194/acp-19-9351-2019, 2019.
 - Emery, C., Liu, Z., Russell, A., Odman, M., Yarwood, G., and Kumar, N.: Recommendations on Statistics and Benchmarks to Assess Photochemical Model Performance, Journal of the Air & Waste Management Association (1995), 67, 10.1080/10962247.2016.1265027, 2016.
- Fierce, L., Onasch, T. B., Cappa, C. D., Mazzoleni, C., China, S., Bhandari, J., Davidovits, P., Fischer, D. A., Helgestad, T.,
- 480 Lambe, A. T., Sedlacek, A. J., 3rd, Smith, G. D., and Wolff, L.: Radiative absorption enhancements by black carbon controlled by particle-to-particle heterogeneity in composition, Proceedings of the National Academy of Sciences of the United States of America, 117, 5196-5203, 10.1073/pnas.1919723117, 2020.
 - Fuller, K. A., Malm, W. C., and Kreidenweis, S. M.: Effects of mixing on extinction by carbonaceous particles, Journal of Geophysical Research-Atmospheres, 104, 15941-15954, Doi 10.1029/1998jd100069, 1999.
- 485 Gao, M., Han, Z., Tao, Z., Li, J., Kang, J.-E., Huang, K., Dong, X., Zhuang, B., Li, S., Ge, B., Wu, Q., Lee, H.-J., Kim, C.-H., Fu, J. S., Wang, T., Chin, M., Li, M., Woo, J.-H., Zhang, Q., Cheng, Y., Wang, Z., and Carmichael, G. R.: 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, Atmospheric Chemistry and Physics, 20, 1147-1161, 10.5194/acp-20-1147-2020, 2020.
- 490 Hess, M., Koepke, P., and Schult, I.: Optical properties of aerosols and clouds: The software package OPAC, Bulletin of the American Meteorological Society, 79, 831-844, Doi 10.1175/1520-0477(1998)079<0831:Opoaac>2.0.Co;2, 1998.
 - Hu, K., Liu, D., Tian, P., Wu, Y., Li, S., Zhao, D., Li, R., Sheng, J., Huang, M., Ding, D., Liu, Q., Jiang, X., Li, Q., and Tao, J.: Identifying the Fraction of Core–Shell Black Carbon Particles in a Complex Mixture to Constrain the Absorption Enhancement by Coatings, Environ Sci Tech Let, 9, 10.1021/acs.estlett.2c00060, 2022.
- 495 Huang, X.-F., Peng, Y., Wei, J., Peng, J., Lin, X.-Y., Tang, M.-X., Cheng, Y., Men, Z., Fang, T., Zhang, J., He, L.-Y., Cao, L.-M., Liu, C., Zhang, C., Mao, H., Seinfeld, J. H., and Wang, Y.: Microphysical complexity of black carbon particles restricts their warming potential, One Earth, 10.1016/j.oneear.2023.12.004, 2023.



500



- IPCC: Climate Change 2021: The Physical Science Basis. Contribution of Working Group I to the Sixth Assessment Report of the Intergovernmental Panel on Climate Change, Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2021.
- Jacobson, M. Z.: Comment on "Radiative absorption enhancements due to the mixing state of atmospheric black carbon", Science, 339, 393, 10.1126/science.1229920, 2013.
- Kang, Z., Ma, P., Quan, J., Liao, Z., Pan, Y., Liu, H., Pan, X., Dou, Y., Zhao, X., Cheng, Z., Wang, Q., Yuan, T., and Jia, X.: Mixing state of refractory black carbon in the residual layer over megacity, Atmospheric Environment, 295, 119558, 10.1016/j.atmosenv.2022.119558, 2023.
- Li, J., Chen, X., Wang, Z., Du, H., Yang, W., Sun, Y., Hu, B., Li, J., Wang, W., Wang, T., Fu, P., and Huang, H.: Radiative and heterogeneous chemical effects of aerosols on ozone and inorganic aerosols over East Asia, The Science of the total environment, 622-623, 1327-1342, 10.1016/j.scitotenv.2017.12.041, 2018.
- Li, J., Han, Z., Wu, Y., Xiong, Z., Xia, X., Li, J., Liang, L., and Zhang, R.: Aerosol radiative effects and feedbacks on
 boundary layer meteorology and PM2.5 chemical components during winter haze events over the Beijing-Tianjin-Hebei
 region, Atmospheric Chemistry and Physics, 20, 8659-8690, 10.5194/acp-20-8659-2020, 2020.
 - Li, M., Liu, H., Geng, G. N., Hong, C. P., Liu, F., Song, Y., Tong, D., Zheng, B., Cui, H. Y., Man, H. Y., Zhang, Q., and He, K. B.: Anthropogenic emission inventories in China: a review, Natl Sci Rev, 4, 834-866, 10.1093/nsr/nwx150, 2017.
 - Li, W., Shao, L., Zhang, D., Ro, C.-U., Hu, M., Bi, X., Geng, H., Matsuki, A., Niu, H., and Chen, J.: A review of single
- 515 aerosol particle studies in the atmosphere of East Asia: morphology, mixing state, source, and heterogeneous reactions, Journal of Cleaner Production, 112, 1330-1349, 10.1016/j.jclepro2015.04.050, 2016.
 - Li, W., Riemer, N., Xu, L., Wang, Y., Adachi, K., Shi, Z., Zhang, D., Zheng, Z., and Laskin, A.: Microphysical properties of atmospheric soot and organic particles: measurements, modeling, and impacts, npj Climate and Atmospheric Science, 7, 65, 10.1038/s41612-024-00610-8, 2024.
- 520 Liu, C., Chung, C., Yin, Y., and Schnaiter, M.: The absorption Ångström exponent of black carbon: From numerical aspects, Atmospheric Chemistry and Physics, 18, 6259-6273, 10.5194/acp-18-6259-2018, 2018.
 - Liu, D. T., Whitehead, J., Alfarra, M. R., Reyes-Villegas, E., Spracklen, D. V., Reddington, C. L., Kong, S. F., Williams, P. I., Ting, Y. C., 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, 184-U132, 10.1038/Ngao2901.2017.
- 525 10.1038/Ngeo2901, 2017.
 - Liu, D. T., He, C. L., Schwarz, J. P., and Wang, X.: Lifecycle of light-absorbing carbonaceous aerosols in the atmosphere, Npj Clim Atmos Sci, 3, 40, 10.1038/s41612-020-00145-8, 2020.
 - Liu, X., Ma, P.-L., Wang, H., Tilmes, S., Singh, B., Easter, R., Ghan, S., and Rasch, P.: Description and evaluation of a new four-mode version of the Modal Aerosol Module (MAM4) within version 5.3 of the Community Atmosphere Model,
- 530 Geoscientific Model Development, 9, 505-522, 10.5194/gmd-9-505-2016, 2016.



535



- Mann, G., Carslaw, K., Spracklen, D., Ridley, D., Manktelow, P., Chipperfield, M., Pickering, S., and Johnson, C.: Description and evaluation of GLOMAP-mode aerosol microphysics model for the UKCA composition-climate model, Geoscientific Model Development Discussions, 3, 10.5194/gmd-3-519-2010, 2010.
- Matsui, H.: Black carbon simulations using a size- and mixing-state-resolved three-dimensional model: 1. Radiative effects and their uncertainties, Journal of Geophysical Research-Atmospheres, 121, 1793-1807, 10.1002/2015jd023998, 2016.
- Matsui, H., Koike, M., Kondo, Y., Fast, J., and Takigawa, M.: Development of an aerosol microphysical module: Aerosol Two-dimensional bin module for formation and Aging Simulation (ATRAS), Atmospheric Chemistry and Physics, 14, 10.5194/acp-14-10315-2014, 2014.
- Mie, G.: Beiträge zur Optik Trüber Medien, Speziell Kolloidaler Metallösungen, Ann. Phys., 25, 377, 10.1002/andp.19083300302, 1908.
 - Mishchenko, M. I., Dlugach, J. M., Yanovitskij, E. G., and Zakharova, N. T.: Bidirectional reflectance of flat, optically thick particulate layers: an efficient radiative transfer solution and applications to snow and soil surfaces, Journal of Quantitative Spectroscopy & Radiative Transfer, 63, 409-432, Doi 10.1016/S0022-4073(99)00028-X, 1999.
 - Peng, J., Hu, M., Guo, S., Du, Z., Zheng, J., Shang, D., Levy Zamora, M., Zeng, L., Shao, M., Wu, Y. S., Zheng, J., Wang,
- 545 Y., Glen, C. R., Collins, D. R., Molina, M. J., and Zhang, R.: Markedly enhanced absorption and direct radiative forcing of black carbon under polluted urban environments, Proceedings of the National Academy of Sciences of the United States of America, 113, 4266-4271, 10.1073/pnas.1602310113, 2016.
 - Obiso, V. and Jorba, O.: Aerosol-radiation interaction in atmospheric models: Idealized sensitivity study of simulated shortwave direct radiative effects to particle microphysical properties, Journal of Aerosol Science, 115, 46-61, 10.1016/j.jaerosci.2017.10.004, 2018.
 - Paasonen, P., Kupiainen, K., Klimont, Z., Visschedijk, A., Denier van der Gon, H. A. C., and Amann, M.: Continental anthropogenic primary particle number emissions, Atmos. Chem. Phys., 16, 6823-6840, 10.5194/acp-16-6823-2016, 2016.
 - Riemer, N., P. Ault, A., West, M., L. Craig, R., and H. Curtis, J.: Aerosol Mixing State: Measurements, Modeling, and Impacts, Reviews of Geophysics, 10.1029/2018RG000615, 2019.
 - Riemer, N., West, M., Zaveri, R. A., and Easter, R. C.: Simulating the evolution of soot mixing state with a particle-resolved aerosol model, Journal of Geophysical Research-Atmospheres, 114, 10.1029/2008jd011073, 2009.

Ran, L., Deng, Z. Z., Wang, P. C., and Xia, X. A.: Black carbon and wavelength-dependent aerosol absorption in the North China Plain based on two-year aethalometer measurements, Atmospheric Environment, 142, 132-144, 10.1016/j.atmosenv.2016.07.014, 2016.

560

550

555

Shen, W. X., Wang, M. H., Riemer, N., Zheng, Z. H., Liu, Y. W., and Dong, X. Y.: Improving BC Mixing State and CCN Activity Representation With Machine Learning in the Community Atmosphere Model Version 6 (CAM6), Journal of Advances in Modeling Earth Systems, 16, 10.1029/2023MS003889, 2024.



565

595



Stevens, R., and Dastoor, A.: A Review of the Representation of Aerosol Mixing State in Atmospheric Models, Atmosphere, 10, 168, 10.3390/atmos10040168, 2019.

- Sun, J., Xie, C., Xu, W., Chen, C., Ma, N., Xu, W., Lei, L., Li, Z., He, Y., Qiu, Y., Wang, Q., Pan, X., Su, H., Cheng, Y., Wu, C., Fu, P., Wang, Z., and Sun, Y.: Light absorption of black carbon and brown carbon in winter in North China Plain: comparisons between urban and rural sites, The Science of the total environment, 770, 144821, 10.1016/j.scitotenv.2020.144821, 2021.
- 570 Toon, O. B., and Ackerman, T. P.: Algorithms for the calculation of scattering by stratified spheres, Appl Opt, 20, 3657-3660, 10.1364/AO.20.003657, 1981.
 - Tuccella, P., Curci, G., Pitari, G., Lee, S., and Jo, D. S.: Direct Radiative Effect of Absorbing Aerosols: Sensitivity to Mixing State, Brown Carbon, and Soil Dust Refractive Index and Shape, Journal of Geophysical Research-Atmospheres, 125, 10.1029/2019JD030967, 2020.
- 575 Wang, J., Liu, D., Ge, X., Wu, Y., Shen, F., Chen, M., Zhao, J., Xie, C., Wang, Q., Xu, W., Zhang, J., Hu, J., Allan, J., Joshi, R., Fu, P., Coe, H., and Sun, Y.: Characterization of black carbon-containing fine particles in Beijing during wintertime, Atmos. Chem. Phys., 19, 447-458, 10.5194/acp-19-447-2019, 2019.
- Wang, Y., Li, W., Huang, J., Liu, L., Pang, Y., He, C., Liu, F., Liu, D., Bi, L., Zhang, X., and Shi, Z.: Nonlinear Enhancement of Radiative Absorption by Black Carbon in Response to Particle Mixing Structure, Geophysical Research Letters, 48, 10.1029/2021gl096437, 2021a.
 - Wang, Y., Hu, R., Wang, Q., Li, Z., Cribb, M., Sun, Y., Song, X., Shang, Y., Wu, Y., Huang, X., and Wang, Y.: Different effects of anthropogenic emissions and aging processes on the mixing state of soot particles in the nucleation and accumulation modes, Atmos. Chem. Phys., 22, 14133-14146, 10.5194/acp-22-14133-2022, 2022.
- Wang, Y. Y., Pang, Y. E., Huang, J., Bi, L., Che, H. Z., Zhang, X. Y., and Li, W. J.: Constructing Shapes and Mixing
 Structures of Black Carbon Particles With Applications to Optical Calculations, Journal of Geophysical Research-Atmospheres, 126, e2021JD034620, 10.1029/2021JD034620, 2021b.
 - Wang, Z. F., Maeda, T., Hayashi, M., Hsiao, L. F., and Liu, K. Y.: A nested air quality prediction modeling system for urban and regional scales: Application for high-ozone episode in Taiwan, Water Air and Soil Pollution, 130, 391-396, Doi 10.1023/A:1013833217916, 2001.
- 590 Wu, Y., Zhang, R., Tian, P., Tao, J., Hsu, S. C., Yan, P., Wang, Q., Cao, J., Zhang, X., and Xia, X.: Effect of ambient humidity on the light absorption amplification of black carbon in Beijing during January 2013, Atmospheric Environment, 124, 217-223, https://doi.org/10.1016/j.atmosenv.2015.04.041, 2016.
 - Wu, Y., Xia, Y., Zhou, C., Tian, P., Tao, J., Huang, R.-J., Liu, D., Wang, X., Xia, X., Han, Z., and Zhang, R.: Effect of source variation on the size and mixing state of black carbon aerosol in urban Beijing from 2013 to 2019: Implication on light absorption, Environmental pollution, 270, 116089, https://doi.org/10.1016/j.envpol.2020.116089, 2021.
 - Xie, C. H., Xu, W. Q., Wang, J. F., Wang, Q. Q., Liu, D. T., Tang, G. Q., Chen, P., Du, W., Zhao, J., Zhang, Y. J., Zhou, W., Han, T. T., Bian, Q. Y., Li, J., Fu, P. Q., Wang, Z. F., Ge, X. L., Allan, J., Coe, H., and Sun, Y. L.: Vertical



620



characterization of aerosol optical properties and brown carbon in winter in urban Beijing, China, Atmospheric Chemistry and Physics, 19, 165-179, 10.5194/acp-19-165-2019, 2019.

- Kie, X., Hu, J., Qin, M., Guo, S., Hu, M., Ji, D., Wang, H., Lou, S., Huang, C., Liu, C., Zhang, H., Ying, Q., Liao, H., and Zhang, Y.: Evolution of atmospheric age of particles and its implications for the formation of a severe haze event in eastern China, Atmospheric Chemistry and Physics, 23, 10563-10578, 10.5194/acp-23-10563-2023, 2023.
 - Xu, P., Chen, Y., and Ye, X.: Haze, air pollution, and health in China, Lancet, 382, 2067, 10.1016/S0140-6736(13)62693-8, 2013.
- 605 Yang, W., Li, J., Wang, W., Li, J., Ge, M.-F., Sun, Y., Chen, X., Ge, B., Tong, S., Wang, Q., and Wang, Z.: Investigating secondary organic aerosol formation pathways in China during 2014, Atmospheric Environment, 213, 133–147, 10.1016/j.atmosenv.2019.05.057, 2019.
 - Yao, Y., Curtis, J. H., Ching, J., Zheng, Z., and Riemer, N.: Quantifying the effects of mixing state on aerosol optical properties, Atmos. Chem. Phys., 22, 9265-9282, 10.5194/acp-22-9265-2022, 2022.
- 610 Yu, F., and Luo, G.: Simulation of particle size distribution with a global aerosol model: contribution of nucleation to aerosol and CCN number concentrations, Atmos. Chem. Phys., 9, 7691–7710, 2009.
 - Yu, F., Luo, G., and Ma, X.: Regional and global modeling of aerosol optical properties with a size, composition, and mixing state resolved particle microphysics model, Atmospheric Chemistry and Physics, 12, 5719-5736, 10.5194/acp-12-5719-2012, 2012.
- 615 Yu, P., Toon, O., Bardeen, C., Mills, M., Fan, T., English, J., and Neely, R.: Evaluations of Tropospheric Aerosol Properties Simulated by the Community Earth System Model with a Sectional Aerosol Microphysics Scheme, Journal of Advances in Modeling Earth Systems, 7, 10.1002/2014MS000421, 2015.
 - Zaveri, R. A., Barnard, J. C., Easter, R. C., Riemer, N., and West, M.: Particle-resolved simulation of aerosol size, composition, mixing state, and the associated optical and cloud condensation nuclei activation properties in an evolving urban plume, Journal of Geophysical Research-Atmospheres, 115, 10.1029/2009jd013616, 2010.
 - Zhang, H., Guo, H., Hu, J., Ying, Q., and Kleeman, M. J.: Modeling atmospheric age distribution of elemental carbon using a regional age-resolved particle representation framework, Environmental science & technology, 2018.
 - Zhang, Y. T., Liu, H., Lei, S. D., Xu, W. Y., Tian, Y., Yao, W. J., Liu, X. Y., Liao, Q., Li, J., Chen, C., Sun, Y. L., Fu, P. Q., Xin, J. Y., Cao, J. J., Pan, X. L., and Wang, Z. F.: Mixing state of refractory black carbon in fog and haze at rural sites
- in winter on the North China Plain, Atmospheric Chemistry and Physics, 21, 17631-17648, 10.5194/acp-21-17631-2021, 2021.
 - Zhao, G., Shen, C., and Zhao, C.: Technical note: Mismeasurement of the core-shell structure of black carbon-containing ambient aerosols by SP2 measurements, Atmospheric Environment, 243, 117885, https://doi.org/10.1016/j.atmosenv.2020.117885, 2020.





- 630 Zhao, G., Tan, T., Hu, S., Du, Z., Shang, D., Wu, Z., Guo, S., Zheng, J., Zhu, W., Li, M., Zeng, L., and Hu, M.: Mixing state of black carbon at different atmospheres in north and southwest China, Atmospheric Chemistry and Physics, 22, 10861-10873, 10.5194/acp-22-10861-2022, 2022.
 - Zhao, G., Tan, T., Zhu, Y., Hu, M., and Zhao, C.: Method to quantify black carbon aerosol light absorption enhancement with a mixing state index, Atmos. Chem. Phys., 21, 18055-18063, 10.5194/acp-21-18055-2021, 2021.
- 635 Zhao, P., Dong, F., Yang, Y., He, D., Zhao, X., Zhang, W., Yao, Q., and Liu, H.: Characteristics of carbonaceous aerosol in the region of Beijing, Tianjin, and Hebei, China, Atmospheric Environment, 71, 389-398, https://doi.org/10.1016/j.atmosenv.2013.02.010, 2013.
 - Zheng, H., Kong, S., Chen, N., and Wu, C.: Secondary inorganic aerosol dominated the light absorption enhancement of black carbon aerosol in Wuhan, Central China, Atmospheric Environment, 119288, https://doi.org/10.1016/j.atmosenv.2022.119288, 2022.
 - Zhou, C. H., Gong, S., Zhang, X. Y., Liu, H. L., Xue, M., Cao, G. L., An, X. Q., Che, H. Z., Zhang, Y. M., and Niu, T.: Towards the improvements of simulating the chemical and optical properties of Chinese aerosols using an online coupled model – CUACE/Aero, Tellus Series B-chemical Physical Meteorology, 64, 91-102, 2012.