Global-regional nested simulation of particle number concentration by combing microphysical processes with an evolving organic aerosol module

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Abstract

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Aerosol microphysical processes are essential for the next generation of global and regional climate and air quality models to determine particle size distribution. The contribution of organic aerosol (OA) to particle formation, mass, and number concentration is one of the major uncertainties in current models. A new global-regional nested aerosol model was developed to simulate detailed microphysical processes. The model combines an advanced particle microphysics (APM) module and a volatility basis-set (VBS) OA module to calculate the kinetic

condensation of low-volatility organic compounds and equilibrium partitioning of semi-volatile organic compounds in a 3-dimensional (3-D) framework using global-regional nested domain. In addition to the condensation of sulfuric acid, the equilibrium partitioning of nitrate and ammonium, and the coagulation process of particles, the microphysical processes of the OAs are realistically represented in our new model. The model uses high-resolution size-bins to calculate the size distribution of new particles formed through nucleation and subsequent growth. The multi-scale nesting enables the model to perform high-resolution simulations of the particle formation processes in the urban atmosphere in the background of regional and global environments. By using the nested domains, the model reasonably reproduced the OA components obtained from the analysis of aerosol mass spectrometry measurements through positive matrix factorization and the particle number size distribution in the megacity of Beijing during a period of approximately a month. Anthropogenic organic species accounted for 67 % of the OA of secondary particles formed by nucleation and subsequent growth, which is considerably larger than that of biogenic OA. On the global scale, the model well predicted the particle number concentration in various environments. The microphysical module combined with the VBS simulated the universal distribution of organic components among the different aerosol populations. The model results strongly suggest the importance of anthropogenic organic species in aerosol particle formation and growth at polluted urban sites and over the whole globe under the influence of anthropogenic sources.

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Key words: IAP-AACM+APM, VBS, organic aerosol, particle number concentration **1Introduction**

The increased concentrations of atmospheric aerosol particles caused by anthropogenic activities have become an important scientific issue due to their substantial climate forcing and health effects (Twomey, 1977; Albrecht, 1989; Charlson et al., 1992; Donaldson et al., 2002; Tsigaridis et al., 2006; IPCC, 2013) on global and regional scales. These effects depend on aerosol size, composition, and mixing state. The direct influence of aerosols on climate is their scattering of solar radiations largely determined by the aforementioned key properties of aerosols (IPCC,

2013). The indirect effects of aerosols involve their ability in affecting cloud microphysical properties and precipitation processes by serving as cloud condensation nuclei (CCN), which are highly dependent on CCN number concentrations (Dusek et al., 2006). Ultrafine particles, despite having a lower mass concentration, have larger health effect because of their ability to easily penetrate the body and their higher number concentrations (Delfino et al., 2005; Kumar et al., 2014). Therefore, it is crucial to gain deep insight into the life cycle of aerosol particles and quantify their sources not only in mass concentration but also in their number concentration.

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There are two sources of atmospheric aerosols: direct emissions from primary sources and secondary formation processes (Seinfeld and Pandis, 2006). Mineral dust particles over desert regions and sea salt particles over oceans are the two major natural sources contributing to particle mass and number concentration regionally (Textor et al., 2006). Anthropogenic activities (e.g., fossil fuel combustion and biomass burning) can directly emit particles and they are the most significant contributors to the aerosols since the industrial revolution (IPCC, 2013). The physical and chemical properties of these primarily emitted particles can be modified by condensation, coagulation, and chemical reactions in the atmosphere (Seinfeld and Pandis, 2006). In addition, new particle formation (NPF) has been reported to be an important contributor to aerosol particles in global various environments (Holmes, 2007; Yu et al., 2008; Yu and Luo, 2009; Kulmala et al., 2013). Field observation studies have also demonstrated that NPF can significantly increase CCN number concentrations (Kuang et al., 2009; Wiedensohler et al., 2009; Yue et al., 2011). Thus, it is necessary to reasonably represent primary emission, their microphysical aging, and new particle formation process in 3-D models.

During the past two decades, numerous models have incorporated microphysical module to describe particle formation processes (e.g., Binkowski and Shankar, 1995; Jacobson, 1997; Stier et al., 2005; Bergman et al., 2012). However, large uncertainties still exist due to the unclear complication of processes and the relevant mechanisms. Intercomparison and evaluation of global aerosol models indicate that constraint of size-resolved primary emission and improved understanding of secondary formations

are required to improve the ability of model to simulate particle number size distribution (PNSD) (Mann et al., 2014). Spracklen et al. (2005) found that the size distribution assumption has a large impact on particle number concentrations in the boundary layer. A comparison between the simulations and single particle soot photometer measurements suggested that the model that was employed had a large bias in simulating the number size distribution of black carbon particles (Reddington et al., 2013). Considerable improvements in the simulation of the particle number concentration and aerosol optical properties were achieved by using an optimized size distribution of primary particles in polluted atmosphere over areas with large emissions (Zhou et al., 2012, 2018). However, much work remains to reduce the primary emissions, uncertainty associated with especially over primary particles-dominated regions (e.g., China) in terms of particle number concentration.

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The main source of uncertainty in simulating NPF at regional and global scales can be attributed to the nucleation mechanism and particle growth rates unexplained. Although sulfuric acid has been identified as a major component and plays a central role in nucleation (Yu and Turco, 2001; Boy et al., 2005; Kirkby et al., 2011), alone it could not explain the NPF rates (Wang et al., 2013; Kulmala et al., 2013). Recent studies have revealed that certain organic vapors are involved in particle nucleation (Metzger et al., 2010; Zhang et al., 2012; Yao et al., 2018) and contribute considerably to particle growth (Kulmala and Kerminen, 2008; Tröstl et al., 2016). Thus, it is no doubt that a reasonable representation of OA is crucial for aerosol models to realistically simulate NPF and particle growth. However, it is still an open question which organic species are possibly involved in new particle formation process. Even the chemical composition and the sources of OA are still uncertain because they contain large number of compounds (Goldstein and Galbally, 2007). To date, OA is still the least understood one among the components of aerosols (Kanakidou et al., 2005; Hallquist et al., 2009). Clearly, OA representation is the major uncertainty contributing to the large knowledge gap in elucidating particle formation processes.

In recent years, much progress has been achieved in simulating the formation of OA and secondary organic aerosol (SOA). The two product (2P) model recommended

by Odum et al. (1996) has been widely used in 3-D models to describe the SOA formation process empirically. The volatility basis-set (VBS) approach was recently developed to represent the oxidation of primary OA (POA) and SOA and the partitioning of OA in different volatilities between gas phase and aerosol phase (Donahue et al., 2006). Many regional models have used VBS to simulate OA and SOA (Shrivastava et al., 2008; Fountoukis et al., 2011; Ahmadov et al., 2012; Zhao et al., 2016; Han et al., 2016). However, the application of VBS in global models is limited because of the large number of tracers required and the uncertainty of the involved parameters (Farina et al., 2010; Hodzic et al., 2016). There are even fewer applications of this unified framework in 3-D global aerosol models to calculate the processes of particle formation. Among the second phase AeroCom aerosol microphysical models, the simplified parameterization and the 2P method are the most common schemes used to represent SOA (Mann et al., 2014). Recently, some models have incorporated VBS in their microphysical module to simulate the aerosol microphysical formation process. Patoulias et al. (2015) developed a new aerosol dynamics model with VBS and explored the contribution of SOA with various volatilities to particle growth in different stages; however, the 3-D modeling was not presented. By assuming equilibrium partitioning for all volatility bins, Gao et al. (2017) implemented VBS in an aerosol microphysics model and examined the effect of semi-volatile SOA (SV-SOA) on the composition, growth, and mixing state of particles. Their box model simulation suggested that the volatility of organic compounds simulate rather different mixing states from those simulated by the coagulation process alone in the scheme treating the primary emission of organics as nonvolatile. Matsui (2017) represented aerosol size distribution with a two-dimensional (2-D) sectional method in a global aerosol model coupled with the VBS scheme, but the size-bin resolution is insufficient to accurately resolve the growth of new particles.

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To our knowledge, there are few 3-D modeling studies using VBS to account for both (1) the kinetic condensation of low-volatility organics and re-evaporation of semi-volatile organics and (2) the size-resolved kinetics of the mass transfer for new

particles. In addition, particle formation in the polluted atmosphere was not well understood (Kulmala et al., 2016; Wang et al., 2017; Chu et al., 2019). Over urban areas in northern China, observation and modeling studies have indicated that anthropogenic SOA contributes a larger fraction to OA than that of biogenic one and play an significant role in particle formation (Yang et al., 2016; Guo et al., 2020; Han et al., 2016; Lin et al., 2016). Simultaneously calculating both anthropogenic and biogenic SOA in microphysical models with a high resolution is crucial to resolve the particle formation processes over urban areas. Furthermore, the previous studies focusing on the sensitivity of particle number concentration to primary emission were based on models without considering the detailed microphysics of organic species (e.g., Spracklen et al., 2006; Chang et al., 2009; Chen et al., 2018; Zhou et al., 2018). Therefore, it is urgently needed to establish a 3-D modeling framework of VBS with an aerosol microphysics module with high size-bin resolution to simulate the particle number size distribution and explore the uncertainties associated with the treatment of primary emission.

In our previous work, a regional model with detailed microphysical processes has been developed to improve the simulation of NPF during summer in Beijing (Chen et al., 2019). In this study, we extend our work to the global scale and doing so to establish a new aerosol model by coupling a VBS OA scheme with a particle microphysics module in a global-regional nested model. The model performance was evaluated against the measurements at a tower and a dataset collected from published papers. In addition, the model's sensitivity to the size distribution of primary emission and volatility distribution of POA were explored to elucidate and quantify the uncertainties associated. This new modeling framework can provide a useful tool to simulate the aerosol microphysical process in both global and regional scales. The description of the model and its development method are introduced in Sect.2. The experiments setup and model input are detailed in Sect.3. The observed data used for evaluating model performance are described in Sect.4. The model results and simulation analysis are presented in Sect.5. Finally, the conclusions and discussions are summarized in Sect.6.

2 Model description

2.1 Host model

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The host model employed in this study is the Atmospheric Aerosol and Chemistry Model developed by the Institute of Atmospheric Physics, Chinese Academy of Sciences (IAP-AACM). The IAP-AACM is a 3-D atmospheric chemical transport model treating chemical and physical processes for gases and aerosols in global and regional scales using multi-scale domain-nesting technique (Wang et al., 2001; Li et al., 2012; Chen et al., 2015). The meteorological parameters input to the IAP-AACM were simulated by the global version of the Weather Research and Forecasting (WRF) model (Skamarock et al., 2008; Zhang et al., 2012). The IAP-AACM model has been successfully used to explore mercury transport (Chen et al., 2015) and simulate the global and regional distribution of gaseous pollutants and aerosol components (Du et al., 2019; Wei et al., 2019). The calculation of some modules in the model has also been optimized recently (Wang et al., 2017, 2019). The model calculates 3-D advection (Walcek, 1998), turbulent diffusion (Byun and Dennis, 1995), gas phase chemical reactions (Zaveri and Peters, 1999), dry deposition at the surface (Zhang et al., 2003), aqueous reactions in the cloud and wet scavenging (Stockwell et al., 1990), and heterogeneous chemical processes (Li et al., 2012). The partition of nitric acid and ammonia into particle phase to form nitrate and ammonium is simulated using a thermodynamic equilibrium model (Nenes et al., 1998). The model calculates the online emission of dimethyl sulfide (Lana et al., 2011), sea salt (Athanasopoulou et al., 2008) and dust (Wang et al., 2000; Luo and Wang, 2006). The simulation results of IAP-AACM have been evaluated against a comprehensive observation dataset and compared with other model results. The model exhibited good performance in reproducing global aerosol components (Wei et al., 2019).

2.2 Advanced particle microphysics module

Advanced particle microphysics (APM) module is an aerosol module that uses the sectional method to represent particle number size distribution. The APM module has been coupled to several 3-D models, such as Goddard Earth Observing System-Chemistry model (Yu and Luo, 2009), Weather Research and

Forecasting-Chemistry model (Luo and Yu, 2011), and the Nested Air Quality Prediction Modeling System (NAQPMS; Chen et al., 2014). In APM, there are two types of aerosol particles: secondary particles (SPs) and primary particles (PPs) with a secondary species coating. The definitions of SPs and PPs in our model differ from those of secondary aerosol and primary aerosol commonly used in the community. SPs indicate they originate from the nucleation and the subsequent growth of newly nucleated particles whereas PPs originate from direct emission. PPs include dust particles, sea salt particles, organic carbon (OC) particles, and black carbon (BC) particles. The APM module has a high size-bin resolution to accurately describe the formation and growth processes of SPs (composed of sulfate, nitrate, ammonium, and organic compounds). SPs are represented by 40 size bins from 0.0012 to 12 µm in dry diameter. Among the PPs, the representation of BC and OC was updated from a modal method in the original version (Yu and Luo, 2009) to a size-bin scheme in the revised version (Chen, 2015). Dust particles in 0.03 to 50 µm are represented by four size bins and sea salt particles in 0.0012 to 12 µm are represented by 20 size bins. SPs are assumed to be internally mixed, and PPs are assumed to consist of a primary core and coating species. SPs and PPs of different categories are externally mixed with each other. In addition to the primary core, the coated species are explicitly simulated in the APM module.

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The basic microphysical processes in the APM module include nucleation, condensation or evaporation, coagulation, and thermodynamic equilibrium partition. The nucleation scheme is ion-mediated nucleation (IMN) (Yu, 2006, 2010), which is physically-based and constrained by laboratory data and has predicted reasonable distributions of global nucleation (Yu et al., 2008). Because of very low saturation vapor pressure, the condensation of H₂SO₄ is explicitly calculated. The semi-volatile inorganic species (nitrate and ammonium) and secondary organic species are simulated through equilibrium partitioning. The bulk mass concentrations of coating species are tracked to reduce the computational cost and the corresponding tracers used are defined as BC sulfate, OC sulfate, sea salt sulfate, and dust sulfate. For coagulation, the APM module not only calculates the self-coagulation of sea salt

particles, BC particles, OC particles, and SPs, but also considers the coagulation scavenging of SPs by four types of PPs. Yu (2011) has further developed the APM module to explicitly calculate the co-condensation of sulfuric acid and low-volatility secondary organic gas (LV-SOG) on SPs and PPs. In the scheme, the production rate of LV-SOG and the semi-volatile OA (SV-OA) input to APM are simulated with the extended 2P SOA formation model. For high calculation efficiency, a pre-calculated look-up table of coagulation kernels is used in the coagulation module. The numerical scheme used is from Jacobson et al. (1994). More details on microphysical processes of APM can be found in Yu and Luo (2009).

2.3 VBS module

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To reproduce the formation and evolution of OA, a 1.5-D VBS approach (Koo et al., 2014) based on 1-D VBS framework but accounting for changes in the oxidation state and volatility of OA in the 2-D VBS space is coupled to the model. Both secondary and primary OAs are distributed in five volatility bins ranging from 10⁻¹ to 10³ μg/m³ in saturation concentration (C*) at 298 K, and the temperature dependence of C* is calculated by the Clausiuse-Clapeyron equation (Sheehan and Bowman, 2001). The compounds distributed in the lowest bin with C^* less than $10^{-1}~\mu g/m^3$ represent the effectively nonvolatile OAs and they are regarded in our model as low-volatility organic compounds that are almost partitioned to the particulate phase. The compounds in the other four bins (i.e., $C^* = 10^0$, 10^1 , 10^2 , and $10^3 \ \mu g/m^3$) are defined as semi-volatile organic compounds that can be partitioned between the gas and particulate phase by using equilibrium assumption (Donahue et al., 2009). To track the oxidation state of OA, four basis sets are used in the scheme: two-basis sets for chemically aged OA from anthropogenic and biogenic sources, and two-basis sets for freshly emitted OA from anthropogenic sources and biomass burning. The molecular properties of primary OA (POA) and SOA in each volatility bin are provided by the parameters calculated by the 2-D volatility scheme (Donahue et al., 2011, 2012).

In this VBS module, gas-phase organic compounds can be aged by extremely reactive hydroxyl (OH) radicals and other oxidants. Volatile organic precursors of

SOA in this study include compounds with terminal olefin carbon bond (R-C=C), and internal olefin carbon bond (R-C=C-R). The associated species in the model are terpenes, isoprene, and aromatics. The aging of POA by OH proceeds at a reaction rate of 4×10^{-11} cm³·molecule⁻¹·s⁻¹ (Robinsonet al., 2007). Considering a single oxidation step would not be able to move the oxidation products of POA into the oxidized OA basis in the volatility bin that is one magnitude lower, the concept of "partial conversion" is used; that is, the oxidation products are a mixture of POA and oxidized POA (OPOA) in the adjacent lower volatility bins (Koo et al., 2014). In addition, the multigenerational oxidation processes of intermediate volatile organic compounds (IVOCs) with OH radicals at a rate constant of $4\times10^{-11} \text{cm}^3 \cdot \text{molecule}^{-1} \cdot \text{s}^{-1}$ are considered in SOA formation. To account for the insufficient reduction in carbon number and volatility decrease of the IVOC product, the SOA mass yields from IVOC are assumed to be lower than that of POA (Yang et al., 2019). Additional IVOCs emissions are assumed to be 4.5, 0.5, and 0.5 times of the POAs emissions for vehicles, other anthropogenic sources, and biomass burning, respectively. IVOC emission is put into the bin of 10⁴ µg/m³ saturation concentration. The VBS module in this study does not consider OA formation through aqueous-phase or heterogeneous reactions although their importance has been suggested in some studies (e.g., Liu et al., 2012; Ervens et al., 2014; Lin et al., 2014). SOA generated from volatile organic compounds (VOCs), IVOCs, and anthropogenic OPOA are assumed to be further oxidized by OH radicals at an aging rate of 2×10^{-11} cm³·molecule⁻¹·s⁻¹ on the base of the work in Koo et al. (2014). The volatilities of multi-generation oxidation products decrease and move down to the adjacent bin with an order of magnitude lower volatility (Donahue et al., 2006). Fragmentation in the 1.5-D VBS module is implicitly considered through reduction in carbon number of the oxidation products. NOx-dependent product mass yields from oxidation of hydrocarbon precursors were determined based on smog chamber data (Murphy and Pandis, 2009; Hildebrandt et al., 2009) The model has 32 pairs of semi-volatile compounds including organic gases (OGs) and the corresponding OAs through equilibrium partitioning. Plus the eight groups of low-volatility OAs, the model has 40 groups of OAs and 32 groups of OGs

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in total. Detailed information on this VBS module can be found in Koo et al. (2014) and Yang et al. (2019).

2.4 Model development

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In our previous work, the VBS module was combined with APM to improve the simulation of NPF process in our regional model (NAQPMS+APM, Chen et al., 2019). Here, we use a similar method to couple the VBS and APM into the global model (IAP-AACM). The newly developed model is named IAP-AACM+APM. In the model, not only the aforementioned basic microphysical processes but also the condensation of LV-SOG and equilibrium partition of SV-OA are calculated following the approaches described in Yu and Luo (2009) and Yu (2011). In addition to the aforementioned tracers of OAs and OGs, a new tracer for LV-SOG is tracked in the IAP-AACM+APM. The sources of sulfuric acid and LV-SOG are photochemical reactions. Their production rates are calculated by the carbon bond mechanism Z (CBM-Z) and VBS module, respectively. The production rate of LV-SOG is equivalent to that of the lowest bin OAs in the VBS module. For simplicity and computing efficiency, the condensation of LV-SOG on SPs of various sizes is calculated along with H₂SO₄ and the low-volatility SOA (LV-SOA) on SPs are merged into one bulk tracer (SP-LV). When necessary (e.g. calculating the condensation growth and coagulation of SPs and the coagulation scavenging of SPs by PPs), SP-LV is redistributed to size-bins according to the surface area of the particles. The condensation of LV-SOG on PPs (i.e., dust, sea salt, BC, and OC particles) is calculated in the same manner as H₂SO₄. The amount of LV-SOA coated on these particles are defined as dust-LV, salt-LV, BC-LV and OC-LV. In this manner, LV-SOAs are distributed approximately proportional to the aerosol surface area. The SV-SOA partitioned to SPs in each bin and the coatings on PPs are assumed to be proportional to the corresponding low-volatility organic aerosol (LV-OA) mass. For OC particles, the coated SV-SOA depends on both the OC-LV and primary organic carbon (POC). The SV-SOA input to APM is the total mass concentration of 32 groups of SV-OAs in the VBS module. The partition of this part of the OA is similar to that of the equilibrium partition theory (Pankow, 1994a,b; Odum et al., 1996). By using the aforementioned treatments, the various microphysical behaviors of OAs with different volatilities are reasonably simulated. The dry deposition at the surface level and wet deposition by precipitation of LV-SOG are modeled using the same scheme as for H₂SO₄. The dry deposition and wet scavenging of the coated LV-OA associated with SPs and PPs are calculated using the same scheme as for the sulfate coated on PPs (Yu, 2011).

The tracers associated with aerosol microphysical processes in the IAP-AACM+APM are listed in Table 1. Compared with the IAP-AACM, 129 newly tracers were added in IAP-AACM+APM; therefore, the computing time of 3-D advection and turbulent diffusion is nearly double that of the IAP-AACM. Among the modules in IAP-AACM+APM, the gas phase reaction module and the microphysical module are the most time-consuming. The newly-developed processes in the IAP-AACM+APM do not add much computing time. The total computing time of the IAP-AACM+APM is less than twice that of the IAP-AACM and is acceptable. The aerosol microphysical module combining VBS with APM in this study can be used in other 3-D models.

3 Model configuration and experiments setting

3.1 Model domain and model inputs

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In this study, we used two nested modeling domains for 1-year simulation in 2010, with the first domain covering the whole globe at 1° resolution and the second domain covering east Asia at 0.33° resolution. The model has 20 vertical layers and the top layer is at 20 km. The simulation from 1 December, 2009, to 31 December, 2010, was used for annual mean analysis and the first 1-month of the simulation was spin-up time and not used in the analysis. In addition, a case study in 2015 using three nested domains, with the third domain of 0.11° resolution, was conducted to evaluate the model performance in simulating OA components and particle number size distribution at a typical urban site. The model domains are demonstrated in Fig.S1.

The IAP-AACM+APM model used the same domain and horizontal grid as for the global WRF. Thus, only vertical interpolation of the meteorological fields from the global WRF model was performed to drive the IAP-AACM+APM. The meteorological fields input to IAP-AACM+APM were updated hourly. The initial and boundary conditions of the global WRF was provided by Final Analysis (FNL) datasets from the National Centers for Environmental Prediction (NCEP) (https://rda.ucar.edu/datasets/ds083.2). The temperature, humidity, wind speed, and pressure in the global WRF were nudged to FNL datasets. For the first domain, a nudging coefficient of 0.0003 for wind, temperature, and water vapor was used in all vertical layers; for the second and third domain, the same nudging scheme was used in vertical layers except those in boundary layer, where nudging was not used. The gridded emission inventory used in the IAP-AACM+APM was an integrated dataset from a publicly available datasets (https://edgar.jrc.ec.europa.eu/htap_v2/index.php) and the multi-resolution emission inventory for China (http://www.meicmodel.org).

3.2 Experiments setting

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As pointed out in Sect.1, the size distribution of primarily emitted particles can directly influence the number concentration of PPs through emission and indirectly change the number concentration of SPs through coagulation-scavenging SPs and competing for condensable gases with SPs. In addition, POA volatility distribution can influence the concentration of OA and the microphysical behavior of OA and thus the particle number concentration. For these reasons, the sensitivity experiments involving size distribution of primarily emitted particles, including BC and POC, and the volatility distribution of POA, were designed to investigate the impacts of these factors on the particle number concentration. One base experiment and four sensitivity experiments were used in our study. Table 2 lists the experiments and their corresponding parameters used in this study. In the base experiment, the volatility distributions of POA from vehicles and biomass burning were based on values from chamber studies (May et al., 2013a,b,c); the factors of other POA emissions were from the estimation of Robinson et al. (2007). The low-volatility POA (LV_POA) experiment and high-volatility POA (HV_POA) experiment used the lower and upper quartiles of the POA volatility distribution factors (May et al., 2013a,b,c; Robinson et al., 2007), respectively. In the OCD0.5 and PPD0.5 experiment, the geometric mean diameter was set as half of the values used in the base experiment for POC, both BC

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4 Observation data

The hourly observation of OA and particle number size distribution (PNSD) in Beijing was used to evaluate the model performance in a typical urban environment. The observation site was located at the Tower Branch of the Institute of Atmospheric Physics (IAP), Chinese Academy of Sciences (CAS) (39°58′N, 116°22′E). The details of the observation site are described in Sun et al. (2015). The observation period was from August 22 to September 30, 2015. OA compositions were measured using a high-resolution aerosol mass spectrometer (HR-AMS, Aerodyne Research Inc.) and an aerosol chemical speciation monitor (ACSM, Aerodyne Research Inc.) at ground level and 260 m, respectively (Zhao et al., 2017). Using positive matrix factorization (PMF) algorithm (Paatero and Tapper, 1994; Paatero, 1997), organic aerosol (OA) were separated into hydrocarbon-like OA (HOA) and oxygenated OA (OOA). A detailed evaluation of PMF results was given in Zhao et al. (2017). PNSD from 15 to 685 nm at ground level and 260 m on the 325 m meteorological tower were measured using two scanning mobility particle sizers. More details on the observation can be found in Du et al. (2017). In the evaluation of PNSD, the observed PNSDs were mapped to the defined size bins of SPs in APM. In the evaluation of particle number concentration in a size range, the number concentrations of all particles in the corresponding size range were summed for comparison. OC concentrations in 2006 from the Interagency Monitoring of Protected Visual Environments (IMPROVE) network (http://vista.cira.colostate.edu/improve) and the China Atmosphere Watch Network (CAWNET) reported by Zhang et al. (2008) were used to compare with the simulated OC of our model. The estimated fractions of OC that is secondary in Zhang et al. (2008) were also used for comparison with the simulations. In addition, a list of surface observations of particle number concentration having at least 1 full year measurements was compiled to verify model performance. Table S1 presents the compiled mean concentrations of condensation nuclei larger than 10 nm (CN10) and the corresponding station information from published papers.

5 Results

425 5.1 PNSD and aerosol components of SPs in Beijing

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The simulated OA concentration was compared with the results of the PMF analysis of the HR-AMS measurements before evaluating the simulated PNSD in Beijing. Here, HOA and OOA components obtained through PMF analysis were been compared with the simulation resultes assuming they are primary and secondary components of OA (i.e., POA and SOA), respectively. Because they were affected by local emission sources (e.g., traffic emission and cooking emission), the observed values of OA were not representative at the ground level. Therefore, only HOA and OOA at 260 m were used for comparison. The third-domain results at 0.11° horizontal resolution, with the other configurations the same as those used in the base experiment, were extracted for the analysis and comparison. First, BC simulations were compared with the observations (Fig.1a), considering that BC is a passive tracer and it is generally co-emitted with POA. The correlation coefficient between the simulated BC and observed BC was 0.70. Because BC is only influenced by emissions, transport and deposition, the agreement between the model and observations in Fig.1a suggests the model represented these processes reasonably well. Figure 1b and 1c reveal the comparison of the simulated and the observed hourly OA components at 260 m. The comparison in Fig.1 highlights the good skill of model in capturing the variation of POA and SOA in our 3-D framework with VBS. The correlation coefficient between the simulated POA and observed HOA was 0.50. Although the measurements at higher level were not susceptible to local emissions, the observed OA concentrations were inevitably influenced by the sources near the measurement site. For example, cooking-related OA, assumed as part of HOA here, have been identified as an important contributor to OA (Zhao et al., 2017). Moreover, nearby traffic emissions would also have a large influence on the observed OA concentrations at the measurement site (Sun et al., 2015). Similar to BC, the temporal variation of POA is mainly influenced by emissions, transport and deposition; thus, the disagreement between the simulated POA and the observed HOA can largely be attributed to the emissions. In additions, PMF analysis has its own uncertainties and deficiencies (Ulbrich et al., 2009). As a result, some observed HOA values were not

reproduced by the IAP-AACM+APM. By contrast, the predicted SOAs and their temporal variation were better agreement with the OOA observation although their concentrations were partially underestimated and some peaks were high. The correlation coefficient between the simulated SOA and observed OOA was 0.52. Overall, our model simulated the POA and SOA concentrations well. The model also reasonably reproduced the concentrations of fine particulate matter in Beijing and its surrounding cities (Fig.S2).

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During the past decades, many field observations have been conducted to study the characteristics of PNSD in Beijing (Wehner et al., 2004; Wu et al., 2007, 2008; Wang et al., 2015). The NAQPMS model has been used to explain the evolution of PNSD in winter in Beijing (Chen et al., 2017). However, 3-D modeling study on these issues are still limited (Kulmala et al., 2016; Wang et al., 2016). In the current study, the observed PNSD at 260 m was used to evaluate the model performance. Figure 2 presents the comparison of the simulated PNSD with the observations. The model realistically reproduced the evolution of PNSD at 260 m at the measurement site. In the observation, there are five cycles of conversion from clean days to pollution days. Once the pollution episode was over, an obvious new particle formation event occurred, such as the events in September 3, 12, 19, and 25. When the pollution level increased, the PNSD shifted to the large diameter side. The model successfully captured the NPF events and the growth of particles in the aforementioned pollution episode. Because the atmosphere at higher level is not susceptible to local sources, the observation at 260 m was more representative than that at the ground level. The number concentration of particles from 100 nm to 685 nm was accurately reproduced, with a normalized bias less than 40% and a correlation coefficient of 0.70. The consistency between the simulation and observation suggests the good performance of model in producing reasonable number concentration of regional aerosol particles, especially in the climate-relevant size range. However, the number concentration of particles from 15 nm to 25 nm was overestimated. On one hand, the measurements have analytical errors (Du et al., 2017). On the other hand, the model also has several uncertainties. First, the model used the monthly mean emissions and therefore could

not simulate the diurnal variation of traffic emission. In addition, the size distribution of primary emissions did not meet the assumed lognormal distribution. For example, traffic sources emit smaller particles than do industrial sources (Paasonen et al., 2013; Kumar et al., 2014). Second, the nucleation scheme also has some uncertainties (Zhang et al., 2010; Yu et al., 2018). Nevertheless, the main features of NPF events and the growth of particles were captured by the model. Generally, our model produced the aerosols of real atmosphere and the simulation results were reasonable.

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The reasonable performance of our model in simulating OA components and PNSD gives us the confidence to further analyze the composition of newly formed particles through nucleation and subsequent growth (i.e., SPs) in our model. Figure 3a shows the simulated mean contribution of sulfate, nitrate, ammonium, and OA to the mass concentration of SPs in September. Figure 3b displays the contribution of LV anthropogenic OA (LV-AOA), LV biogenic OA (LV-BOA), SV biogenic OA (SV-BOA), and SV anthropogenic OA (SV-AOA) to the mass concentration of OA in SPs. OA was the major component of SPs, followed by sulfate, nitrate, and ammonium (Fig.3a). Among the components of OA in SPs, AOA accounted for 67%, substantially larger than the 33% of BOA, suggesting the dominant role of AOA in particle growth. In terms of volatility, LV-OA comprised 67%, of which LV-AOA was responsible for 50% and LV-BOA for 17%. Our model calculated the gas-phase concentration of LV-SOG and its kinetic condensation on size-resolved SPs. The large fraction of LV-AOA in the OA of SPs indicates their critical role in the growth of SPs. Furthermore, LV-AOA is an indicator of aged atmosphere, and its large contribution to OA suggested the influence of regional transport of OA and precursors of OA from surrounding areas to Beijing. The aging and growth during the lifetime of SPs in the atmosphere could greatly enhance their regional impact. In addition to the local emissions of OA precursors (Guo et al., 2014), our results also highlight the importance of regional sources of OA precursors in the growth of new particles.

5.2 Global and regional distribution of OA

There are two important characteristics of OA that influence particle growth and particle number concentration: (1) the concentration of OA and (2) the condensation

behavior of OA. The concentration of OA is dependent on OA sources and sinks. The condensation behavior of OA is closely related to the separation of POA and SOA and their volatility distribution. Therefore, these properties of OA are given as the background to discuss global and regional particle number concentration. Figure 4 presents the surface distribution of OC concentration and the fraction of secondary OC (SOC) in the base experiment. In our model, OA is formed by primary emission and the partitioning of gas-phase species onto preexisting OA. Therefore, the distribution of OC is well correlated with the amount of primary emission and SOA precursors. Globally, high concentrations of OC are located in continental regions with large emissions. Over China and India, OC concentration can exceed 10 μg/m³ because of the high emissions from intense anthropogenic activities. In the tropical region of Africa, OC concentrations are larger than 5 µg/m³ because of the biomass burning. Over North America and Europe, OC concentrations are below 3 µg/m³. The model successfully reproduced this spatial difference reflected by the observations in North America and China. The highest concentrations are located in central-eastern China. In the second domain (Fig.4c), the highest concentrations of simulated OC can exceed 15 µg/m³ over some areas in the Sichuan Basin and North China Plain. The model reproduced the OC concentrations over North America and the west-east gradient of OC concentrations over China. However, the absolute concentrations of OC at most sites in China were underestimated by ratios between 40% and 75%.

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The discrepancy over China can be explained by the following reasons: (1) the time difference between the simulation and observation could lead to this discrepancy; (2) the OC emission over China had large uncertainties (Zhao et al., 2013; Li et al., 2017) and the emission (3.54 Tg C yr⁻¹) used in our study are 47% lower than a top-down constrained emission (6.67 Tg C yr⁻¹) (Fu et al., 2012); (3) distributing POA emissions to the five volatility bins can lead to a lower OC concentration than that when assuming POA to be non-volatile (Donahue et al., 2009); (4) the OA pathways included and the parameters used in the model still have uncertainties (e.g., SOA product yields and lacking aqueous SOA formation); (5) the model resolutions are not high enough to capture the hot points in cities with small urban areas,

especially those cities in western China (e.g., Lhasa and Dunhuang). In Sect.5.1, using the same model configuration but with different emissions, the model can accurately reproduce the BC and OA concentration in Beijing, which indicates that the emission is the most influential factor responsible for the inconsistent model performance in OC simulation for the year 2010. The observation-constrained OC emission by including non-traditional emission, incorporating new SOA formation pathways, and increasing the model resolution would help reduce the simulation bias of OC over China.

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In Fig.4b, it can be seen that SOC dominated most regions of the globe with the fraction above 70%. Over China, POC dominated the eastern regions whereas SOC dominated the western regions. In the eastern regions, the higher primary emissions led to the lower fraction of SOC in OC even though the SOC concentrations are higher than those in western regions. In the VBS in our model, there are three pathways forming SOA, i.e., oxidation of POA, oxidation products of anthropogenic and biogenic VOC and IVOC. The simulations of most previous studies (Kanakidou et al., 2005; Tsigaridis et al., 2014) have indicated that biogenic SOAs (BSOAs) are dominant over the global scale because their major sources from the oxidation of biogenic VOCs. However, our simulations (Fig.5) indicated that anthropogenic SOA (ASOA) is as important as the biogenic one, especially over areas with large anthropogenic emissions. Over some areas in India and eastern China, ASOA concentrations can exceed 7 µg/m³, significantly greater than the BSOA concentrations (< 3 µg/m³). Even over South America and Africa, ASOA has concentrations of 1~3 µg/m³ because of the large contribution of IVOC and POA emitted from biomass burning. Although a recent study indicated that the VBS representation could not capture the physicochemical dependencies of SOA formation on dominant pathway from isoprene (Jo et al., 2019), considering the possible underestimation of BSOA could not reduce the importance of ASOA. The higher concentrations of ASOA than BSOA are also demonstrated by other studies (Matsui et al., 2014; Lin et al., 2016; Zhao et al., 2016). For example, adding an additional SOA correlated with the CO emission can improve the observed OA concentration

(Spracklen et al. 2011). In the second domain simulation (Fig.5c and 5d), it is more clearly seen that ASOA has the higher concentrations than BSOA over China. In North China Plain, concentrations of ASOA were above 3 µg/m³ while concentrations of BSOA were below 1 µg/m³. Previous modeling studies using VBS (Han et al., 2016; Lin et al., 2016) have also suggested that ASOA is dominant in North China. Observation analysis indicated that ASOA was the greatest one among the contributors of SOA sources, which differs considerably from the reported cases in developed countries (Ding et al., 2014; Li et al., 2017; Tang et al., 2018). In addition, our simulation considered the SOA formation from IVOC, which has been proved to be a large contributor to SOA (Zhao et al., 2016; Yang et al., 2019). Clearly, it is the allowing of POA to be volatile and including the SOA formation from IVOC that constitute the larger sources of ASOA. In the case of lower simulated OC concentration than observation, the underestimation of SOC fraction in source regions (Fig.4d) indicated that the model underestimated ASOA and the underestimation of ASOA is a significant cause of the negative bias in OC over China. The exact contribution of ASOA to SOA would be greater than estimated in our simulation. The substantial contribution of ASOA to SOA suggests the critical role of ASOA in particle growth over areas with intense anthropogenic emissions, which will be discussed in Sect.5.4.

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The volatility distribution of SOA is a factor controlling not only the mass concentrations of OA but also the size distributions of aerosol particles through microphysical processes. The VBS framework can simulate the volatility distribution of OA in five saturation concentration bins. Study of Riipinen et al. (2011) suggested that approximately half of the condensing mass needs to be distributed proportional to the aerosol surface area to explain the observed aerosol particle growth. The condensation of this part of OA is governed by gas-phase concentration rather than the equilibrium vapour pressure, which is how our model calculates the growth of LV-SOA to particles. The volatility distribution of SOA is an important factor affecting the global and regional distribution of particle number concentrations. Figure 6 shows the surface layer spatial distributions of SV-SOA and LV-SOA

concentrations. Globally, high SV-SOA concentrations are mainly located in the continental source regions. By contrast, LV-SOA distribution is more homogeneous and its contribution to SOA is lower in source regions. The differences between spatial distributions of SV-SOA and LV-SOA are mainly caused by different formation processes. The major source of SV-SOA is the oxidation of VOCs and its distribution is almost consistent with the source areas of VOCs. However, LV-SOA is formed from the further oxidation of SV-SOGs. Multi-generation aging processes enable LV-SOA to form both in the source regions and downwind regions. The continental areas with higher emission have a lower contribution of LV-SOA. In downwind regions, LV-SOA has a higher concentration than does SV-SOA. Even over source areas, such as North America and Europe, LV-SOA also has a higher concentration than does SV-SOA. These results indicate that the multi-generation aging of OA in the VBS produce a higher concentration of LV-SOA and consequently the wider spread of OA, which has a large impact on the role of OA in particle formation processes. Over China, SV-SOA has a concentration of 3-10 µg/m³ and is dominant over source areas in the eastern region. LV-SOA has a concentration of 2-5 μg/m³ in the eastern region and 0.6-2 μg/m³ in the western region. Measurement analysis suggested that the OA and SOA in Beijing in China are more volatile than those of cities in Europe and North America (Xu et al., 2019). Our study indicated that, in addition to the different emission sources, the greater volatility of SOA is also caused by the lower contribution of LV-SOA to SOA although the concentration of LV-SOA over eastern China is higher than that over Europe and North America.

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5.3 Global and regional distribution of particle number concentration

Figure 7 displays the simulated surface layer horizontal spatial distributions of annual mean number concentrations of CN10 and the fraction of CN10 that is secondary. The observed CN10 values provided in Table S1 are also illustrated in Figs.7a and 7c for comparison. High concentrations of CN10 in the surface layer are located in the regions with large anthropogenic emissions (Fig.7a). The highest concentrations of annual mean CN10 are over central-eastern China and the Sichuan basin, and their values can be larger than 10000 cm⁻³. Over the eastern United States,

most areas of developed European countries, and India, the values of annual mean CN10 are over 5000 cm⁻³. Over South America and South Africa, CN10 concentrations are also high because of the biomass burning emission. Affected by continental sources and ship emissions, CN10 concentrations over the coastal regions and adjacent seas close to the continent can be over 1000 cm⁻³. Over the polar regions and the oceans far from continents, CN10 concentrations are lower than 300 cm⁻³. The model accurately reproduced the aformentioned spatial variation of CN10 represented by observations in different environments. By a more specific comparison (Fig.8), where the simulation values are compared in a scatter plot with corresponding observations at 34 sites given (Table S1), the simulations of annual mean concentration of CN10 agree quite well with the observations, within a factor of two for most of the sites. The spatial pattern of CN10 over the second domain (Fig.7c) is similar to that of the corresponding region in the first domain (Fig.7a), but the gradients of CN10 are characterized more precisely because of the higher horizontal resolution. For example, the high concentrations of CN10 over southern Hebei are clearly depicted in Fig.7c than in Fig.7a. The observed annual mean CN10 concentration (12000 cm⁻³) in Shangdianzi in eastern China was five times greater than that of Waliguan (2030 cm⁻³) in western China. The corresponding simulated CN10 concentrations, 14380 cm⁻³ and 2780 cm⁻³, well reflected this regional difference.

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Both SPs formed through nucleation and subsequent growth and direct emission of PPs can contribute to atmospheric particle number concentration. It is important to quantify the contribution of these two sources in different parts of the globe. In Fig.7b, it can be seen that SPs are dominant in most parts of the globe except for regions with large primary emissions, such as eastern China, India, and southern Africa. The low contribution of SPs in these regions is due to the strong scavenging of secondary particles by primary particles and the low nucleation rate caused by the competition of PPs for condensable gases. Although secondary aerosol species are high in these regions (Fig.6), they tend to act as coating species on PPs rather than form new particles; thus, PPs are dominated. This spatial pattern is consistent with the results of

previou study (Yu and Luo, 2009). However, the fractions of SPs in CN10 are lower than those in CN3 reported in Yu and Luo (2009) because of the dominant contribution of secondary nucleation to particles in 3 to 10 nm. A boundary from northeast to southwest China can be seen to separate the areas dominated by SPs from that by PPs over China (Fig.7d). This phenomenon is also caused by the large difference in emissions between the western region and eastern region of the country.

5.4 The mixing state of OAs and their growth to new particles

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In addition to particle number concentration, mixing state of aerosols is also necessary to evaluate aerosol impacts on climate. The VBS framework treats the emitted OA with diverse volatilities and thus allows them to be partitioned among different aerosol particles through condensation. In addition, the evolving volatility due to oxidation in the atmosphere makes the microphysical behavior of POA different from that of the nonvolatile POA. In our model, semi-volatile organics are temperature-driven partitioned through the equilibrium assumption while low-volatility species is kinetically condensed on the particles. Figure 9 presents the fraction of organic species residing in aerosols of different types (i.e., SPs, sea salt, dust, BC, and OC) defined in our model. Most of the organic species reside in OC, SPs, and BC particles, suggesting the intense mixing of anthropogenic aerosol species (Fig.9). In the Southern Hemisphere, the fractions of organic species residing in SPs are above 30%, larger than that of OC particles. In the Northern Hemisphere, organic species mainly reside in OC particles because of the higher concentration of POA and the subsequent partition. The fractions of organic species residing in SPs are lower, but still considerable, indicating the important role of organic species in forming particles over the whole globe. Because of differences in emissions and the associated microphysical processes, there are distinct spatial variations of organic species distribution among different continents. Over the United States, 30-40 % of OA resides in SPs. By contrast, this fraction is below 20 % over China. In China, significant differences also exists between the western and eastern regions. The dominant contribution of semi-volatile species to OA (Fig.6) and their partition proportional to LV-OA lead to a higher fraction of organic species residing in OC

particles over eastern China. The mixing of natural aerosols and organic species is also demonstrated in Fig.9. Over most areas of the globe, 15 % of organic species are distributed in dust particles, which could greatly modify the properties of the dust particles and thus their climate forcing over these regions (Huang et al., 2019).

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Previous studies have indicated that organic species are the major components of aerosols (e.g., Zhang et al., 2007; Jimenez et al., 2009) and low-volatility organic species can greatly enhance the growth of new particles (e.g., Yu, 2011; Tröstl et al., 2016). Our results also indicated the substantial distribution of organic species in SPs. For this reason, the contribution of LV-SOG to the growth of SPs was analyzed. Figure 10 shows the ratio of LV-SOG to H₂SO₄ and the ratio of low-volatility organic species to sulfate that reside in SPs. The concentration of LV-SOG is a factor of ~1.5-10 higher than that of H₂SO₄ over many parts of the continents and the adjacent oceans but is lower in East Asia, the eastern United States, southern Europe, and northern Africa where emissions of SO₂ are high. Especially, over the areas in Sichuan Basin and eastern China (Fig.10c), the concentrations of H₂SO₄ are considerably higher than those of LV-SOG. Different from the simulation of Yu (2011), our results included anthropogenic LV-SOG and therefore the ratios of LV-SOG to H₂SO₄ are higher, especially in the regions influenced by continental sources and oceans with ship emissions. In Fig.10b and Fig.10d, the contribution of low-volatility organic species to the growth of SPs, presented as the concentration ratio of low-volatility organic species to sulfate, is higher in the Southern Hemisphere and lower in the Northern Hemisphere where continental sources of SO₂ are larger. Despite being lower, the contribution remains considerable (approximately 10-20 %) over Europe and North America. Similar to the contribution of ASOA to SOA, LV-SOA residing in SPs is dominated by anthropogenic contributions over POA source areas (as is the case in Beijing; Fig.3). The condensation growth of SPs through low-volatility organic species can enhance their survival rate and therefore could increase the contribution of SPs to particle number concentration. These results highlight the importance of ASOA in new particle growth over polluted regions, such as eastern China and India.

725 5.5 Sensitivity of particle number concentration to volatility of POA

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In the VBS, POA is treated as volatile species and allowed to be aged by oxidation in the atmosphere; thus, it is necessary to explore the uncertainties associated with this treatment of volatility distribution. In addition, the size distribution of POA and the associated microphysical processes are also modified because of this treatment. Therefore, the sensitivity of particle number concentration to the volatility of POA and the assumed size distribution of PPs are discussed here. Figure 11 displays the change ratio of CN10 number concentrations in the LV_POA experiment, HV POA experiment, PPD0.5 experiment, and OCD0.5 experiment relative to that in base experiment. Overall, CN10 concentrations changed little when POA volatilities were in the inter-quartile range of measurements (Fig.11a and b). When using the low volatility distribution of POA, PPs number concentrations were increased by 5-10% over most areas in the Northern Hemisphere; the concentrations decreased by 5-10 % when the high volatility distribution of POA was used. By contrast, SPs number concentrations only exhibited minor changes over the areas with the strongest emissions. Because of the dominant contribution of SPs, CN10 had no clear change in most regions of the globe. By contrast, the size distribution of emitted PPs has large influence on the CN10 concentration. When the median diameter of BC and OC was set as half the size used in the base experiment, concentrations of CN10 increased by 50-150 % over the areas with large emission sources of BC and OC, which were too high to match the observations shown in Fig.7a. For example, the CN10 in the PPD0.5 experiment was greatly overestimated when compared with the observed concentration (12000 cm⁻³) in Shangdianzi in eastern China. Therefore, halving the median diameter of BC and OC in the PPD0.5 experiment could not represent the real situation. However, halving the median diameter of OC only leads to the increase of CN10 by 10-50 % over eastern China. Over the other areas with high emission, no observations were available for comparison. Considering the other factors affecting the simulation of CN10, we cannot confirm that the assumed median size of OC is too small in the PPD0.5 experiment. Moreover, the emitted OC particles are volatile and can re-evaporate after dilution (Robinson et al., 2007; Donahue et al.,

2009). The assumption of OC size distribution should consider OC volatility. To clarify this matter, it is necessary to measure the size distribution of freshly emitted primary particles and compare the model results with observation in polluted atmosphere dominated by PPs.

6 Discussion and conclusion

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Laboratory studies and field observations have highlighted the importance of the sources of OA, volatility distribution, and the microphysical behavior of organic species in particle formation processes; however, OA processes are still poorly represented and this lack of knowledge contributes considerably to model uncertainties in simulating aerosol microphysical properties. In this study, a new global-regional nested aerosol model was developed to simulate detailed microphysical processes in the real atmosphere. The new model combines the APM module and a VBS module to simulate the microphysical processes of OA. In the model, the OA in the lowest volatility bin is treated as non-volatile or low-volatility species and their condensation is simulated using the kinetic method. The OA in other volatility bins is simulated using equilibrium partitioning. Using this framework, both the condensation of secondary inorganic species (i.e., sulfuric acid, nitrate, and ammonium) and the condensation of organic species with various volatilities (i.e., low-volatility organic compounds and semi-volatile organic compounds) are simultaneously simulated, which represents a major advance of our new model. The concentration of low-volatility organics is separately calculated and the condensation of H₂SO₄ and LV-SOG on size-resolved SPs is explicitly simulated, along with the condensation of LV-SOG on PPs. Therefore, the growth of LV-SOG to new particles and the aging of PPs by organic species were represented in a realistic way.

Compared with most models in the second phase AeroCom (Tsigaridis et al., 2014; Mann et al., 2014) and recently developed new models (e.g., Yu, 2011; Patoulias et al., 2015; Gao et al., 2017), our model includes more comprehensive sources of SOA, especially anthropogenic SOA, by using the VBS framework. In addition, allowing POA to evaporate and re-condense onto the particles makes its microphysical behavior resemble that of SOA and therefore gives new meaning to the

POA–SOA split, which has a substantial effect on global CCN formation (Trivitayanurak and Adams, 2014). The flexible framework of APM combined with VBS produces the different distribution of organic species in aerosols (i.e., the mixing state of OA), which has been found to cause substantial difference in the radiative effects of aerosols (Zhu et al., 2017). Box model analyses revealed that LV-SOA has a large fraction in the growing nucleation mode particles (Pierce et al., 2011). The comprehensive thermodynamic-kinetic approach treating the condensation and the partitioning of organic species that originated from biogenic and anthropogenic sources enble us to investigate the full role of organic species in the growth of new particles, which is vital for understanding the formation processes of particles relevant to radiative forcing and clouds (Shrivastava et al., 2017).

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The model with three nested domains was applied to simulate the aerosol components and PNSD in the Megacity Beijing during a period of approximately 1 month. The simulation results were evaluated by the observations at high level of the IAP tower, which is more representative than ground level observations in the regional scale. The simulated BC and OA components agreed well with the PMF analysis of the AMS measurements. The evolution of PNSD and NPF events were also nicely reproduced by the model. Our modeling analyses revealed that AOA accounts for the largest part of the OA of SPs and thus contribute considerably to the growth of SPs in Beijing. Molteni et al. (2018) indicated that highly oxygenated organic compounds that formed from anthropogenic VOCs can substantially contribute to NPF in urban areas. Observations in Beijing suggested that anthropogenic VOCs are major constituents of SOA (Ding et al., 2015; Yang et al., 2016). For the first time, the contribution of AOA to new particles was quantified because of the mixing state our model resolved in our study. Although the exact role of AOA in NPF is not perfectly clear, our study explicitly demonstrated the critical role of AOA in NPF in Chinese megacities, which can help elucidate the mechanism of more frequent occurrences of NPF events than theoretical prediction in polluted atmosphere (Kulmala et al., 2017; Chu et al., 2019). Comparisons with the observations collected from published data revealed that the model well reproduced the annual mean concentration of the observed OC at continent sites in America and the spatial pattern of OC in China. Because of the re-evaporation and oxidation of POA and the additional emission of IVOC, ASOA becomes dominant in SOA over POA source areas. At sites in different environments over the globe, the model produced the reasonable CN10 concentrations within a factor of two of observed values for most of the sites. We noted that LV-SOG, especially anthropogenic SOA, has a large contribution to new particle growth over areas with intense anthropogenic emissions, such as eastern China. The global simulation of Kelly et al. (2018) indicated that including large anthropogenic SOA sources could achieve results consistent with observations over mid-latitudes of the Northern Hemisphere. Simulation over East Asia also indicated that most of the OAs were from anthropogenic sources (Matsui et al., 2014). Together with these studies, our modeling results further provided the direct evidence of AOA in particle formation processes not only in Chinese megacities but also in other regions influenced by anthropogenic sources in the global scale.

Sensitivity analyses indicated that CN10 concentration changed only a little in the regions with the highest POA emission and had no clear change in most regions of the globe when POA volatilities were in the inter-quartile range of measurements. Although the size distribution of primary emitted particles has a large effect on the simulation of CN10, as suggested by other studies (e.g., Spracklen et al., 2006; Chang et al., 2009; Zhou et al., 2018), the simulation of the base experiment gave the better agreement with the observations than the sensitivity experiments and the conclusions will not be changed. Even so, the importance of the size distribution of primary emitted particles should be emphasized. Global model results in a related study suggested the high sensitivity of CCN to the assumed emission size distribution (Lee et al., 2013). Recently, Xausa et al. (2018) found that using the size-segregated primary particle number emissions can help make the number concentration of accumulation mode particles more closer to the measurements. The current simulation indicates to the importance of parameterization of the size distribution of emitted OC particles after considering their re-evaporation and condensation. Therefore, the

primary emissions must be constrained both in their size distribution and volatility. In addition, the simulated properties of OA were also determined by the parameters of the VBS module, the emissions inventory and meteorological fields input to the model, and the physicochemical processes in the model. Although our model provided the reasonable calculations comparable with the available observations and model results of other authors, it has room for further improvements in the future. For example, the size-resolved emissions of anthropogenic PPs may be used as the model input to reduce the associated uncertainties. The fixed parameters in VBS make it difficult to represent the real formation pathway of SOA and capture the response of SOA to emission changes (Jo et al., 2019). More accurate parameterizations considering the key physicochemical dependencies should be incorporated to update the VBS module in our model. More nucleation schemes may also be implemented into the model to investigate the influence of nucleation schemes on the aerosol number concentrations because the uncertainties from the nucleation scheme remain large (Dunne et al., 2016). Aqueous-phase formation processes of SOA have an evident influence on the particle properties and total SOA mass (Ervens et al., 2011), and these processes can close the gap between simulations and observations (Lin et al., 2014). Finally, the description of aerosol microphysical processes should be refined by including the aqueous formation of SOA in our model.

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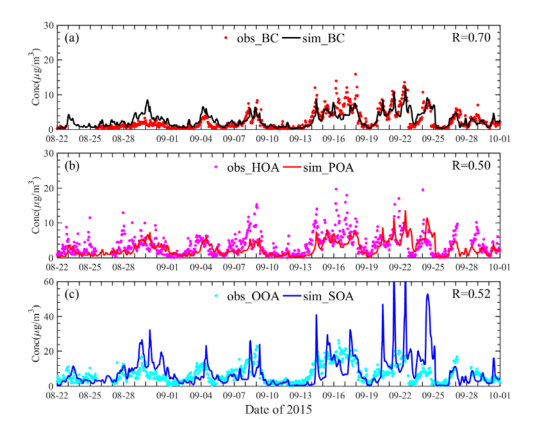


Fig.1 Comparison of the simulated and the observed (a) black carbon, (b) primary organic aerosol, and (c) secondary organic aerosol at the 260m height in Beijing from August 22 to September 30, 2015. All the observations were shown with dot points and the simulations with lines.

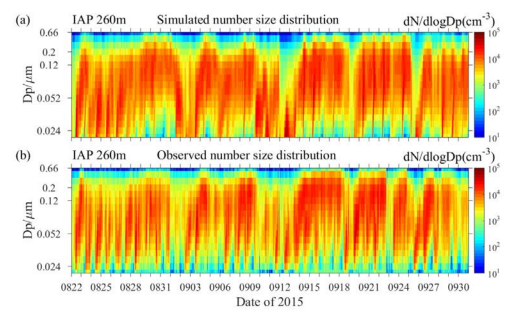


Fig.2 (a) Simulated and (b) observed particle number size distribution at high level (260 m) in Beijing from August 22 to September 30, 2015.

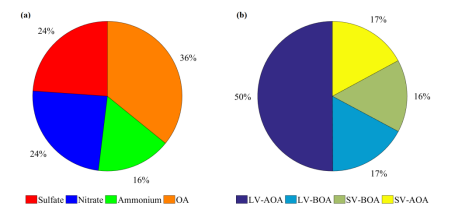


Fig.3 (a)The mean contribution of sulfate, nitrate, ammonium, and OA to the mass concentration of SPs and (b) the mean contribution of LV-AOA, LV-BOA, SV-BOA, and SV-AOA to the mass concentration of OA in SPs in September, 2015.

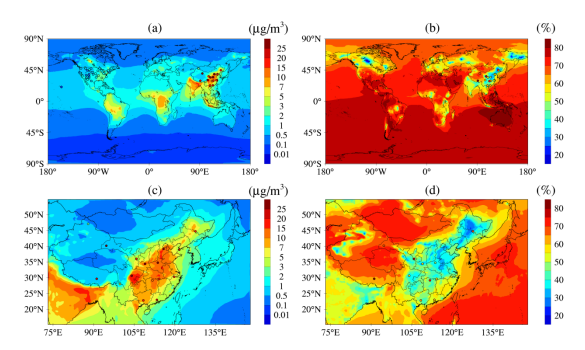


Fig. 4. Surface layer horizontal spatial distributions of organic carbon concentrations (left panel) and the fraction of OC that is secondary (right panel) over the first domain (top panel) and second domain (bottom panel). Observed OC and estimated fraction of secondary OC collected in Sect.4 are overlapped with shaded circles on the plots for comparison.

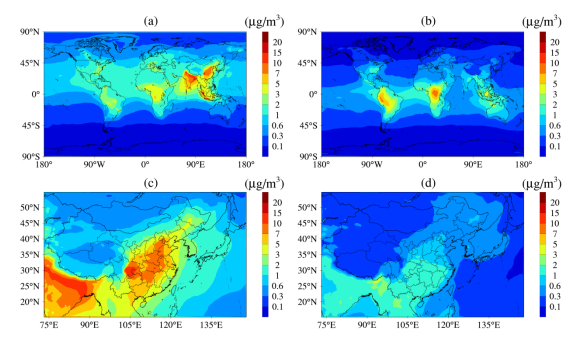


Fig. 5. Surface layer horizontal spatial distributions of ASOA concentrations (left panel) and BSOA concentrations (right panel) over the first domain (top panel) and second domain (bottom panel).

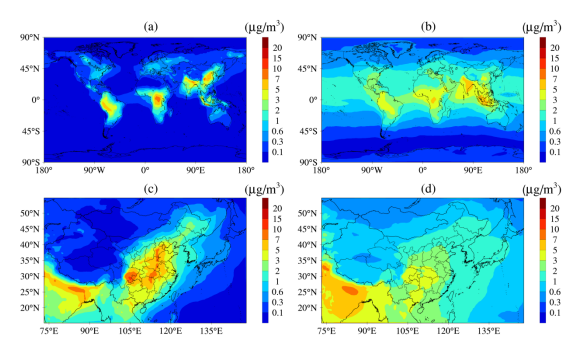


Fig. 6. Surface layer horizontal spatial distributions of SV-SOA concentrations (left panel) and LV-SOA concentrations (right panel) over the first domain (top panel) and second domain (bottom panel).

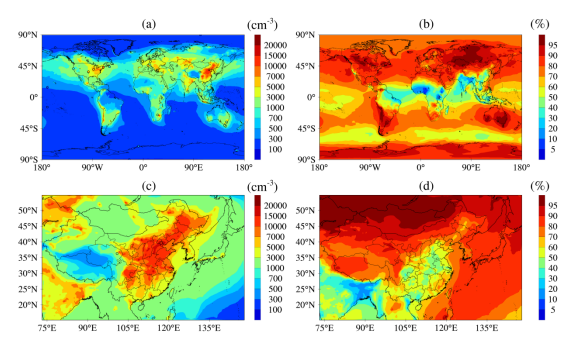


Fig. 7. Surface layer horizontal spatial distributions of annual mean number concentrations of CN10 (left panel) and fraction of CN10 that is secondary (right panel) over the first domain (top panel) and second domain (bottom panel). Observed CN10 values in Table 3 are also overlapped with shaded circles on the plots for comparison.

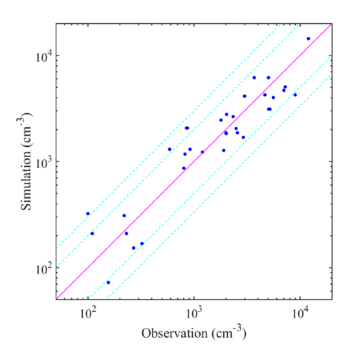


Fig. 8. Comparison of simulated and observed annual mean number concentrations of particles condensation larger than 10 nm at 34 sites listed in Table S1. The solid carmine line shows a 1:1 ratio and the dashed turquoise lines show ratios of 3:1, 2:1, 1:2, and 1:3.

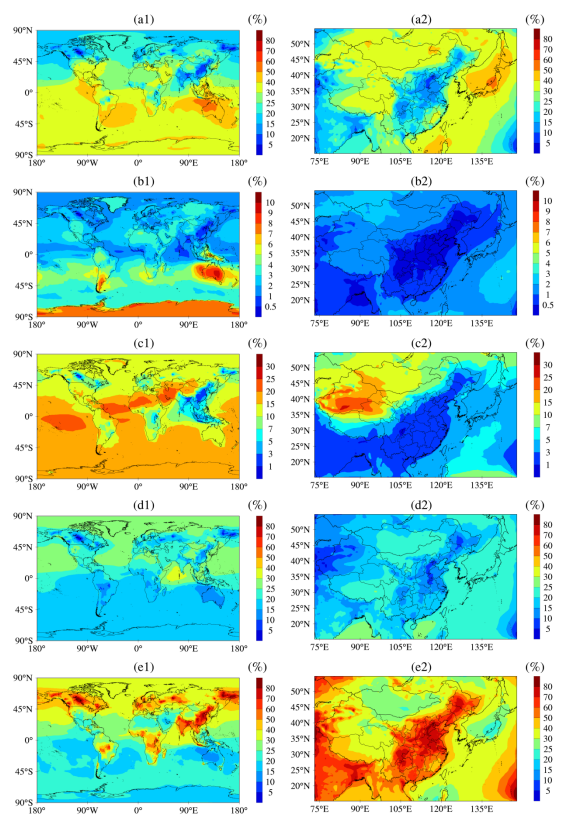


Fig. 9. Surface layer horizontal spatial distributions of the fraction of organic species that reside in SP (a1 and a2), sea salt (b1 and b2), dust (c1 and c2), BC (d1 and d2), and OC (e1 and e2) particles over the first domain (left panel) and second domain (right panel).

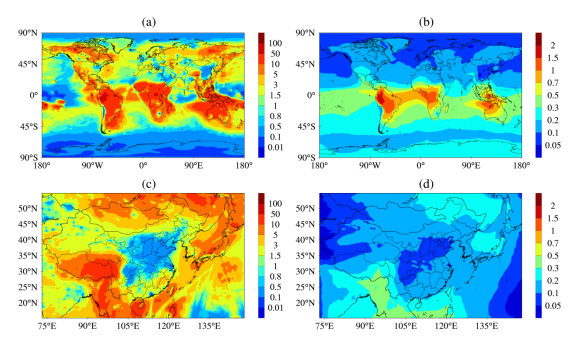


Fig. 10. The ratio of LV-SOG to H₂SO₄ (left panel) and the ratio of LV-OA to sulfate (right panel) that reside in SPs. Top panel is for the first domain and bottom panel for the second domain.

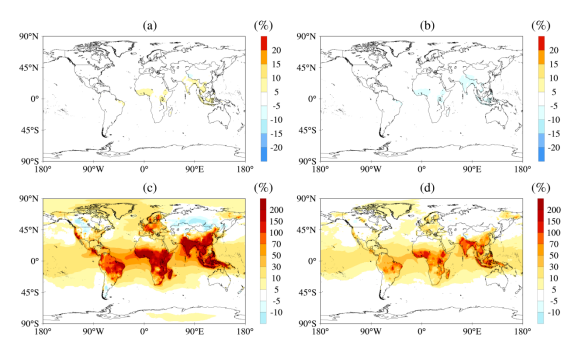


Fig. 11. Relative change of number concentrations of CN10 in (a) LV_POA experiment, (b) HV_POA experiment, (c) PPD0.5 experiment, and (d) OCD0.5 experiment to that in base experiment.

Table 1 The newly added tracers for simulation in microphysical processes

Experiments	Primary size	Volatility distribution
BASE	60, 150 (1.80, 1.80) for	0.27, 0.15, 0.26, 0.15, 0.17; 0.167, 0.167, 0.243,
	BC and POC	0.197, 0.226; 0.2, 0.1, 0.1, 0.2, 0.4
LV_POA	60, 150 (1.80, 1.80) for	0.34, 0.21, 0.3, 0.1, 0.05; 0.234, 0.217, 0.27, 0.157,
	BC and POC	0.122; 0.25, 0.15, 0.15, 0.2, 0.25
HV POA	60, 150 (1.80, 1.80) for	0.16, 0.21, 0.21, 0.19, 0.33; 0.11, 0.093, 0.217,

	BC and POC	0.217, 0.363; 0.15, 0.05, 0.05, 0.2, 0.55
OCD0.5	30,75 (1.80, 1.80) for	0.27, 0.15, 0.26, 0.15, 0.17; 0.167, 0.167, 0.243,
	POC	0.197, 0.226; 0.2, 0.1, 0.1, 0.2, 0.4
PPD0.5	30,75 (1.80, 1.80) for	0.27, 0.15, 0.26, 0.15, 0.17; 0.167, 0.167, 0.243,
	BC and POC	0.197, 0.226; 0.2, 0.1, 0.1, 0.2, 0.4

Data availability. All of the observation in this paper are provided in the manuscript. The simulation data can be available from the authors upon request (chenxsh@mail.iap.ac.cn, zifawang@mail.iap.ac.cn).

Author contribution. XC developed the model, performed the simulations and analysis, and prepared the manuscript with contributions from all co-authors. FY provided the code of APM module and modified the manuscript. WY coupled the VBS module and modified the manuscript. YS, WD, and JZ provided the observation data in Beijing and modified the manuscript. HC prepared the emission data and modified the model code. YW, LW, HD, ZW, QW, JL, and JA modified the manuscript. ZW guided the study and modified the manuscript.

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Competing interests. The authors declare that they have no conflict of interest.

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