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An improved framework for efficiently modeling organic aerosol (OA) considering primary OA evaporation and secondary OA formation from VOCs, IVOCs, and SVOCs[†]

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Organic aerosols (OA) constitute an important fraction of fine particulate matter (PM2.5) air pollution, yet accurate and efficient OA modeling within chemical transport models (CTM) remains a challenge. Volatility basis set (VBS) schemes for OA have demonstrated improved performance in simulating OA, particularly for primary organic aerosol (POA), but their computational complexity impedes application to advanced modeling tasks, such as detailed source apportionment. Conversely, simpler "two-product" schemes are efficient and compatible with source apportionment techniques but many of them tend to overestimate POA by treating it as non-volatile. Either VBS or 2-product schemes can perform well for secondary organic aerosol (SOA) depending upon the data and assumptions used to model SOA formation from precursors. In this study, we update the Comprehensive Air Quality Model with extensions (CAMx) "SOAP" 2-product modeling framework by (1) treating POA as semivolatile using an efficient scheme, (2) adding SOA formation from semivolatile organic compounds (SVOCs), and (3) adopting SOA yields derived from the widely-used Community Multiscale Air Quality (CMAQ) AERO7 scheme. The first update allows temperature-dependent partial evaporation of POA to SVOC, which is subsequently oxidized in the gas phase. For the latter two updates, SOA yields are updated to emulate the AERO7 scheme based on an offline conceptual model. We implemented these changes within the existing SOAP2 scheme of CAMx to create a new scheme called "SOAP3". A series of CTM simulations were conducted with the SOAP3 scheme to simulate OA and its components in China during July and November 2018. Results were validated against surface observations and compared to the SOAP2 and AERO7 schemes. Compared to SOAP2, SOAP3 substantially reduced POA proportions (by 10-24%) and increased SOA concentrations (by 45–193%) for selected regions. SOAP3 performs more like the AERO7 scheme than SOAP2 in terms of the simulated OA components and improved accuracy compared to observations. Uncertainties and limitations of the current SOAP3 scheme are also discussed. Our study demonstrates a feasible and readily implemented methodology for improving two-product OA modeling, which is currently employed in many CTMs.

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Environmental significance

Organic aerosol (OA) is a major contributor to ambient $PM_{2.5}$ and has different chemical components originating from diverse sources. Models are an essential tool for understanding OA sources, but model accuracy is important, and detailed model source attribution is highly desirable. Our new OA modeling approach can better predict the impacts of human activities and natural emissions on air quality and climate. This improved understanding can inform policy decisions aimed at reducing $PM_{2.5}$ levels, protecting human health, and mitigating climate change.

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1. Introduction

Human exposure to fine particulate matter $(PM_{2.5})$ is associated with adverse health outcomes and is reported to cause 4.14 million deaths worldwide annually (95% confidence interval: 3.45 to 4.80).¹ Consequently, $PM_{2.5}$ is a focus of air quality management efforts over the past decades worldwide. Organic aerosol (OA) is an important contributor to PM_{2.5}, with relative mass contribution ranging from 20-90% across different regions.² OA sources can be complex to characterize because the primary and secondary contributions are not necessarily distinct.3,4 The relative contributions of primary OA (POA) and secondary (SOA) vary spatially and temporally. SOA is formed in the atmosphere when gas-phase reactive organic compounds (ROC) are oxidized and form products that can migrate to the particle-phase either by condensation⁵ or via heterogeneous reaction.6 When atmospheric conditions favor secondary pollution events, SOA can account for the majority (60-90%) of OA.7-10 Traditionally, POA emissions were considered nonvolatile, *i.e.*, existing permanently in the particle phase.^{11,12} However, for many sources, this characterization oversimplifies the reality that the emitted ROC has many constituents with differing vapor pressures that consequently partition between the gas phase and an organic-aerosol phase according to ambient conditions.5,13 A state-of-the-science viewpoint considers ROC emissions as a continuum in volatility³ that spans from the most (*i.e.*, volatile organic compounds, VOCs) to least volatile compounds (i.e., POA).

Chemical transport models (CTMs) are crucial tools for understanding the sources of OA and evaluating the effectiveness of air quality management policies. The widely adopted volatility basis set (VBS) approach to OA modeling13 follows the "volatilitycontinuum" conceptual model by grouping OA precursor emissions into volatility bins defined by saturation vapor pressure (C^*) ranging from $10^{-3} \ \mu g \ m^{-3}$ (least volatile) to >10⁶ $\ \mu g \ m^{-3}$ (most volatile). However, the computational complexity of the VBS approach restricts its application in advanced modeling techniques like online source apportionment¹⁴ and sensitivity analysis.15 Quantifying OA source contributions is particularly important for air quality planners to develop effective emission control strategies. Some modeling studies have addressed this need by combining a VBS approach with "brute-force" sensitivity analysis,16-18 which defines the change in simulated SOA concentration resulting from an emission change as representing the source contribution. The "brute-force" method, however, is practically cumbersome (i.e. number of simulations scales linearly with the number of source contributions) and may not be suitable to retrieve source contributions when the relationship between concentration and emissions is non-linear.15,19,20 On the other hand, tagged species methodology, such as the Particulate Source Apportionment Technology (PSAT) implemented in the Comprehensive Air Quality Model with Extensions²¹ (CAMx) is an efficient tool for source apportionment and is considered more appropriate under non-linear situations.^{20,22} Unfortunately, the application species-tagging to VBS is challenging due to its complexity and would be computationally expensive.

The two-product SOA scheme, proposed by Odum *et al.*,²³ is simpler and more efficient than VBS and has been coupled with tagged species source tracking.^{21,24} The two-product model describes SOA formation from ROC oxidation using two semivolatile products, usually with high and low volatility that are derived by fitting chamber experiments.²³ Two-product schemes were developed to model SOA and have been implemented in CTMs in combination with a simple non-volatile POA scheme^{24,25} which tends to overestimate OA near emission sources.²⁶

This study aims to develop an efficient "volatilitycontinuum" modeling scheme that can support detailed source apportionment to assist air quality management. To do this, we start with an existing two-product scheme and make targeted modifications that improve coherence with the "ROCcontinuum" concept. Meanwhile, we retain the simplicity and computational efficiency of the two-product scheme to preserve compatibility with efficient source-tracking methodologies (work in progress). Our results demonstrate a feasible and readily implemented methodology for improving the existing two-product OA modeling employed in many CTMs.

2. Methods

2.1 Development of the CAMx SOAP3 scheme

CAMx is a widely used air quality model and provides a twoproduct scheme for SOA modeling, which is referred to as SOAP2.2 (hereafter SOAP2). The SOAP2 scheme, illustrated by the grey lines in Fig. 1, represents the oxidation of anthropogenic VOCs (AVOCs, including benzene, toluene, and xylene), intermediate volatility organic compounds (IVOCs), and biogenic VOCs (BVOCs, including isoprene, monoterpenes, and sesquiterpenes) into condensable gases (CGs), followed by equilibrium partitioning between CG and SOA:

SOA precursor + oxidant
$$\rightarrow$$
 CGs \leftrightarrow SOA (1)

Two condensable products, one more-volatile (CG1 for anthropogenic and CG3 for biogenic) and one less-volatile product (CG2 for anthropogenic and CG4 for biogenic), are utilized. In addition, non-volatile products (SOPA for anthropogenic and SOPB for biogenic, respectively) from the oxidation step instantly condense to form SOA. POA is treated separately from SOA and considered non-volatile in SOAP2.

The Community Multiscale Air Quality (CMAQ) model (version 5.30) includes the "AERO7" which tracks SOA formation from AVOCs, BVOCs and IVOCs using a combination of VBS, two-product and one-product approaches²⁷ (Table S3†). For instance, AERO7 uses a VBS approach with four bins to represent SOA formation from AVOCs and a VBS with seven bins for monoterpenes. Isoprene and sesquiterpene oxidation products are parameterized with two and one semivolatile products, respectively. IVOC in AERO7 is represented by pcVOC, which oxidizes with OH to form a low-volatility condensable vapor (pcSOG, $C^* = 10^{-5} \ \mu g \ m^{-3}$) with a molar yield of 1. The SOA aging process in the CMAQ AERO7 scheme involves particle-



Fig. 1 Schematic diagram of the SOAP3 scheme (updates to the existing SOAP2 scheme are presented in blue)

phase oligomerization (applied to SOA from isoprene, sesquiterpenes, and aromatics) or hydrolysis (applied to SOA from monoterpene-derived organic nitrates). POA is represented using a VBS in AERO7 with gas-phase OH oxidation to form SOA (see details in Section S1[†]).

CMAQ with AERO7 is widely-used, has been applied for OA over East Asia and achieved acceptable model performance,^{28,29} and has gone through frequent model updates with respect to OA formation.^{27,30–33} Therefore, we built the new SOAP3 scheme upon the existing SOAP2 scheme utilizing parameters derived from the AERO7 scheme. The updates made in the SOAP3 scheme are highlighted in blue in Fig. 1, and the key differences between SOAP2 and SOAP3 are summarized in Table 1. It is important to note that although SOAP3 was developed to emulate the AERO7 scheme, the SOAP3 and AERO7 schemes have fundamentally different structures. Another difference is that SOAP3 includes condensed-phase SOA photolysis in contrast to AERO7. The impact of SOA photolysis on simulated SOA concentrations is discussed in later section.

2.1.1 Partial evaporation of POA to SVOCs. The first SOAP3 update treats POA as being semivolatile instead of non-volatile in SOAP2. A new species, semi-volatile organic compounds (SVOCs), is added to the SOAP3 scheme to represent the fraction of POA that promptly evaporates to the gas phase after emission (Fig. 1). SVOCs reside in the gas phase and can undergo oxidation to form condensable products and therefore SOA. The evaporation of POA to SVOCs is parameterized as a temperature-dependent function (E = f(T), where *E* is the fraction of POA that evaporates to SVOCs) that mimics the POA treatment in the AERO7 scheme, as described below.

In the AERO7 scheme, POA emissions are allocated to five bins with volatility (*i.e.*, effective saturation concentration, C^*) ranging from 0.1 µg m⁻³ to 1000 µg m⁻³ at 298 K (Table S1†). For each bin (*i*), C^* can be adjusted to the local ambient temperature using the Clausius Clapeyron equation (eqn (2)).³⁴

$$C_i^* = C_{i,0}^* \frac{T_0}{T} \exp\left[\frac{\Delta H_i^{\text{vap}}}{R} \left(\frac{1}{T_0} - \frac{1}{T}\right)\right]$$
(2)

where C_i^* and $C_{i,0}^*$ are the saturation concentrations for volatility bin *i* at *T* and T_0 (=298 K), respectively. *R* is the universal gas constant, and ΔH_i^{vap} denotes the enthalpy of vaporization for volatility bin *i*. The detailed values for each parameter are given in Table S1.† For each bin, the particle phase fraction of POA is calculated using Pankow's partitioning theory:^{5,13,23}

$$F_i = \frac{C_{\text{OA}}}{C_i^* + C_{\text{OA}}} \tag{3}$$

where F_i is the fraction of POA in the particle phase for bin *i*, and C_{OA} is the ambient OA concentration. A C_{OA} of 50 µg m⁻³ was used to represent the high concentrations near the emission source. The influence of different OA concentrations on POA evaporation is discussed in Section 3.4. By summing over the five bins, the total fraction of POA that exists in the particle phase is calculated as:

$$1 - E = \sum_{i} f_i F_i \tag{4}$$

where f_i is the volatility distribution factor used to allocate POA emissions to each volatility bin (Table S1[†]). Take an ambient temperature of 290 K as an example. At 290 K, it is calculated that 45% of POA emissions exist in the particle phase. We

Species	Existing SOAP2 scheme	Newly developed SOAP3 scheme		
POA	Non-volatile	Semivolatile, with temperature-dependent evaporation fraction derived from AERO7		
AVOCs/IVOCs	Oxidized into CG1/CG2 and non-volatile product (SOPA)	Same as SOAP2 but with updated SOA yields derived from AERO7		
SVOCs	Not considered	Added in SOAP3, with SOA yields derived from AERO7		
BVOCs	Oxidized into CG3/CG4 and non-volatile product (SOPB)	Same as SOAP2 but with updated SOA yields derived from AERO7; adding SOA formation pathways from heterogeneous uptake of GLY/ MGLY and monoterpenes + NO ₃		

 Table 1
 Comparison of OA treatment in SOAP2 vs. SOAP3



Fig. 2 (a) Polynomial fitting between particle fraction of POA (1 - E) and temperature with $C_{OA} = 50 \ \mu g \ m^{-3}$; (b) fitting of SOAP3 SOA formation from SVOCs to AERO7 over a range of total OA concentration from 0.1 $\mu g \ m^{-3}$ to 50 $\mu g \ m^{-3}$.

performed the above calculation at various ambient temperatures to obtain the overall POA gas-particle partitioning as a function of temperature and fitted these results to a polynomial function between the particle fraction of POA (*i.e.*, 1 - E) and the ambient temperature (Fig. 2a). The high R^2 values (0.994) indicate that a polynomial function captures the relationship quite well, which SOAP3 applies to allocate the POA emissions between the gas phase (*i.e.*, SVOCs) and the remaining particle phase (*i.e.*, POA) according to the local temperature.

2.1.2 Adding SOA formation from SVOCs. POA that evaporates to SVOCs in the SOAP3 scheme undergoes oxidation and partitioning similar to the scheme for AVOCs and IVOCs:

$$SVOCs + OH = \alpha_{1,SVOCs} \times CG1 + \alpha_{2,SVOCs} \times CG2 + \alpha_{3,SVOCs} \times SOPA$$
(5)

SOA molar-vield coefficients (under high/low NO., conditions) in the SOAP3 scheme

where $\alpha_{1,\text{SVOCs}}$, $\alpha_{2,\text{SVOCs}}$, and $\alpha_{3,\text{SVOCs}}$ are stoichiometric coefficients. The rate constant for SVOC + OH is set to be 4×10^{-11} cm³ per (molecules per s). We derived SVOCs α values to reproduce the SOA mass yields simulated by the AERO7 scheme. To do so, we first calculate SOA yields from the evaporated fraction of POA after a representative amount of OH exposure (*e.g.*, 10^{10}

molecules per cm³) in the AERO7 scheme, using an offline conceptual model²⁸ (also see example calculation in Section S1†). This calculation is performed for a wide range of total OA concentrations from 0.1 μ g m⁻³ to 50 μ g m⁻³ (Fig. 2b). Meanwhile, the corresponding SOA yields from SVOCs in the SOAP3 scheme at different OA concentrations are calculated as:

SOA yield from SVOCs =
$$\sum_{i}^{N} \alpha_{i,\text{SVOCs}} \left(1 + \frac{C_{i}^{*}}{C_{\text{OA}}} \right)^{-1}$$
 (6)

where *N* now equals 3 for eqn (5) (compared to 10 in AERO7). A scatter plot of SOAP3-generated and AERO7-generated SOA mass at different OA concentrations can be established based on the α_i values. The three coefficients ($\alpha_{1,\text{SVOCs}}$, $\alpha_{2,\text{SVOCs}}$, and $\alpha_{3,\text{SVOCs}}$) were fitted to obtain the highest R^2 value and slope closest to one (Fig. 2b). The final stoichiometric coefficients were converted from g g⁻¹ to ppm ppm⁻¹ (Table 2) for model implementation.

2.1.3 Adjusting SOA yields from AVOCs and IVOCs. The AVOCs oxidation and partitioning process in the existing SOAP2 scheme is as follows:

$$AVOCs + OH = \alpha_{1,AVOCs} \times CG1 + \alpha_{2,AVOCs} \times CG2 + \alpha_{3,AVOCs} \times SOPA$$
(7)

	CG1	CG2	SOPA	
<i>C</i> * (µg m ⁻³)@300 К	14	0.31	0	SOA mass yields (g g $^{-1})$ at C_{OA} = 10 $\mu g \text{ m}^{-3}$
Benzene	0.1874/0	0/0	0.0036/0.1314	0.160/0.370
Toluene	0.0921/0	0.0123/0	0/0.126	0.082/0.300
Xylene	0.0637/0	0/0	0.005/0.174	0.047/0.360
IVOCs	0	0	0.964	1.000
SVOCs	1.79	1.73	0.16	1.813
	CG3	CG4	SOPB	
<i>C</i> * (µg m ⁻³)@300 К	26	0.45	0	SOA mass yields at $C_{OA} = 10 \ \mu g \ m^{-3}$
Isoprene	0.038	0.0076	0	0.047
Monoterpenes ^a	0.1665	0.038	0.031	0.159
Sesquiterpenes	1.405	0.1133	0	0.440

^a See Section S2 for organic nitrates yields from monoterpenes + NO₃.

Table 2



Fig. 3 Fitting of SOAP3 SOA formation from (a) benzene, (b) toluene, (c) xylene, (d) isoprene, (e) monoterpenes, and (f) sesquiterpenes to AERO7 over a range of total OA concentration from 0.1 μ g m⁻³ to 50 μ g m⁻³.

In SOAP3, we only updated the stoichiometric coefficients $(\alpha_{1,AVOCs}, \alpha_{2,AVOCs}, \alpha_{3,AVOCs})$ based on the AERO7 parameterization. To do this, we first calculated the AERO7 SOA yields from different AVOCs (*e.g.*, benzene, toluene, and xylene) at different OA concentrations (range from 0.1 to 50 µg m⁻³) using the same method illustrated by eqn (6), with relevant information listed in Table S3.† Similarly to SVOCs, the three coefficients were fitted to obtain results closest to AERO7 under high and low NO_x conditions (Fig. 3a–c). In AERO7, SOA aging from AVOCs is represented by oligomerization processes in the particle phase to form non-volatile products with a rate of 9.49 × 10⁻⁶ s⁻¹ (equivalent to a lifetime of 20.5 h), which is adopted in SOAP3.

IVOCs emissions have been reported to be an important precursor of SOA.^{35–38} The IVOCs oxidation and partitioning process in the existing SOAP2 scheme is similar to SVOCs (eqn (5)) and AVOCs (eqn (7)). The current values of α_1 , α_2 , and α_3

used in the SOAP2 scheme underestimate SOA yields from IVOC as compared to AERO7 (Section 3.2). Therefore, we updated the coefficients of IVOCs by fitting the SOA yields based on the AERO7 scheme. In AERO7, IVOCs are oxidized by OH to form an extremely low-volatility product ($C^* = 10^{-5} \,\mu g \,m^{-3}$) with a molar yield of 1.³² Accordingly, we set the molar yield from IVOCs as 1 in SOAP3 by setting $\alpha_{3,IVOCs}$ to 1 and $\alpha_{1,IVOCs}$ and $\alpha_{2,IVOCs}$ to 0.

2.1.4 Adjusting SOA yields from BVOCs. The SOA formation from BVOCs in the existing SOAP2 scheme is similar to AVOCs:

BVOCs + oxidant =
$$\beta_1 \times CG3 + \beta_2 \times CG4 + \beta_3 \times SOPB$$
 (8)

where β_1 , β_2 , and β_3 are stoichiometric coefficients. As shown in Section 3.2.4, SOA formed from BVOCs (*i.e.*, BSOA) in the existing SOAP2 scheme is significantly underestimated compared to AERO7. To resolve these discrepancies, we made

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several updates to the SOA formation from BVOCs in the new SOAP3 scheme. First, we updated the stoichiometric coefficients for isoprene, monoterpenes, and sesquiterpenes in SOAP3 by fitting them to AERO7 results using the same method as AVOCs (Fig. 3e and f). Second, while testing, we noticed that the organic nitrates from monoterpenes oxidation by NO₃ in CMAQ AERO7 represent an important contribution to BSOA aging by hydrolysis. In contrast, in the existing SOAP2 scheme, the formation of organic nitrates is significantly underestimated. Therefore, we adjusted the yield coefficients of organic nitrates in SOAP3 and accounted for the aging effects due to hydrolysis in AERO7 (see details in Section S2[†]). Lastly, SOA formation via heterogeneous uptakes of glyoxal (GLY) and methylglyoxal (MGLY) on the aerosol surface, which is also found to be a non-negligible pathway in AERO7, was now added in CAMx SOAP3 (see details in Section S2[†]).

2.2 Model configurations

We tested the new SOAP3 scheme and compared the results with the existing SOAP2 scheme in CAMx and the AERO7 scheme implemented in CMAQ using the same model configuration as our previous study.²⁸ The CAMx model configuration utilized the CB6 photochemical gas-phase mechanism,³⁹ with a static two-mode coarse/fine (CF) PM chemistry option incorporating the ISORROPIA inorganic gas-aerosol partitioning scheme,⁴⁰ along with the Regional Acid Deposition Model (RADM) for aqueous phase chemistry, the Zhang dry deposition option,⁴¹ and wet deposition. For CMAQ simulations, the model configuration included the CB6 gas-phase mechanism, RADM aqueous phase chemistry, ISORROPIA inorganic particulate thermodynamics, and the AERO7 aerosol scheme.27 SOA concentrations in China during July (summer) and November (autumn) 2018 were simulated, covering the entire country with a spatial resolution of 36 km (Fig. 4). Meteorological fields were simulated using the Weather Research and Forecasting (WRF) model (version 4.0)42 with configurations detailed in our previous studies.36 The anthropogenic emissions comprised the Multi-resolution Emission Inventory of China (MEIC, https:// www.meicmodel.org, accessed on December 1, 2022) for conventional air pollutants and the S/IVOCs emission inventory developed by Wu et al.43 BVOCs emissions were calculated using a recent offline version of the Model of Emissions of Gases and Aerosols from Nature (MEGAN version 3.2, https:// aqrp.ceer.utexas.edu/projects.cfm, accessed on September 25, 2022). For this study, the partial evaporation of POA emissions to SVOCs was treated outside CAMx as a preprocessing step (offline scheme) using gridded monthly averaged 2 m temperature simulated by the WRF model. POA evaporation is being moved inside CAMx (in-line scheme) for the next public release version of CAMx.

Simulations were conducted over all of China for July and November 2018, with a horizontal resolution of 36 km (Fig. 4). For each model/scheme, we conducted three parallel simulations with different configurations of the input emissions (Table S4†). For the "Base" scenarios, all SOA precursors, including VOCs, IVOCs, and SVOCs, were included. For the "No_SVOCs" scenarios, SVOCs emissions were excluded, and similarly, in the "No_S/IVOCs" scenarios, only VOCs emissions were considered for SOA formation. Different OA components, including POA, BSOA, and SOA formed from AVOCs (ASOA),



Fig. 4 Modeling domain with definitions of key regions and locations of 1477 national monitoring sites (in black diamonds), 13 sites with OC/EC observations (in blue triangles) and 2 AMS sites (in red stars).

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IVOCs (IVOC-SOA), and SVOCs (SVOC-SOA), were either direct outputs (*e.g.* POA, BSOA) or calculated as differences between two sets of scenarios. For example, IVOC-SOA is calculated as the difference of SOA between the "Base" and "No_SVOCs" scenario while SVOC-SOA is the difference between the "No_S-VOCs" and "No_S/IVOCs" scenario. We compared results for five key regions in China (Fig. 4), including the Beijing–Tianjin– Hebei (BTH), Yangtze River Delta (YRD), Pearl River Delta (PRD), Sichuan Basin (SCB), and Fenwei Plain (FWP).

2.3 Model performance evaluation

We evaluated the simulated concentration of $PM_{2.5}$, primary organic carbon (POC), and secondary organic carbon (SOC) following the same approach in our previous study.²⁸ The simulated concentrations of $PM_{2.5}$ were compared against surface observations of hourly $PM_{2.5}$ at 1477 national monitoring sites. Hourly observed OC/EC at 13 monitoring sites (see detailed locations in Table S5†) were used to evaluate simulated POC and SOC by applying the minimum OC/EC ratio method:⁴⁴

$$SOC = OC - (OC/EC)_{pri} \times EC$$
(9)

$$POC = OC - SOC \tag{10}$$

where $(OC/EC)_{pri}$ represents the minimum value of OC/EC ratio during the observation period (see Table S5†). We calculated the values of SOC/OC from the observed data and compared it with the simulated values of the SOAP2, SOAP3, and AERO7 schemes.

In addition, SOA factors resolved using the positive matrix factorization (PMF) analysis of measurements obtained with an aerosol mass spectrometer (AMS) at a supersite in Shanghai⁴⁵ and an Aerodyne time-of-flight aerosol chemical speciation monitor (ToF-ACSM) in Beijing⁴⁶ were used to further validate the model performance of simulated SOA. The assessment of model performance was conducted using well-established statistical metrics, including the correlation coefficient (*R*), mean bias (MB), normalized mean bias (NMB), normalized mean error (RMSE), and fractional bias (FB).

3. Results and discussion

3.1 Model performance of the SOAP3 scheme

Fig. 5 shows the spatial distribution of monthly averaged PM_{2.5} simulated by the SOAP3 scheme with observed values from 1477 national monitoring sites superimposed. The simulation results generally align with the spatial distributions and seasonal variations of observed PM₂ 5. Both months show slight overestimation, with MB (NMB) of 1.8 μ g m⁻³ (7%) in July and 0.6 μ g m⁻³ (1%) in November. Regionally, an overestimation of PM_{2.5} is observed over SCB (Table S6[†]), which is also found in our previous study.28 PM2.5 is underestimated over PRD and FWP in July and over BTH and FWP in November. We further evaluated simulated POC, SOC, and the ratio of SOC to OC (SOC/OC) at limited observation sites (Fig. S1-S5[†]). The model performance for simulated SOC varies across different sites (NMB ranging from -39.8% to 55.8% in July and -51.4% to 66.7% in November), with a general pattern of underestimation in November. In terms of SOC/OC (Fig. S3[†]), the original SOAP2 scheme notably underestimated observed SOC/OC, especially in November, while the results from the AERO7 scheme were closer to the observations. The newly developed SOAP3 scheme demonstrates substantial improvements, yielding slightly lower values compared to AERO7. However, all schemes failed to capture observed SOC/OC for specific sites in November, indicating the presence of other uncertainties, such as those related to emissions. Fig. S6[†] further compares the observed and modeled POA fraction at various monitoring sites, including observations collected from other studies (e.g. Chen et al.⁴⁷). The results indicate a comparable or slightly lower POA fraction between the observed values and those simulated by SOAP3, with the exception of four sites in the BTH region. The time series plots (Fig. S4[†]) reveal that the simulated POC concentrations demonstrate good performance for sites in the YRD region (specifically JS_CZ, SH, and SZ). However, for sites in the BTH region (namely BJ, TJ, and HB_CZ), the simulated POC concentrations and the fraction of POA/OA are notably higher than the observed values, regardless of the models used. This may suggest that the over-predictions were due to bias in the POA emissions in the BTH region.



Fig. 5 Comparison of the monthly average simulations and observations of PM_{2.5} by 1477 national monitoring sites, (a) July 2018, (b) November 2018.



Fig. 6 Simulated and AMS-based concentrations (a) by ToF-ACSM in Beijing (July 2018) and (b) by AMS in Shanghai (November 1 to 16, 2018).

Fig. 6 shows the simulated and AMS-based SOA concentration at the ToF-ACSM site in Beijing (July) and the AMS site in Shanghai (November). The original SOAP2 scheme underestimated the observed SOA concentrations, with an NMB of –60% in Beijing and –69% in Shanghai. In contrast, the AERO7 scheme showed a slight underestimation at one site (NMB of -7% for Beijing) and an overestimation at the other site (NMB of 6% for Shanghai). With the new SOAP3 scheme, the large underestimation of SOA in the SOAP2 scheme was greatly mitigated, resulting in NMB values of -11% and -24% at the Beijing and Shanghai sites, respectively. While not entirely comparable, the performance of the SOA simulation with SOAP3 met the "goal" benchmarks for OC and OM proposed by Huang et al.48 for the application of CTMs in China. The "goal" benchmarks, defined as NMB < \pm 35% for OC/OM, were derived from the top 33rd percentile values collected from existing studies. However, it is worth noting that during the initial half of the observation period in Shanghai, all schemes failed to capture the observed SOA concentrations. The exact cause for this underestimation remains elusive, as the biases in the simulated meteorological conditions were insufficient to explain the phenomenon. One potential explanation could be unaccounted emissions, highlighting the need for future investigation in future studies.

3.2 OA concentrations simulated by SOAP3

Fig. 7 shows the domain-averaged absolute concentrations of OA components simulated by the SOAP3 scheme and the other two schemes (*i.e.*, the original SOAP2 scheme and AERO7 scheme) for the selected five key regions. The corresponding spatial distributions for each component are presented in Fig. S7 and S8.† With the newly developed SOAP3 scheme, the simulated OA concentrations ranged from 7.5 μ g m⁻³ (PRD) to 11.7 μ g m⁻³ (FWP) in July and from 8.8 μ g m⁻³ (SCB) to 14.8 μ g m⁻³ (YRD) in November (Table S7†). Spatially, high OA concentrations were observed over BTH and YRD in July and November. Compared to the original SOAP2 scheme, SOAP3 simulated OA concentrations generally show a substantial increase of 29–77%. As discussed below, the changes in total OA concentrations from SOAP2 to SOAP3 reflect the contribution of

two opposing effects of reduced POA proportion (by 10–24%) and increased SOA concentration (by 45–193%) from different precursors in the SOAP3 scheme. On the other hand, compared to the AERO7 scheme, the SOAP3 simulated OA concentrations are lower by 4–21% (except for FWP in July and SCB in both months).

3.2.1 POA evaporation. POA simulated by SOAP2 and AERO7 show similar spatial and seasonal variations, although differing in magnitudes. In July, SOAP2 shows significantly higher values than AERO7, while in November, SOAP2 indicates slightly lower values. Both models simulate high POA concentrations over Northeast China, BTH, northern YRD, and Sichuan Basin, which closely follows the spatial distribution of POA emissions. With the SOAP2 scheme, domain-averaged POA concentrations ranged from 1.3 μ g m⁻³ (PRD) to 3.0 μ g m⁻³ (BTH) in July and from 3.0 μ g m⁻³ (PRD) to 9.8 μ g m⁻³ (BTH) in November. With AERO7, the simulated POA concentrations ranged from 1.0 μ g m⁻³ (PRD) to 1.9 μ g m⁻³ (BTH) in July and from 2.9 μ g m⁻³ (PRD) to 10.9 μ g m⁻³ (BTH) in November. Despite the higher POA concentrations in SOAP2 due to nonvolatile treatment, AERO7 compensates by introducing SVOCs emissions, which are not accounted for in SOAP2. The inclusion of SVOCs in AERO7, allocated partially to non-volatile bins, contributes to higher POA concentrations. Consequently, in November, where less POA evaporation occurs due to lower temperature, AERO7 shows slightly higher simulated POA concentrations compared to SOAP2.

The newly developed SOAP3 scheme incorporates a temperature-dependent evaporation function, allowing POA to partially evaporate into the gas phase. With this feature, POA concentrations simulated by SOAP3 are lower than SOAP2 by 21–29% in July, bringing results closer to those of AERO7. The SOAP3 simulated POA concentrations ranged from 1.0 μ g m⁻³ (PRD) to 2.2 μ g m⁻³ (BTH) in July and from 2.8 μ g m⁻³ (PRD) to 10.6 μ g m⁻³ (BTH) in November. Similarly to the AERO7 scheme, SVOCs are now incorporated into SOAP3, with the nonvolatile fraction of SVOCs emissions offsetting the reductions in POA concentrations due to evaporation, particularly in November when evaporation effects are weaker than in July.

3.2.2 SOA formation from the newly added SVOCs. SOA formation from the gas-phase portion of SVOCs is included in



Fig. 7 Domain-averaged concentrations of OA components simulated with different schemes in (a) July and (b) November 2018.

the SOAP3 scheme, with SOA yields based on the AERO7 scheme. As shown by Fig. 7 and Table S7,† the SVOC-SOA concentrations simulated by the SOAP3 scheme ranged from only 0.2 μ g m⁻³ (PRD) to 0.6 μ g m⁻³ (BTH) in July and from 0.5 μ g m⁻³ (BTH) to 0.9 μ g m⁻³ (YRD) in November, making it an overall small fraction (<10%) of the total OA concentrations. Meanwhile, the AERO7 scheme also indicates a low contribution of SOA formation from the SVOCs emissions, with even lower magnitudes. The small discrepancies between AERO7 and SOAP3 are explained in Section S1 of the ESI.†

The relatively small contribution from SVOC-SOA was also noticed in our previous study²⁸ but is different from Miao *et al.*,⁴⁹ who found a dominant role of SVOC-SOA (30–39% in July) to total OA concentrations over eastern China. The uncertainties associated with the estimated SVOCs emissions could be causing these differences in addition to the differences in model parameterizations.

3.2.3 Increased SOA formation from IVOCs. IVOCs emissions are considered an important SOA precursor. In the original SOAP2 scheme, however, the simulated IVOC-SOA concentrations were significantly lower (by 2–5 times) than those simulated by AERO7. For instance, in BTH, the domain-averaged IVOC-SOA concentration simulated in the SOAP2 scheme was 1.1 μ g m⁻³ in July and 0.6 μ g m⁻³ in November.

When the AERO7 scheme is applied, the corresponding values are 3.6 μ g m⁻³ and 2.8 μ g m⁻³, respectively. In the newly developed SOAP3 scheme, the SOA yields from IVOCs emissions were adjusted to mimic the AERO7 scheme. With this change, the SOAP3 simulated IVOC-SOA concentrations increase substantially, as compared to SOAP2, by 207–232% in July and 222–307% in November. Domain-averaged IVOC-SOA simulated with the SOAP3 scheme ranged from 0.9 μ g m⁻³ (PRD) to 3.6 μ g m⁻³ (BTH) in July and from 1.8 μ g m⁻³ (BTH) to 3.6 μ g m⁻³ (YRD). These values are much closer to those predicted by the AERO7 scheme. Although magnitudes differ, all three schemes exhibit similar seasonal variations with higher values in November (except for BTH and FWP). Regionally, elevated IVOC-SOA concentrations were observed in BTH, FWP, and YRD in July and shifted southward in November.

3.2.4 SOA formation from traditional VOCs. Simulated ASOA concentrations are relatively consistent across the three schemes compared to the other SOA components. With the SOAP3 scheme, domain-averaged ASOA concentrations ranged from 0.3 μ g m⁻³ (PRD) to 1.0 μ g m⁻³ (BTH) in July and from 0.8 μ g m⁻³ (BTH) to 1.5 μ g m⁻³ (YRD) in November. Spatially, higher levels of ASOA were predominantly observed in the BTH and FWP regions in July, whereas in November, elevated ASOA concentrations were mainly observed in the YRD and PRD

regions (Fig. S7 and S8†). Similar seasonal and spatial variations were observed for the other two models.

BSOA simulated by the SOAP2 and AERO7 schemes showed similar spatial (higher concentrations in southern regions, e.g., PRD and YRD) and seasonal (higher values in summer) variations, consistent with existing studies.50-52 However, the magnitudes differ significantly, with AERO7 exhibiting much higher values. In July, for instance, domain-averaged BSOA concentration simulated by the AERO7 scheme ranged from 4.4 $\mu g m^{-3}$ (BTH) to 8.5 $\mu g m^{-3}$ (YRD), approximately two times the value simulated by SOAP2. The much higher BSOA in the CMAQ AERO7 scheme, compared to the SOAP2 scheme, is likely associated with higher SOA yields from monoterpene53 and more SOA pathways included.30,33 With updated SOA yield coefficients and additional formation pathways (Section S2[†]) in the SOAP3 scheme, simulated BSOA concentrations significantly increased and exhibited greater alignment with the AERO7 results. In July, domain-averaged BSOA concentrations ranged from 4.0 μ g m⁻³ (BTH) to 7.1 μ g m⁻³ (YRD), while in November, they ranged from 0.8 μ g m⁻³ (BTH) to 5.9 μ g m⁻³ (PRD).

3.3 Relative contribution of different OA components

Fig. 8 compares the relative contribution of OA components predicted by SOAP3, SOAP2, and AERO7. In July, BSOA emerges as the most abundant component for all five regions, with relative contributions ranging from 35.1% in BTH to 68.5% in PRD with SOAP3. This trend holds for the other two schemes (except BTH in SOAP2), and AERO7 anticipates slightly higher BSOA contributions (37.7% in BTH to 76.3% in PRD). Meanwhile, IVOC-SOA contribution is comparable to BSOA in BTH and FWP under the SOAP3 scheme, reaching 31.7% and 30.4%, respectively. These values significantly exceed SOAP2 predictions and align with AERO7. The relative contribution from POA ranges from 13.1% (PRD) to 19.2% (BTH) with SOAP3, falling in between SOAP2 and AERO7 and showing closer agreement with AERO7. ASOA stands as a moderate contributor with SOAP3, ranging from 4.4% (PRD) to 8.6% (BTH), which is slightly lower than AERO7.

November sees POA ascend to the dominant OA component for all regions except PRD, reaching 39.0% (SCB) to 72.8% (BTH) with SOAP3. These values are considerably lower than SOAP2. BSOA remains dominant in PRD with 42.2%. IVOC-SOA retains its importance, ranging from 12.7% (BTH) to 24.0% (YRD). Both SOAP3 and AERO7 identify SVOC-SOA as the least abundant component, with SOAP3 predicting a range of 2.1% (PRD) to 5.4% (BTH) in July and 3.4% (BTH) to 7.1% (SCB) in November. Overall, AERO7 predicts the highest BSOA contribution, while SOAP2 predicts the highest POA contribution. The modifications implemented in SOAP3 position it between the other two schemes and much closer to the AERO7 scheme.

3.4 Uncertainties and limitations

Compared to the original SOAP2 scheme, the updated SOAP3 scheme exhibited substantial improvement in capturing the observations. However, there are still some uncertainties and limitations associated with the current implementation. Firstly, the evaporation of POA emissions in the current version of SOAP3 was implemented offline (i.e., outside CAMx) using monthly averaged temperatures instead of in-line using hourly temperatures. POA emissions exhibit diurnal variations, with higher emissions during the daytime and lower emissions during the nighttime. The higher daytime temperatures are expected to result in more POA evaporation, thus leading to lower POA concentrations. The full implementation of the dynamic POA evaporation in response to variations in hourly temperatures is under development and will be available in the public release of the latest CAMx version. The current POA partitioning scheme simplifies the dynamics of POA evaporation and further aging and, therefore, does not fully account for POA aging. This may cause the scheme to overestimate the POA/ SOA ratio in aged air masses.

Secondly, when deriving the temperature-dependent POA evaporation, a C_{OA} of 50 µg m⁻³ was used to calculate the particle phase fraction of POA. Fig. S9† demonstrates the impact of different OA concentrations on the partition of POA between gas and particle phases. Higher OA concentrations are expected to lead to higher fractions in the particle phase. At T =



Fig. 8 Domain-averaged OA proportion in July and November 2018 (including POA, ASOA, BSOA, IVOC-SOA, and SVOC-SOA).

290 K, the particle-phase fraction increases from 32% to 53% when C_{OA} increases from 10 µg m⁻³ to 100 µg m⁻³. We conducted a CAMx sensitivity simulation (case 1) using a POA evaporation function derived with a C_{OA} of 20 µg m⁻³ (Fig. S10[†]) and found more POA evaporation into the gas-phase, as expected, resulting in POA reductions of 14.7-16.8% in July and 14.6-15.7% in November for different regions. The impact on SOA was minimal (-0.1% to 1.0% in July and 2.2-5.8% in November) because of compensating impacts of having less POA for SOA to condense on being offset by having more gasphase precursors to produce SOA. However, it should be pointed out that the impact of C_{OA} on POA evaporation is less pronounced compared to the variations with temperature. Moreover, explicitly incorporating the dependence on C_{OA} , as in VBS scheme, could introduce additional uncertainty due to the introduction of a feedback loop where underpredicted C_{OA} would lead to more evaporation, thus resulting in further underestimated C_{OA} .

Thirdly, SOA yields have many uncertainties including the effect of NO_x on yields and SOA production from the IVOC category which has chemically diverse constituents. We investigated sensitivity to these two SOA yield uncertainties by conducting another CAMx sensitivity simulation (case 2) with two changes: (a) reducing the SOA yields from IVOC by half and (b) reducing the SOA yields of BVOCs (i.e., isoprene, monoterpene, sesquiterpenes) under high-NO_x conditions by 30% based on chamber studies by Sarrafzadeh et al.54 and Wildt et al.55 that carefully investigated this issue. As shown by Fig. S11,† the SOA yields from BVOCs in case 2 are reduced by 18.1-22.7% in July and less than 10% in November (except for PRD, 15.1%). The SOA yields from IVOC in case 2 are reduced by 49.6-50.8% and these reductions are similar among different regions and seasons. Consequently, the relative importance of different OA components changed slightly in case 2. For instance, for BTH, YRD and FWP in July, POA exceeds the IVOC-SOA, becoming the second most abundant OA component following BSOA. In November, the SOA yields from IVOC are less than BSOA in SCB and YRD. Thus there is a continuing need to refine SOA yield parameterizations for IVOC, BVOC and the effect of NOx on vields.

Lastly, condensed-phase SOA photolysis has been shown to exhibit a substantial effect on SOA formation.⁵⁶⁻⁵⁸ For example, by applying an empirical SOA photolysis rate (J_{SOA}) of 0.04% \times J_{NO2} (NO2 photolysis rate) in GEOS-Chem, Hodzic et al.⁵⁶ showed a decrease of 40-60% in SOA yields at mid-latitudes in the summer. This condensed-phase SOA photolysis is not considered in the CAMQ AERO7 scheme and but was implemented in the SOAP3 scheme. A sensitivity simulation was conducted on top of the current SOAP3 scheme by turning off the SOA photolysis. As illustrated by Fig. S12,† particle-phase SOA photolysis reduced the simulated SOA concentrations by 14% (PRD) to 33% (FWP) in July and by 17% (BTH) to 22% (SCB) in November. We consider representing SOA photolysis to be necessary in SOAP3 although the current parameterization is simple and can be improved as more information becomes available.

4. Conclusions and recommendations

Existing schemes for modeling OA in CTMs have strengths and limitations inherent to their formulation. VBS schemes unify the representation of POA and SOA, can describe complex relationships between emissions composition, atmospheric conditions and OA concentration, but incur relatively greater computational cost due to their complexity. The combination of a two-product SOA scheme with non-volatile POA is relatively simple and efficient, compatible with powerful modeling tools for detailed source contribution and sensitivity analysis, but fails to describe the partial evaporation of POA following emission and dilution to ambient conditions. In this study, we improved the CAMx SOAP2 scheme (non-volatile POA with twoproduct SOA) to a new SOAP3 scheme, resulting in better performance in simulating OA components while retaining computational simplicity. Improvements for SOAP3 are derived by parameterizing behaviors simulated by the CMAQ AERO7 scheme. First, by virtue of the fact that POA is semivolatile, we developed temperature-dependent partial evaporation of POA to SVOCs for SOAP3. Second, we added atmospheric oxidation of SVOCs to SOA by fitting parameters to the POA aging scheme in AERO7. We also updated SOA formation from IVOCs and VOCs by fitting parameters to the AERO7 scheme. Finally, we added formation pathways from monoterpene nitrates and heterogeneous uptake of GLY/MGLY to improve the simulation of BSOA. As a result, POA concentrations decreased (21-29%) with SOAP3 compared to SOAP2 in July. The SOA production from IVOCs increased by 2-3 times with SOAP3 compared to SOAP2. The newly-added SOA production from SVOCs contributed a small part of OA (2-7%) with SOAP3. In summary, SOAP3 performs more like the CMAQ AERO7 scheme than SOAP2 in terms of the simulated OA components and improved accuracy in comparison with observations. While we acknowledge that our results are subject to uncertainties, especially uncertainties associated with the precursor emissions (e.g. Ling et al., 59 Huang et al., 28 and Chen et al.47), our primary focus is developing an efficient modeling framework and we recommend that this framework should be maintained to account for improvements in science and community knowledge.

In terms of computational performance, a direct comparison of computational time between CAMx SOAP3 and CMAQ AERO7 simulations would be uninformative because many model processes, not only the SOA scheme, influence computational performance. The efficiency of the SOAP3 scheme relative to VBS schemes is evident from the smaller number of model species which influences computational requirements throughout the model. SOAP3 requires 12 model species (POA, SVOC, CG1, CG2, CG3, CG4, SOA1, SOA2, SOA3, SOA4, SOPA, and SOPB) to represent OA, whereas the CAMx 1.5D VBS OA scheme employs 50 model species and the CMAQ AERO7 OA scheme employs more than 60 model species. Furthermore, the efficiency of SOAP3 extends to advanced tagged-species source apportionment techniques like CAMx Particulate Source Apportionment Technology (PSAT) which is widely-used to track contributions from many source sectors and/or regions in a single simulation (*e.g.* Li *et al.*,⁶⁰ Liu *et al.*,⁶¹ Lu *et al.*,⁶² and Wang *et al.*⁶³). SOAP3 requires fewer PSAT tagging species than a VBS-style framework such as CAMx 1.5D VBS and the PSAT implementation is much simpler for SOAP3.

The current implementation of SOAP3 has several limitations that can be addressed by follow-up work. In this study, we modeled partial evaporation of POA offline using monthly average gridded temperatures, which will be improved by implementing an in-line calculation using hourly gridded temperatures. Sensitivity simulations indicate a non-negligible impact of photolysis on simulated SOA concentrations, highlighting the need for further investigation. We developed parameters for SOAP3 by analyzing behaviors of the CMAQ AERO7 scheme, but these parameters could also be derived by analyzing different schemes utilized in other CTMs, such as WRF-Chem⁶⁴ and CHIMERE.⁶⁵ Nevertheless, our study demonstrates a feasible and readily implemented methodology for improving the existing two-product OA modeling framework employed in many CTMs.

Data availability

The code for SOAP3 and data will be available upon request to the corresponding authors.

Conflicts of interest

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

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