Analysis of secondary organic aerosol simulation bias in the Community Earth System Model (CESM2.1)

Yaman Liu1,2, Xinyi Dong1,2, Minghuai Wang1,2, Louisa K. Emmons3, Yawen Liu1,2, Yuan Liang1,2, Xiao Li1,2, and Manish Shrivastava4

1School of Atmospheric Science, Nanjing University, Nanjing, China
2Joint International Research Laboratory of Atmospheric and Earth System Sciences & Institute for Climate and Global Change Research, Nanjing University, China
3National Center for Atmospheric Research, Boulder, CO, USA
4Pacific Northwest National Laboratory, Richland, Washington, USA

Correspondence: Xinyi Dong (dongxy@nju.edu.cn)

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Abstract. Organic aerosol (OA) has been considered as one of the most important uncertainties in climate modeling due to the complexity in presenting its chemical production and depletion mechanisms. To better understand the capability of climate models and probe into the associated uncertainties in simulating OA, we evaluate the Community Earth System Model version 2.1 (CESM2.1) configured with the Community Atmosphere Model version 6 (CAM6) with comprehensive tropospheric and stratospheric chemistry representation (CAM6-Chem) through a long-term simulation (1988–2019) with observations collected from multiple datasets in the United States. We find that CESM generally reproduces the interannual variation and seasonal cycle of OA mass concentration at surface layer with a correlation of 0.40 compared to ground observations and systematically overestimates (69 %) in summer and underestimates (−19 %) in winter. Through a series of sensitivity simulations, we reveal that modeling bias is primarily related to the dominant fraction of monoterpene-formed secondary organic aerosol (SOA), and a strong positive correlation of 0.67 is found between monoterpene emission and modeling bias in the eastern US during summer. In terms of vertical profile, the model prominently underestimates OA and monoterpene concentrations by 37 %–99 % and 82 %–99 %, respectively, in the upper air (> 500 m) as validated against aircraft observations. Our study suggests that the current volatility basis set (VBS) scheme applied in CESM might be parameterized with monoterpene SOA yields that are too high, which subsequently results in strong SOA production near the emission source area. We also find that the model has difficulty in reproducing the decreasing trend of surface OA in the southeastern US probably because of employing pure gas VBS to represent isoprene SOA which is in reality mainly formed through multiphase chemistry; thus, the influence of aerosol acidity and sulfate particle change on isoprene SOA formation has not been fully considered in the model. This study reveals the urgent need to improve the SOA modeling in climate models.

1 Introduction

As one of the most important contributors (20 %–90 %) to total fine atmospheric particles (Kanakidou et al., 2004), organic aerosol (OA) plays an important role in the climate system by affecting the radiation budget (Ghan et al., 2012). OA consists of primary organic aerosol (POA, also called primary organic matter, POM) emitted directly from biomass burning, fossil fuel combustion, biological compounds, etc. and secondary organic aerosol (SOA) formed via the oxidation of volatile organic compounds (VOCs) (Hallquist et al., 2009; Tsiganidis et al., 2014; Shrivastava et al., 2017). Chamber studies have revealed the important role of biogenic VOCs such as monoterpenes (Docherty and Ziemann, 2003; Kristensen et al., 2016; Signorell and Bertram, 2009) and isoprene (ISOP; Kroll et al., 2005; Nguyen et al., 2010, 2014; Paulot et al., 2009) in SOA production, and the contribution...
of anthropogenic VOCs such as aromatic compounds emitted from vehicle emissions and solvents also has important influence in urban areas (Nakao et al., 2012; Sato et al., 2010).

Understanding the atmospheric burden and spatiotemporal distributions of OA is one of the key priorities in atmospheric research because of the central roles it plays in regulating both climate and air quality. The radiative forcing effect of OA has been assessed with climate models through tremendous efforts during the past decades (Ghan et al., 2012; Myhre et al., 2013; Sporre et al., 2020; Chen and Gettelman, 2016), yet the limited capability of climate models in terms of simulating the productions and depletions of OA introduces large uncertainties. Substantial divergences were reported for models employed in the framework of Aerosol Comparisons between Observations and Models (AeroCom) phase II project even with the same set of emissions input. The burden of OA varied greatly for 28 models in the range of 0.6−3.8 Tg, with OA lifetime ranging from 3.8 to 9.6 d (Tsigeridis et al., 2014). As OA loading and properties of aerosols varied, the estimated radiative forcing of OA ranged from −0.06 to −0.01 W m$^{-2}$ among the 16 participating models (Myhre et al., 2013), revealing the fundamental uncertainty of OA simulation.

Modeling discrepancies largely come from the lack of a consensus in the representation of chemical composition and formation processes of OA among different models (Tsigeridis et al., 2014; Goldstein and Galbally, 2007). Although laboratory and chamber studies have revealed thousands of new reactions and new species related to VOCs and SOA, these reactions and species are usually simplified and grouped into a few functions and lumped with fewer species in the models to make it possible for simulating. Many unclear SOA formation processes have to be approximated as the knowledge is still under development (Canakidou et al., 2004; Hallquist et al., 2009). Thus, different models may use different simplified functions, lumped species definitions, and approximation methods to represent the overall SOA-related processes. For example, SOA chemistry was represented with the two-product method (Lack et al., 2004; Heald et al., 2008) in the earlier model (Lamarque et al., 2012). Since the late 2000s, volatility basis set (VBS) methods (Donahue et al., 2006; Robinson et al., 2007) have been widely adopted by different models due to the advantages over two-product method for considering the volatility (Lack et al., 2004; Heald et al., 2008). Since the 2010s, pilot studies started to include reactive uptake of isoprene epoxides (IEPOX) formation of SOA through aqueous-phase reactions into regional (Shrivastava et al., 2019; Karambelas et al., 2014) and global models (Marais et al., 2016; Zheng et al., 2020; Jo et al., 2019). Despite the tremendous efforts in the early stage, models still underpredicted measured SOA mass concentration by 1−2 orders of magnitude. Hodzic et al. (2016) therefore suggested corrected stronger yields of SOA formation which took into account the influence of vapor wall losses in chamber studies which was considered in

Tilmes et al. (2019). However, due to the different gas-phase chemistry, dry and wet deposition schemes, and heterogeneous chemistry schemes, the simulated OA may be different for the same VBS configuration in different models (Hodzic et al., 2016; Tilmes et al., 2019). These attempts not only validated the parameterizations and chemical pathways derived from measurement studies but also extended the understanding of SOA formation on a scale broader than the chamber. Once formed, most of OA undergoes chemical aging (Zhang et al., 2007) with volatility and hygroscopicity changing, but such processing is poorly understood due to inadequate relevant observations. While some models consider species-dependent aging reactions and the subsequent volatility change, some simply apply a constant aging rate (Tsigeridis et al., 2014; Jo et al., 2013; Donahue et al., 2006; Zhao et al., 2016). In the remote areas of the United States (US), OA and organic carbon (OC) concentrations show opposite bias in half of the participating models (Tsigeridis et al., 2014), indicating the models lack a consensus representation of SOA production and depletion.

To reveal the uncertainties associated with OA simulation in climate models, we evaluate a recent version of Community Earth System Model version 2.1 (CESM2.1) in this study with multiple observational datasets in the US. The model has been widely applied for OA climate effect assessment purposes (Gettelman et al., 2019; Glotfelty et al., 2017; Tilmes et al., 2019; Jo et al., 2021), and a significant portion of improvements has been implemented in the latest version regarding the chemical mechanisms (Tilmes et al., 2019). In the previous CESM version (Lamarque et al., 2012), SOA chemistry was represented with the two-product method (Lack et al., 2004; Heald et al., 2008). The next big update was reported by Tilmes et al. (2019), in which the two-product method was replaced by VBS following the work by Hodzic et al. (2016). Although CAM-chem has been applied in many studies including the AeroCom program (Tsigeridis et al., 2014), the evaluation of simulated OA concentration has not been well documented or thoroughly discussed. We focus on the validation over the US because it has long-term surface measurements and flight campaigns that provide solid observational data. We first evaluate the spatiotemporal characteristics of the simulation bias, then probe into the chemical mechanism to identify the origins through a series of sensitivity runs, and finally demonstrate the urgent need to both improve the current parameterization of the SOA production scheme and implement a more comprehensive production mechanism in climate model.

2 Methods

2.1 Model

Community Earth System Model is a coupled earth system model composed of atmosphere, ocean, land, sea-ice,
land-ice, river, and wave models (Danabasoglu et al., 2020). CESM2 (versions 2.0 and 2.1) includes two versions of model top: the Whole Atmosphere Community Climate Model version 6 (WACCM6) with 72 vertical layers up to about 150 km and the Community Atmosphere Model version 6 (CAM6) with 32 vertical layers up to about 40 km. CAM6 has a simplified chemistry and simplified OA scheme, while CAM6 with a comprehensive chemistry and comprehensive OA scheme is called CAM6-Chem which is updated compared to previous versions. The Model for OZone and Related chemical Tracers (MOZART) chemical mechanism covering the troposphere and stratosphere (referred to as MOZART-TS1) is used in CAM6-Chem. Emmons et al. (2020) reported the updates of MOZART-TS1 in CESM2.1, including the oxidation of isoprene and terpenes, organic nitrate speciation, and aromatic speciation and oxidation, and thus improved the representation of ozone ($O_3$) and SOA precursors. The most recently released CESM2.2 includes a new version (TS2) of MOZART tropospheric chemical mechanism with updates for isoprene and terpene chemistry (Schwantes et al., 2020) aiming at further improving $O_3$ simulation. In this study we also briefly compare the results between TS1 and TS2, as will be discussed in Sect. 3.3.

Both biogenic and anthropogenic VOCs are considered in CAM6-Chem with improved gas-phase chemical mechanisms (Emmons et al., 2020) and new SOA representation (Tilmes et al., 2019). CAM6-Chem applies the volatility basis set (VBS) scheme (Bergström et al., 2012; Hodzic et al., 2016; Shrivastava et al., 2011, 2013; Donahue et al., 2006; Robinson et al., 2007) by lumping SOA precursors based on their volatility bins with simulate SOA production. In CAM6-Chem, SOA and the condensable gas-phase SOA precursors (SOAG) are categorized into five bins with the saturation concentrations ($C^*$) of 0.01, 0.1, 1.0, 10.0, and 100.0 $\mu$g m$^{-3}$ (Tilmes et al., 2019). Compared with the simple SOA scheme which proportionally calculated SOAG based on emissions of precursors (Liu et al., 2012), this VBS scheme was demonstrated to improve CESM performance with a smaller bias of OA concentration over remote regions when evaluated against aircraft observations (Shrivastava et al., 2015; Tilmes et al., 2019). The VBS approach relies on empirical parameterizations fitting to chamber experiments; thus, the parameters vary between models. The current CAM6-Chem configures the VBS scheme and parameters by following the work of Hodzic et al. (2016) with GEOS-Chem as the host model, which differs significantly from CAM-Chem in terms of SOA-related modules such as gas-phase chemistry, aerosol dynamics, and dry and wet depositions. Consequently, the same scheme and parameterization employed by Hodzic et al. (2016) may result in different performance within CAM-Chem. Tilmes et al. (2019) provided a comprehensive comparison between the simple SOA scheme and the VBS scheme in terms of simulated OA burden and radiative forcing, and the simulations were validated against two flight campaigns, yet the evaluation against ground surface measurements at a different temporal scale (e.g., annual, seasonal) over different geophysical areas has not been thoroughly discussed.

### 2.2 Observations

Surface measurements from the Interagency Monitoring of Protected Visual Environments (IMPROVE) network and multiple aircraft campaigns in the Northern Hemisphere are incorporated in this study to validate model performance and also facilitate the analysis of SOA trend. IMPROVE is a long-existing program currently managed by the US Environmental Protection Agency (EPA) and is designed to measure the chemical composition of ambient fine particles and its spatial and temporal information (Solomon et al., 2014). We used 1988–2019 daily data for seasonal cycle and long-term trend evaluation. As IMPROVE measures organic carbon (OC) mass instead of OA, the concentration of OA observations is derived with the ratio of OA to OC (OA/OC) which is determined by the aging process. IMPROVE data have been widely employed in many studies for model evaluation purposes (Hodzic et al., 2016; Shrivastava et al., 2015; Tsagaridis et al., 2014). The OA/OC of IMPROVE data varies between sites, and the mean value used in this study is 1.8 (1.79–2.02) as recommended by Malm and Hand (2007). It should be noticed that not all sites have observations for the whole study period, and thus only those (140 sites, locations shown in Fig. 1) with more than 10 years of continuous data are used in this study to avoid measurement bias.

Free troposphere measurements from a total of five aircraft field campaigns are employed to validate simulated vertical OA profiles including CalNex (California Nexus; Ryerson et al., 2013), DC3 (Deep Convective Clouds and Chemistry; Barth et al., 2015), SENEX (Southeast Nexus; Warneke et al., 2016), SEA4CRS (Studies of Emissions and Atmospheric Composition, Clouds and Climate Coupling by Regional Surveys; Toon et al., 2016), and FRAPPE (Front Range Air Pollution and Photochemistry Experiment; Flocke et al., 2020). Flights were located in the area of the contiguous United States (CONUS) and took place between 2010 and 2014 with more details presented in Table 1 and Fig. 3.

### 2.3 Simulation configurations

To reproduce the observed meteorological conditions and allow for the direct comparison with OA measurements, a specified dynamic simulation (SD) is conducted from 1987 to 2019 with a spin-up time of 1 year; thus, our discussion can focus on chemical mechanism performance of CAM6-Chem. This experiment (referred as CAM-Chem-SD) uses the FCSD component set in which CAM6 physics, troposphere/stratosphere chemistry (MOZART-TS1) with VBS SOA scheme, historical emission, and offline meteorological field are applied. In detail, temperature, horizontal winds, and surface fluxes are nudged to Modern-Era Retrospective
analysis for Research and Applications (MERRA2) fields for detailed comparisons to field experiments and specific observations. The horizontal resolution is $0.9^\circ \times 1.25^\circ$, and vertical resolution is 32 levels with model top at $\sim 45$ km (Emmons et al., 2020). Prescribed historical sea surface temperatures (SSTs) are used in the FCSD component set. Anthropogenic and biomass burning emissions from 1987 to 2014 are from the standard Coupled Model Intercomparison Project round 6 (CMIP6) (Eyring et al., 2016) simulations, and emissions after 2014 are from the SSP585 scenario which is based on the shared socioeconomic pathway 5 (SSP5) (O’Neill et al., 2017) and forcing levels of Representative Concentration Pathways 8.5 (RCP8.5). Biogenic emissions are calculated with the Model of Emissions of Gases and Aerosol from Nature (MEGAN) in CESM (Guenther et al., 2012; Emmons et al., 2020). Moreover, another simulation of CESM2.2 which is the latest released version of CESM, referred to as CAM-Chem-SD(TS2), is conducted with MOZART-TS2 gas-phase emission model.
chemistry (Schwantes et al., 2020) from January 2013 to February 2014 with the first 2 months as spin-up time. The FCSD component set is also used in CAM-Chem-SD(TS2). Except for the difference of gas-phase chemistry, the SOA scheme is also improved in CAM-Chem-SD(TS2) compared with CAM-Chem-SD. The NO$_3^-$ dependence of SOA formation in CAM-Chem-SD(TS2) is not considered in CAM-Chem-SD. Thus, we compared CAM-Chem-SD and CAM-Chem-SD(TS2) to investigate the impact of NO$_3^-$ dependence on SOA formation.

The current VBS scheme in CAM6-Chem represents SOAG production from nine precursors through 15 reactions. The 15 reactions and chemical formulas of related species from Emmons et al. (2020) are shown in Tables S1 and S2 in the Supplement. To identify their contributions to total SOA mass and associated simulation uncertainties, we conducted 15 sensitivity experiments with one of the reactions turned off in each experiment. The 15 sensitivity simulations and a 14-month baseline simulation are set up from January 2010 to evaluate the contribution of each reaction to total SOA production in the VBS scheme. To exclude the influence of potential extreme meteorological conditions or emission inputs, these sensitivity runs are configured with the FC2010climo component set and Newtonian relaxation time of 3 h. The FC2010climo component set is the same as the FCSD component set except that the emissions are a 10-year average used for each year of the simulation. As one reaction is turned off for each sensitivity experiment, the difference between base and the sensitivity experiment represents the influence of this specific reaction (Table 2). For instance, the no ISOP-OH-SOAG experiment excludes SOAG produced by ISOP + OH reaction (Reaction 1 in Table S2), with other configurations the same as the base run (CAM-Chem-climo) experiment.

### 3 Results and discussion

#### 3.1 Evaluation against IMPROVE

We first validate the surface OA simulation with IMPROVE data for the CAM-Chem-SD scenario and find an overall significant overestimation with the results and statistics presented in Fig. 1 and Table 3. As demonstrated in Fig. 1, CESM2.1 shows substantial simulation bias with normalized mean bias (NMB) varying from $-73.87\%$ to $176.47\%$ at different sites with daily data pairs compared to IMPROVE measurements. The model noticeably overestimates annual average surface OA concentration over continental US (CONUS) with a mean bias (MB) of $0.41 \mu g \cdot m^{-3}$ and NMB of $20.27\%$. It also shows large regional differences as demonstrated by the sharp contrast between the two subdomains: the eastern US (EUS; $25-49^\circ$ N, $65-95^\circ$ W) and the western US (WUS; $25-49^\circ$ N, $95-125^\circ$ W) as presented in Fig. 1. Prominent overestimation is found over EUS with NMB more than $100\%$ at nine sites, while moderate underestimation is shown over WUS. Our validation suggests that the updated SOA representation (by applying VBS) helps improve the performance of CAM6-Chem compared to its earlier versions. CAM4-Chem was reported to underestimate OC by $\sim36\%$ compared with IMPROVE dataset with a correlation coefficient of $0.41$ (Lamarque et al., 2012), and CAM5-Chem was found to overestimate with NMB by $24\%$ at urban sites and $217\%$ at remote sites (Tsigidis et al., 2014). The AeroCom phase II multi-model ensemble mean showed NMB of $-48\%$ (range $-85\%$ to $24\%$) at urban locations and $-70\%$ (range $-38\%$ to $217\%$) at remote locations on a global scale, suggesting that the performance of CESM2.1 is consistent with other climate models.

Annual and seasonal variations of surface OA from the model and IMPROVE are presented in Fig. 2 to examine the model’s capability of reproducing the temporal variation. The model generally reproduces the annual decreasing trend of OA concentration for the whole CONUS domain and is in good agreement with observations in spring and fall but substantially overestimates in summer and slightly underestimates in winter, as shown in Fig. 2a. The monthly average of the simulation is found to have larger overestimation in warmer months (May–September) than cooler months, as shown in Fig. 2b. In EUS, the simulation predominantly overestimates surface OA concentration in summer by $4.26 \mu g \cdot m^{-3}$ ($131.15\%$) but successfully reproduces the temporal change with a strong correlation with observations of 0.60 (Table 3), as shown in Fig. 2c. Compared with the CONUS domain, surface OA concentration from the simulation at EUS shows an even greater overestimation during

<table>
<thead>
<tr>
<th>Index</th>
<th>Experiment ID</th>
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<tbody>
<tr>
<td>B1</td>
<td>CAM-Chem-SD</td>
</tr>
<tr>
<td>B2</td>
<td>CAM-Chem-SD (TS2)</td>
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<tr>
<td>B3</td>
<td>CAM-Chem-climo</td>
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<tr>
<td>E1</td>
<td>no ISOP + OH</td>
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<td>E2</td>
<td>no ISOP + O$_3$</td>
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<td>E3</td>
<td>no ISOP + NO$_3$</td>
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<td>E15</td>
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Table 2. CESM experiments used in this study. SVOC signifies the semi-volatile organic precursor of VBS SOA.
warmer months, as shown in Fig. 2d. In WUS, simulated OA shows a slow decreasing trend in summer ($-0.02 \mu g \, m^{-3}$ per decade), while the observations indicate an ascending trend ($0.23 \mu g \, m^{-3}$ per decade). The large interannual variations from 1999 to 2019 are shown in observed surface OA concentration mainly due to the influence of wildfires (Malm et al., 2017). The simulated surface OA concentration also has large interannual variation but does not show an increasing trend due to lower value after 2017. It needs to be emphasized that historical emissions are used from 1987 to 2014 and SSP585 emissions after 2014 in the CAM-Chem-SD simulation, which means the emissions do not exactly match the observed condition after 2014. The model shows smaller bias
in WUS but also a poor correlation of 0.36 (Table 3) in summer, as shown in Fig. 2e and f.

The large modeling bias of OA in summer over EUS may be attributed to the discrepancy in simulating SOA because POA is proportionally determined by emissions from fossil fuel and biofuel that have smaller seasonal diversity (see Fig. S1 in the Supplement). Biomass burning also contributes POA, but wildfire or prescribed fire emission is minimal in EUS (van der Werf et al., 2017). Production of SOA is represented by the MOZART-TS1 and VBS module as mentioned in Sect. 2. MOZART-TS1 has been demonstrated in a few studies (Emmons et al., 2020) to overestimate summertime surface O$_3$ over the southeastern US; thus, it may subsequently introduce uncertainties while simulating the VOC chemistry. Besides, the performance and uncertainty of the VBS scheme in CAM6-Chem remain largely undocumented regarding whether the VBS scheme tends to overestimate or underestimate. With the same VBS scheme but implemented in GEOS-Chem, Hodzic et al. (2016) reported a simulation bias of 34.8% through the validation against IMPROVE data, and the bias was explained by evaporation of OC from IMPROVE samples (Kim et al., 2015) and uncertainties in the boundary layer parameterization. The performance of CAM-Chem-SD seems better as our simulation bias for the CONUS domain is 20.27%. But with the larger biases in simulated OA spatial and temporal trends shown in Figs. 1 and 2, it is critical to realize that the overall evaluation statistic is averaged from large overestimations and underestimations from sites across the CONUS domain.

### 3.2 Evaluation against flight campaigns

We find systematic underestimation in the upper air through the model evaluation against aerosol mass spectrometry (AMS) measurements from flight campaigns, with the profiles shown in Fig. 3 and evaluation statistics summarized in Table 1. These campaigns were conducted over the land area of North America during 2010–2014, and most of the measurements were collected in summer with the rest of the data in late spring and early fall. Simulations are paired with measurements by choosing the nearest model grid at the corresponding time at an hourly scale. OA concentrations are underestimated in upper air (> 500 m) by −20% to −70% compared to the aircraft measurements. The model shows larger bias when compared with SENEX, DC3, and SEAC4RS that mostly cover EUS, while the vertical profiles measured in WUS (CalNex and SENEX) are reproduced bet-

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**Table 3.** The correlation coefficient (CC), mean bias (MB), and normalized mean bias (NMB) between observations (five field campaigns and IMPROVE surface measurements) and CAM-Chem-SD.

<table>
<thead>
<tr>
<th>Observations</th>
<th>CC</th>
<th>Mean obs. (µg m$^{-3}$)</th>
<th>Mean sim. (µg m$^{-3}$)</th>
<th>MB (µg m$^{-3}$)</th>
<th>NMB (%)</th>
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</tbody>
</table>
The most prominent bias is found in the comparison with DC3 campaign at $\sim 7$ km (geometric altitude above mean sea level) due to the aircraft samples of smoke plumes from the High Park fire west of Fort Collins (Barth et al., 2015), suggesting that fire emission plume structure is not properly represented because of the coarse model grid resolution. In general, the model is found to notably underestimate upper-air OA for all flight campaigns with larger discrepancies over EUS. We also examine the simulated and observed isoprene and monoterpenes profiles during the SEAC4RS campaign, as shown in Fig. 3g. These two biogenic VOCs (BVOCs) are reproduced well between the surface and $\sim 2$ km altitude but are also substantially underestimated at $> 2$ km height, suggesting oxidation of these VOCs that is too fast at high altitudes or uncertainties in the parameterizations of convective transport and vertical mixing. Validation against the recent ATom (Atmospheric Tomography Mission aircraft campaign) measurements from Wofsy et al. (2018) also suggested underestimation in the upper air over marine areas (see Fig. S2 and Table S3), consistent with the recent multi-model evaluation results reported in Hodzic et al. (2020). The photolytic depletion of SOA is represented in the VBS scheme of CAM6-Chem by following Hodzic et al. (2016) which demonstrated this stronger removal process can help lower the GEOS-Chem model overestimation of upper-air OA over remote areas (mostly over ocean). But the underestimations of high-altitude SOA within CAM6-Chem model over CONUS land area are different from GEOS-Chem and are likely related to model-to-model differences in physics, aerosol chemistry, and wet removal.

3.3 Uncertainties in the VBS schemes

This section reveals the contributions from different SOA precursors and probes into the uncertainties within the current VBS scheme in CAM6-Chem. We first investigate the seasonality and diagnose the components of simulated OA to narrow down the most important candidates causing the modeling bias. Figure 4a–c present the CAM-Chem-climo-simulated surface OA concentration with the contributions from POA and five bins of SOA (SOA1–SOA5) over CONUS, EUS, and WUS. On an annual average scale POA consists of $35.6\%$ of the total OA with a larger contribution in winter ($\sim 50\%$), while SOA dominates in summer ($\sim 77.1\%$) over CONUS. Our CAM-Chem-climo simulation suggests that the overall ambient OA concentration is determined by SOA for the CONUS domain, which is also consistent with observation-based studies (Yu et al., 2004; Zhang et al., 2018). Figure 4a shows that the absolute contribution from POA is relatively less dynamic than SOA through the year, although both show higher values in summer, suggesting that the seasonality of surface OA is primarily determined by SOA. The five volatility bins show consistent seasonality with high concentrations in warmer months, but the two least volatile bins SOA1 and SOA2 show more prominent seasonal variations, and these two bins also contribute most to total SOA. The dominant influences of SOA1 and SOA2 on total OA presented by CAM6-Chem is consistent with other VBS-based models (Farina et al., 2010).
Figure 4. (a)–(c) Seasonal variation in surface OA concentrations from CAM-Chem-climo simulation at all IMPROVE sites located over (a) CONUS, (b) EUS, and (e) WUS. The contributions of POA and SOA in five VBS bins are shown in filled areas. The 14-month average contribution of POA and SOA are shown as the number in the legend. (d)–(f) Seasonal variation of SOA formed by 15 different pathways over (d) CONUS, (e) EUS, and (f) WUS. The contribution of 15 reactions are shown in filled areas, and the 14-month average contribution is shown as the number in the legend. (g)–(i) The contributions from each pathway to the five SOA bins over (g) CONUS, (h) EUS, and (i) WUS in July 2010. The SOA concentrations in five bins are shown in filled bars.

We then examine the contribution from each of the 15 VBS reactions (listed in Table S1) to identify the most influential species over CONUS, EUS, and WUS, as shown in Fig. 4d–f. The SOA formed by each reaction is calculated as the difference between CAM-Chem-climo and the corresponding sensitivity experiment described in Table 2. The 16 sensitivity experiments (Table 2) are designed for avoiding nonlinear uncertainty of SOA formation mechanisms. For instance, SOA_{MTERP+O3} is calculated by the difference between SOA from CAM-Chem-climo scenario and SOA from “no MTERP + O3” scenario. In CAM-Chem-climo scenario, the reaction is simulated to calculate monoterpene (MTERP) oxidation by O_3 and produce SOAG (Table S1). In “no MTERP + O3” scenario, the reaction is still simulated as usual, but SOAG yield is set to zero to avoid the potential nonlinear influence of turning off the reaction. The sensitivity simulations demonstrate that dominant contributions over CONUS and EUS are from three BVOC-formed types of SOA and lumped monoterpenes (MTERP), followed by isoprene (ISOP) and sesquiterpenes (BCARY), in summer, while the intermediate volatility organic compounds (IVOCs) and aromatics (benzene, toluene, and xylenes) are mainly emitted by anthropogenic sources and play a more important role in winter. The contributions from the 15 reactions over WUS show similar results except that SOA formed by ISOP shows less of a contribution compared to SOA formed by BCARY due to less SOA_{ISOP+OH} in summer. SOA formed from aromatics also shows much smaller seasonal variation compared to BVOC-formed SOA because anthropogenic emissions are relatively stable throughout the year, while biogenic emissions are substantially more intensive during the growing season than other periods. Besides, SOA formed by MTERP dominates total SOA vertical distribution (~34.5%), as shown in Fig. S3. The sensitivity exper-
ments demonstrate that monoterpenes play the most important role in determining both surface and vertical simulated OA concentrations in summer on average over the CONUS domain.

We further diagnose that the SOA concentration in July formed from these reactions because of the prominent seasonality (larger bias in summer) and regional difference (larger bias in EUS) revealed in earlier sections. Figure 4g to i present the contributions from each pathway to the five SOA bins in July over CONUS, EUS, and WUS. Among the 15 pathways to form SOA, the reaction of MTERP + O₃ forms most SOA, followed by MTERP + OH and ISOP + OH over CONUS and EUS, while SOA formed by MTERP + OH contributes most over WUS, which is consistent with the overestimation of O₃ over EUS in MOZART-TS1 as mentioned in Emmons et al. (2020). We also find that MTERP-formed SOA is mostly in the second VBS bin SOA2 with low volatility, but most of the SOA formed by ISOP + OH is in the fourth VBS bin SOA4 with high volatility.

The contribution of monoterpenes to SOA is consistent with other measurement studies (Xu et al., 2018; Zhang et al., 2018) which demonstrated that a significant fraction of observed OA was monoterpenes-generated SOA over the southeastern US. Zhang et al. (2018) indicated monoterpenes-derived SOA contributed to ∼42% of total AMS PM₁ OA during the SOAS (Southern Oxidant and Aerosol Study) field campaign. As Sect. 3.1 reveals substantial overestimation in summer and minor bias in winter, it is thus very likely that monoterpenes-related VBS reactions may be primarily responsible for the modeling uncertainty. We have found a

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**Figure 5.** The relationship between simulated surface OA bias (y axis) and surface MTERP emissions (x axis) in December–February (DJF) (a, e, i), March–May (MAM) (b, f, j), June–August (JJA) (c, g, k), and September–November (SON) (d, h, l) over CONUS (a–d), EUS (e–h), and WUS (i–l). The correlation coefficient of OA bias and MTERP emission is shown as the number in the top right of each plot.
strong correlation between monoterpane emission and modeling bias, as shown in Fig. 5. The CAM-Chem-SD simulation bias increases alongside biogenic monoterpane emission with a correlation of 0.41 in summer and −0.12 in winter over the CONUS domain. This relationship is particularly strong (correlation of 0.67) for summer over EUS (Fig. 5g) with higher biogenic VOCs and almost negligible in winter over WUS (correlation of −0.05) with lower biogenic emission. Moreover, we also find biogenic VOC flux dominates OA bias with a higher correlation in summer over EUS, while anthropogenic VOC flux dominates OA bias over WUS, as shown in Fig. S4. Despite the fact that isoprene biogenic emission flux shows the same correlation as monoterpane biogenic emission flux over EUS, the prominent seasonality of modeling bias and the strong correlation with summertime emission suggests that monoterpane SOA dominates the performance of CAM6-Chem over abundant biogenic emission regions.

The mass yields of VBS in CAM6-Chem were parameterized following Hodzic et al. (2016) which adjusted the values used in GEOS-Chem (Jo et al., 2013) to account for the wall-loss. These parameters are shown in Table 4. The adjustment by Hodzic et al. (2016) resulted in stronger SOA production and led to overestimation in surface OA concentration compared with AMS global network and biogenic sources contributing to the total SOA production. It is certainly reasonable to take the wall-loss effect into account when making the chamber measurements. But it also should be noticed that those measurements were conducted in an artificial environment with predefined chemical species that may vary significantly from the real meteorological condition and atmospheric chemistry regime. Thus, the parameters reported in the chamber studies need to be carefully interpreted and adjusted when applied in atmospheric models. The current VBS scheme in CAM6-Chem differs from Hodzic et al. (2016) by considering the differences of oxidants but does not consider the NOx dependence of monoterpane oxidation. The mass yields applied in CAM6-Chem are generally in the middle of the low and high NOx conditions used by Hodzic et al. (2016) but still show substantial overestimation due to monoterpane SOA as mentioned above, suggesting that the mass yields in CAM6-Chem require further adjustment.

This simplified representation of VBS parameterization also affects simulation bias because BVOCs have been demonstrated to be closely influenced by NOx concentration (Shrivastava et al., 2017). Since high NOx usually suppresses the formation of monoterpane SOA, observed OA would be lower at places with high NOx concentration than those with low NOx concentration. Subsequently, the overestimation due to monoterpane SOA would be more severe at places with a high NOx level. Figure 6a to d present the simulation bias at two typical IMPROVE sites representing the high (Agua Tibia; 33.5°N, 117.0°W) and low (Lake Sugema; 40.7°N, 92.0°W) NOx conditions, respectively. With comparable levels of monoterpane emissions, simulation bias at Agua Tibia is substantially higher than that at the Lake Sugema especially during September and October when NOx concentrations are at a maximum. We further investigate the relationship between OA bias and NOx concentration at low MTERP biogenic emissions, as shown in Fig. 6e. OA bias increases with a higher NOx concentration at the same MTERP biogenic emissions in summer at 53 sites. A recent study (Jo et al., 2021) reported the development of CAM6-Chem for implementing the NOx-dependent yields, along with several other updates including SOA derived from isoprene epoxydiols (IEPOX) through heterogeneous chemistry, isoprene emission adjustment, biogenic VOC deposition, and more detailed gas-phase chemistry of isoprene. We also evaluated a simulation from CESM2.2 (CAM6-Chem configured with MOZART-TS2 which includes the above-mentioned updates except for the heterogeneous chemistry of IEPOX-SOA) against IMPROVE observations through a full year simulation (March 2013–February 2014). We find that these updates moderately lower the simulated OA concentration, but surface OA is still substantially overestimated in summer. The observed OA was 2.4 μg m⁻³ at IMPROVE sites over the CONUS domain in July 2013, while simulated OA by CESM2.1 and CESM2.2 were 5.1 and 4.2 μg m⁻³, respectively (Fig. S5). Thus, we suggest that the monoterpane SOA yield in current CESM2.1 and CESM2.2 might be con-

<table>
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<th>Log(C*)</th>
<th>Condition</th>
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<th>Hodzic et al. (2016)</th>
<th>CAM-Chem</th>
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</thead>
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<tr>
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<td>Low NOx</td>
<td>High NOx</td>
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<tr>
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<td></td>
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</tr>
</tbody>
</table>

Table 4. Mass yield coefficient of monoterpane in JO et al. (2013), Hodzic et al. (2016), and CAM-Chem in VBS bins. C* is the saturation concentration (μg m⁻³).
Figure 6. 2010 observed surface OA concentration (red dots) and simulated POA (dark filled area) and SOA (gray filled area) at the Agua Tibia site (33.5°N, 117.0°W) (a) and Lake Sugema site (40.7°N, 92.0°W) (b); simulated surface NOx concentration (blue lines) and MTERP emission from MEGAN (green lines) at the Agua Tibia site (c) and Lake Sugema site (d). The 1988–2019 summertime surface OA bias (colorful dots) at specific NOx concentration and MTERP emission flux (e) over 53 sites where MTERP emission fluxes are between $3 \times 10^{-10}$ and $6 \times 10^{-10}$.

figured too high and may lead to surface OA overestimation, and the VBS parameterization for other BVOCs in CESM may also need further adjustment for reducing OA bias.

In addition to the monoterpene parameterization, the representation of isoprene SOA by pure gas-phase VBS may also introduce critical uncertainty in the model simulation since over certain regions like the southeastern USA, isoprene SOA is mostly formed from IEPOX via irreversible multiphase chemistry that is closely affected by gas-aqueous phase transfer and acid-catalyzed reactive uptake, for which
neither has been considered in CESM2.1. A few recent studies have revealed that IEPOX-SOA is influenced by anthropogenic emissions because of the changes in inorganic aerosol acidity and sulfate particles (Shrivastava et al., 2019; Zheng et al., 2020). Both modeling and measurement evidence has suggested that the steadily decreasing anthropogenic NOx and SO2 emissions have led to the reduction of biogenic SOA over the western and southeastern US during the past decade, yet CAM-Chem-SD has difficulty in reproducing this trend, as shown in Fig. 7. The IMPROVE data show a decreasing trend of 0.527 µg m\(^{-3}\) per decade at Cape Romain National Wildlife Refuge (NWR), but the CAM-Chem-SD simulated trend is 0.098 µg m\(^{-3}\) per decade. Ridley et al. (2018) indicated that the reduction of anthropogenic emissions due to EPA regulations on vehicular sources and power generation can explain more than two-thirds of the decline in OA over CONUS from 1990 to 2012. Marais et al. (2017) showed that SO2 emissions controlled biogenic OA concentration by sulfate–BVOCs interaction in summer over EUS.

4 Summary and conclusions
The goal of this study is to understand the performance of CESM2.1 and reveal the remaining biases for the simulation of SOA. Through the validation against surface measurements and flight campaigns over the US, we have found that CESM2.1 is able to capture interannual and seasonal variation of surface OA concentrations with a correlation coefficient of 0.41, but it systematically overestimates surface OA concentration in summer by 68.78%. Larger summertime bias is found over the eastern US (overestimates by 131.15%) where BVOC emissions are more intensive than the western US. Opposite to the overestimation near the surface, consistent underestimations by −20% to 70% of OA in the upper air are found in the validation against all five flight campaigns.

Our analysis suggests that it is likely that simulated monoterpene SOA production is parameterized with yields that are too high and may be the most influential factor that affects the modeling bias for three reasons: first, monoterpene SOA contributes most (46.3%) to the total SOA in summertime, while other anthropogenic POAs or BVOCs have a smaller impact on the severe overestimation. The large contribution of monoterpene SOA simulated by CAM6-Chem is consistent with other measurement and modeling studies, but the current VBS configuration adopted from GEOS-Chem may require further adjustment. Isoprene may also play an important role in modeling uncertainty, but the influence is likely less significant than monoterpene as the isoprene-derived SOA consists of 17.0% of total SOA in summer. Second, the simulation bias showed a strong spatiotemporal correlation with monoterpene emissions as demonstrated by the large overestimation in summer over the eastern US, and larger overestimation of OA is found at places with a higher NOx condition under the same monoterpene emissions level. Third, the overestimation of OA at surface layer and the underestimation of OA and monoterpene in the free troposphere suggest that both the production and photolytic removal processes might be parameterized too strongly.

It should be noticed that our analysis of OA simulating bias mainly focused on SOA production, while other related processes such as vertical transport, removal through wet and dry depositions, and evaporation also affect the simulated OA concentration. Model treatments for these processes may also introduce uncertainties in the simulation results. The uncertainty related to VBS, however, might play a more important role than other processes according to the evaluation results shown in the above section. For example, CESM has been demonstrated to reproduce well the vertical profiles of meteorological variables (He et al., 2015). A recent sensitivity study (Gaubert et al., 2020) showed that
the systematic underestimation of CO vertical profile was mainly due to bias in anthropogenic emissions other than the vertical transport scheme. For dry and wet depositions, CESM showed slightly higher estimations than other models with the AeroCom II program, but the estimated lifetime was consistent with the multi-model mean (Tsigaridis et al., 2014). Evaporation of OA was significantly slower than the rapid reaction rates of VOCs and the oxidants (Vaden et al., 2014).\[\text{\ldots}\]

The updated VBS scheme under different NO\textsubscript{x} between near- and above-surface OA bias needs further discussed in this study were all conducted in warm months. It is also necessary to note that the five aircraft campaigns discussed in this study were all conducted in warm months. The simulated performance in winter and the discrepancy between near- and above-surface OA bias needs further investigation. The updated VBS scheme under different NO\textsubscript{x} conditions and multiphase isoprene-derived SOA chemistry have been recently considered in other global and regional models which showed better performance in SOA simulation (Hodzic et al., 2016; Shrivastava et al., 2019; Jo et al., 2021), revealing the urgent need for improving process-level representations of SOA formation and removal in models.


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\textbf{Y. Liu et al.: Community Earth System Model (CESM2.1)}

\textbf{Author contributions.} XD and YamL conceived the idea and designed the model experiments. YamL performed the simulations, conducted the analysis, and wrote the paper. MW, XD, YamL, and XL provided conceptualization for the paper. YuL, YamL, and MS provided guidelines for the simulations in CESM v2.1. YamL helped in IMPROVE data downloading and processing. XD, YamL, LEK, MS, and MW edited the paper.

\textbf{Competing interests.} The authors declare that they have no conflict of interest.

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