



## Modeling biogenic and anthropogenic secondary organic aerosol in China

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**Abstract.** A revised Community Multi-scale Air Quality (CMAQ) model with updated secondary organic aerosol (SOA) yields and a more detailed description of SOA formation from isoprene oxidation was applied to study the spatial and temporal distribution of SOA in China in the entire year of 2013. Predicted organic carbon (OC), elemental carbon and volatile organic compounds agreed favorably with observations at several urban areas, although the high OC concentrations in wintertime in Beijing were under-predicted. Predicted summer SOA was generally higher (10–15  $\mu\text{g m}^{-3}$ ) due to large contributions of isoprene (country average, 61%), although the relative importance varies in different regions. Winter SOA was slightly lower and was mostly due to emissions of alkane and aromatic compounds (51%). Contributions of monoterpene SOA was relatively constant (8–10%). Overall, biogenic SOA accounted for approximately 75% of total SOA in summer, 50–60% in au-

tumn and spring, and 24% in winter. The Sichuan Basin had the highest predicted SOA concentrations in the country in all seasons, with hourly concentrations up to 50  $\mu\text{g m}^{-3}$ . Approximately half of the SOA in all seasons was due to the traditional equilibrium partitioning of semivolatile components followed by oligomerization, while the remaining SOA was mainly due to reactive surface uptake of isoprene epoxide (5–14%), glyoxal (14–25%) and methylglyoxal (23–28%). Sensitivity analyses showed that formation of SOA from biogenic emissions was significantly enhanced due to anthropogenic emissions. Removing all anthropogenic emissions while keeping the biogenic emissions unchanged led to total SOA concentrations of less than 1  $\mu\text{g m}^{-3}$ , which suggests that manmade emissions facilitated biogenic SOA formation and controlling anthropogenic emissions would result in reduction of both anthropogenic and biogenic SOA.

## 1 Introduction

Fast economic development and rapid industrialization and urbanization in the past several decades not only significantly increased the level of air pollution in China but also lead to higher aerosol loadings in the downwind regions. A significant fraction of the aerosols is made up of organic carbonaceous material (Sun et al., 2014). Globally, organic aerosols (OA) from both biogenic and anthropogenic sources play a significant role in affecting global climate by directly scattering and absorbing solar radiation and indirectly by altering cloud and rain formation processes. It was estimated that China contributed to approximately 16 % of global organic aerosol loading in 2000 (Saikawa et al., 2009; Tsigaridis et al., 2005), and its contributions are likely higher nowadays due to increased air pollution levels in recent years. During severe haze events, it was found that secondary organic aerosol (SOA), which is formed in the atmosphere from gas-to-particle conversion of primary precursor volatile organic compounds (VOCs), could account for a significant fraction of the total observed organic aerosol (Huang et al., 2014a; Carlton et al., 2009; Tsigaridis and Kanakidou, 2003).

To better quantify the impacts of SOA on human health and climate and to design more efficient control strategies for improving air quality in China, improved understanding of SOA is urgently needed, including its precursors, formation processes and physicochemical properties. While a large amount of experimental work, both in the field (Boström et al., 2002; Aramandla et al., 2011) and in the laboratory (Lai et al., 2016), has been carried out in recent years, only a small number of modeling analyses of SOA in China have been reported. In order to determine the contributions of SOA to total organic carbon (OC) burden in China, receptor-oriented analysis with specific SOA tracers (Akagi et al., 2011), or statistical analysis with minimal elemental-carbon-to-OC (EC / OC) ratio (Peng et al., 2013; Wang et al., 2015), have been attempted. Recently, advanced techniques such as the positive matrix factorization analysis of organic aerosol data from aerosol mass spectrometers have also been applied to determine the amount of SOA and its precursors (Huang et al., 2014b; Sun et al., 2014; Wang et al., 2016)

Mechanistic modeling analyses of SOA formation in China were seldom reported in the literature until the last 5 years or so. The complexity of the models ranges from relatively simple box models (Ciccioli et al., 2014) to full-blown 3-D Eulerian models (Hellén et al., 2012; Setyan et al., 2012; Jiang et al., 2012; B. Li et al., 2015; Zhao et al., 2016). Most of the studies focused on the Pearl River Delta (PRD) region in southern China (Ciccioli et al., 2014; B. Li et al., 2015; Setyan et al., 2012) and eastern China (Hellén et al., 2012) and were for short time periods. While computationally intensive, a full-year simulation with a fully coupled meteorology–chemistry model for the entire Chinese region was also attempted (Jiang et al., 2012).

The SOA modules used in these studies applied the relatively simple traditional two-product approach (Ciccioli et al., 2014) or the more complex volatility basis set (VBS) approach to better represent multi-generation oxidation of SOA precursors in the gas phase and their aging processes in the aerosol phase (Hellén et al., 2012). The two-product approach has been shown in many previous studies to under-predict SOA. While the VBS approach and many of its variants have been shown to predict higher SOA concentrations, a recent study showed that the VBS parameters adjusted to fit individual SOA precursors in photochemical chamber experiments cannot produce satisfying results for simulating chamber experiments with mixed precursors (Zhao et al., 2015). In addition, both two-product and VBS approaches assume equilibrium partitioning of semivolatile products; however, recent studies revealed that reactions on particle surfaces or with particle water under acidic conditions of certain compounds, such as dicarbonyls (e.g. glyoxal) and epoxydiols from isoprene, lead to rapid formation of low volatile oligomers, organosulfates and organonitrates (George et al., 2015). The heterogeneous SOA formation pathway from aqueous uptake of glyoxal (GLY) has been studied for the PRD region, and it was found that including glyoxal surface uptake leads to higher isoprene SOA (iSOA) fraction (B. Li et al., 2015).

Isoprene epoxydiols (IEPOX) and methacrylic acid epoxide (MAE) formed from isoprene oxidation under low and high-NO<sub>x</sub> conditions (Lin et al., 2013; Surratt et al., 2010), respectively, have been reported to form low-volatile organic products in aerosol water under strong acidic conditions (Karambelas et al., 2014; Pye et al., 2013). Some of the recent SOA modeling studies included more detailed treatment of isoprene oxidation in the gas phase and updated SOA modules by using either a surface-controlled uptake approach (J. Li et al., 2015; Ying et al., 2015) or a more mechanistic approach for the aqueous reactions (Pye et al., 2013). It was found that IEPOX accounted for as much as 34 % of the total iSOA in summer (Ying et al., 2015). Most of these modeling studies were for North America, and whether these processes are important to SOA formation in China remains unclear.

In this study, we simulated ground-level SOA concentrations in China during the entire year of 2013 using a regional 3-D air quality model. The model includes a treatment of the isoprene gas-phase chemistry that leads to the production of IEPOX and MAE. It also includes an updated SOA mechanism with updated two-product yields and SOA formation from reactive surface uptake of dicarbonyls and IEPOX and MAE. This is the first time a regional SOA model with recent advances in iSOA treatment has been applied in China. This study provides new insights into the importance of biogenic and anthropogenic emissions in SOA formation in China under different meteorological and emission conditions, a better evaluation of the current levels of SOA concentrations, and

the contributions of anthropogenic and biogenic sources to SOA loading throughout the country.

## 2 Method

### 2.1 Model description

The Community Multiscale Air Quality (CMAQ) model (v5.0.1), developed by the United States Environmental Protection Agency (US EPA), with an updated gas-phase photochemical mechanism and a revised SOA module, was applied in this study. A complete description of the SOA module was described in greater detail in Ying et al. (2015), and a short summary is provided below. The CMAQ model was also updated to include heterogeneous reactions of NO<sub>2</sub> and SO<sub>2</sub> on particle surface to form secondary nitrate and sulfate (Ying et al., 2014).

The gas-phase photochemical mechanism applied in this study is based on the SAPRC11 (S11) photochemical mechanism (Carter and Heo, 2012), an update of the widely used SAPRC07 (S07) mechanism. The S11 mechanism was further modified to include a more detailed isoprene oxidation scheme to predict the formation of IEPOX and MAE, as originally described by Xie et al. (2013) and Lin et al. (2013). In addition, this modified S11 mechanism was designed to track GLY and methylglyoxal (MGLY) from six major groups of precursors using a tagged-species approach. The six groups of precursors are (1) aromatics with OH reaction rates less than  $2 \times 10^4 \text{ ppm}^{-1} \text{ min}^{-1}$  (ARO1), (2) aromatics with OH reaction rates greater than  $2 \times 10^4 \text{ ppm}^{-1} \text{ min}^{-1}$  (ARO2), (3) isoprene (ISOP), (4) monoterpenes (TERP), (5) sesquiterpenes (SESQ) and (6) primary emissions of GLY and MGLY.

Two major SOA formation pathways were included in the SOA module. The first is the equilibrium partitioning of semivolatile products from oxidation of long-chain alkanes (ALK5), ARO1, ARO2, ISOP, TERP and SESQ, and subsequent formation of nonvolatile oligomers based on the traditional two-product approach as implemented in version 6 of the aerosol module in CMAQ (AERO6). Oligomers from different precursors were also treated using the precursor-tagging approach so that a complete analysis of precursor contributions to SOA could be conducted. In this study, the SOA mass yields were updated from the ones used in Ying et al. (2014, 2015) to account for vapor wall loss during chamber experiments based on data provided by Zhang et al. (2014). The original and updated mass yields are listed in Table 1. No changes of the saturation concentrations ( $C^*$ ) of the products were made, even though the mass yields were updated. This partial correction only represents an upper bound estimation of the wall-loss effect. It should also be noted that while the correction factors proposed Zhang et al. (2014) for aromatic compounds by were based on the same chamber experiments that the CMAQ yields were based on, the correction factors for alpha-pinene and isoprene re-

ported by Zhang et al. (2014) were based on a different set of chamber experiments than were used to derive the yields used in CMAQ. This could lead to additional uncertainties in the yields used in this study. The second SOA formation pathway in the model is the surface-controlled irreversible reactive uptake of GLY, MGLY, IEPOX and MAE. Uptake coefficients ( $\gamma$ ) for GLY and MGLY were fixed at  $2.9 \times 10^{-3}$ , taken from Fu et al. (2008), and an acidity-dependent uptake coefficient for IEPOX and MAE was based on the parameterization described by J. Li et al. (2015). The treatment of GLY and MGLY SOA formation as an irreversible process provided an upper-limit estimation of SOA formation from these two precursors, as it was reported that reactive uptake of GLY and MGLY can be reversible (Galloway et al., 2009) and dissolved GLY and MGLY can react with oxidants to form products with volatile products (Lim et al., 2010). The resulting  $\gamma_{\text{isoepox}}$  in the summertime (June, July and August) in this study are approximately  $4\text{--}10 \times 10^{-4}$  in most areas (see Fig. S1 in the Supplement). These  $\gamma_{\text{isoepox}}$  values are in the range mentioned by Pye et al. (2013), which were based on more mechanistic calculations. The isoprene SOA yields in the CMAQ model were taken from Kroll et al. (2006), whose study was conducted under dry conditions (RH < 10 %). Thus, the yields derived from the experiments do not include contributions from reactive surface uptake, which only occurs on wet particles.

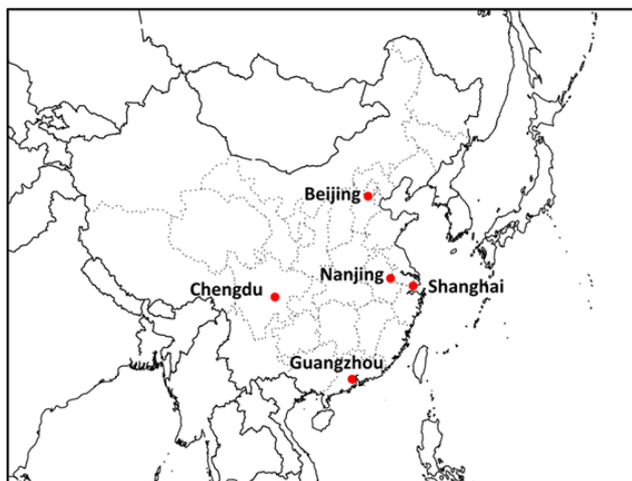
### 2.2 Model application

The modified CMAQ model was applied to simulate surface concentrations and determine precursor contributions to SOA in China for the entire year of 2013. The model domain covers China and surrounding areas in eastern and Southeast Asia, with a horizontal resolution of 36 km (see Fig. 1). The meteorology inputs needed to drive the air quality simulations were generated by the Weather Research and Forecasting (WRF) model v3.6.1 using the NCEP FNL (National Centers for Environmental Prediction Final) Operational Model Global Tropospheric Analyses dataset for boundary and initial conditions. The anthropogenic emissions in China were based on the Multi-resolution Emission Inventory for China (MEIC, v1.0) (<http://www.meicmodel.org>) with a resolution of  $0.25^\circ \times 0.25^\circ$  resolution and a matching speciation of VOCs for the SAPRC mechanism. Emissions in other countries were based on the Regional Emission inventory in Asia version 2 (REAS2) (Kurokawa et al., 2013). Biogenic emissions were generated using the Model for Emissions of Gases and Aerosols from Nature (MEGAN) v2.1 with year-specific leaf area index (LAI) data from the 8-day Moderate Resolution Imaging Spectroradiometer (MODIS) LAI product (MOD15A2). The Fire Inventory from the National Center for Atmospheric Research (FINN) was used for open biomass burning emissions (Wiedinmyer et al., 2011). Seasonal average emissions of SOA precursors (ALK5, ARO1, ARO2, ISOP, TERP and SESQ) are shown in Fig. S2. In-

**Table 1.** Updated aerosol mass yields used in this study that account for the wall-loss using scaling factors based on Zhang et al. (2014). Saturation concentrations ( $C^*$ ) for each product were also listed.

|                   | Updated    |            | Original <sup>c</sup> |            |         |          |
|-------------------|------------|------------|-----------------------|------------|---------|----------|
|                   | $\alpha_1$ | $\alpha_2$ | $\alpha_1$            | $\alpha_2$ | $C_1^*$ | $C_2^*$  |
| ALK               | 0.0865     | –          | 0.0865                | –          | 0.02    | –        |
| ARO1 <sup>a</sup> | 0.2545     | 0.7623     | 0.2253                | 0.6764     | 8.024   | 119.3166 |
| ARO2 <sup>b</sup> | 0.2545     | 0.7653     | 0.2253                | 0.6764     | 8.024   | 119.3166 |
| TERP              | 0.1811     | 0.5905     | 0.1393                | 0.4542     | 14.792  | 133.7297 |
| ISOP              | 0.0634     | 0.5104     | 0.0288                | 0.232      | 0.617   | 116.01   |
| SQT               | 1.537      | –          | 1.537                 | –          | 24.984  | –        |

<sup>a</sup>  $k_{OH} < 2 \times 10^4 \text{ ppm}^{-1} \text{ min}^{-1}$ . <sup>b</sup>  $k_{OH} \geq 2 \times 10^4 \text{ ppm}^{-1} \text{ min}^{-1}$ . <sup>c</sup> As used in Ying et al. (2015).



**Figure 1.** Location of major cities (Beijing, Nanjing, Shanghai, Chengdu and Guangzhou) mentioned in this study.

line processing was used to generate dust and sea salt emissions during CMAQ simulations. More details of the emission processing and meteorology model result evaluation can be found in J. Hu et al. (2016).

### 3 Results

#### 3.1 Model evaluation

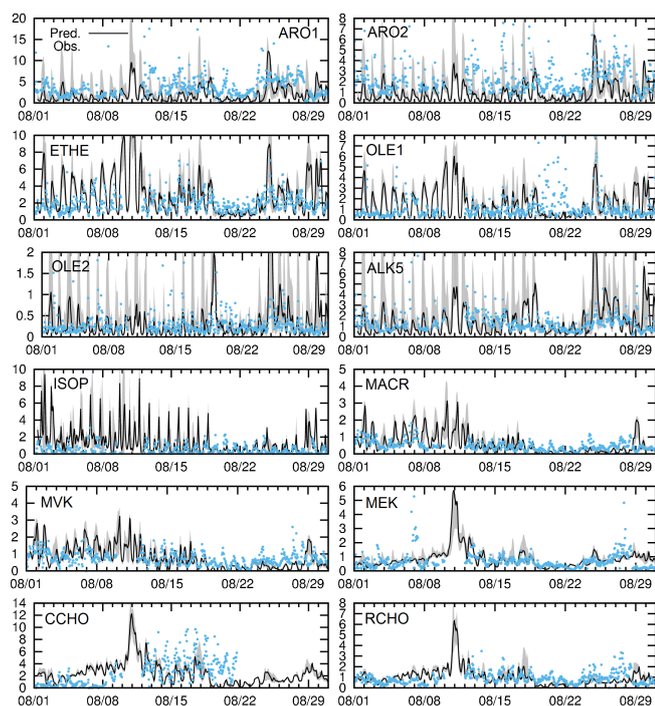
The capability of the model to predict ozone and  $\text{PM}_{2.5}$  mass concentrations has already been evaluated by comparison with observations at 60 major urban areas, and it is described in much greater detail in J. Hu et al. (2016). In summary, model performance of ozone (1 h peak and 8 h average) and  $\text{PM}_{2.5}$  mass concentrations (daily average) were considered generally within the model performance criteria recommended by the US EPA. Model performance is generally better in the more economically developed regions such as the Yangtze River Delta (YRD) and North China Plain (NCP) because of more accurate emission inventories than

the less developed regions such as the northwest. Nonetheless, as ozone and a significant portion of the  $\text{PM}_{2.5}$  mass are formed from photochemical reactions of the precursors, the general capability of the model to reproduce these species suggests that the oxidation capability of the atmosphere and precursor concentrations essential for SOA predictions were reasonably predicted – even though the SOA formation processes are much more complex and less understood.

In this section, predicted concentrations of several VOCs, as well as OC and EC (elemental carbon), are compared with observations available during this period to provide more supporting evidence for the SOA estimations.

##### 3.1.1 Volatile organic compounds

Predicted hourly concentrations of lumped primary VOCs (ARO1, ARO2 and ALK5) and ISOP, which act as SOA precursors in the S11 mechanism are compared with observations made at the Nanjing University of Information Science and Technology (NUIST) in August 2013 and shown in Fig. 2. The VOC monitoring site is located on the rooftop of the science building on campus of NUIST ( $33.205^\circ \text{N}$ ,  $118.727^\circ \text{E}$ ) and is about 15 m above the ground. Measurement of these VOC species followed the same protocol as that used at the Photochemical Air Monitoring Stations (PAMS) of the US EPA (Lonneman, 1994) (<https://www3.epa.gov/ttnamti1/pamsmain.html>). The observed concentrations of 54 detailed VOC species are used to calculate the concentrations of the lump S11 species. In addition to these SOA precursors, predicted ethene (ETHE), lumped primary olefins (OLE1 and OLE2), acetaldehyde (CCHO), and lumped higher aldehydes (RCHO), which have significant contributions to the oxidation capacity of the atmosphere, are also compared with observations. While both CCHO and RCHO can be directly emitted, in the polluted urban areas most of them are formed from oxidation of other VOCs. Also included in the comparison, were methacrolein (MACR) and methyl vinyl ketone (MVK), oxidation products from isoprene. Methyl ethyl ketone (MEK), which can be both emitted as a primary species and formed secondarily from oxida-



**Figure 2.** Predicted and observed August VOC concentrations in Nanjing. Observed concentrations are based on individual VOCs lumped into the SAPRC-11 model species. Shaded areas represent the range of predictions in the nine grid cells with the monitor site at the center. Units are ppb.

tion of a number of VOCs and their oxidation products (such as MACR and MVK), are also included in the evaluation. The predicted concentrations at the grid cell where the monitor is located are used to compare with observations. To illustrate the spatial variability of the VOC species, the ranges of the VOC concentrations in the nine grid cells, with the monitor site at the center, are also shown in the comparison.

As shown in Fig. 2, predicted anthropogenic primary emitted species (ETHE, OLE1, OLE2, ARO1, ARO2 and ALK5) have large spatial variations. This indicates that their emissions are likely much higher in urban areas but significantly lower in rural areas. Day-to-day variations of these species are generally well captured. For example, the decrease of ETHE as well as other primary species during 20–24 August and the subsequent rapid increase on 25 August are correctly predicted. This day-to-day variation is due to changes in the meteorological conditions. Emissions remain constant in the same month except for the weekday and weekend variations, and other factors that cause day-to-day variations of emissions (such as plume rise) within the same month were not considered. Observed OLE2 concentrations show erratically high concentrations during 20–22 August, which are likely due to impact of local emissions near the sampling site. For ARO1 and ARO2, the two major SOA precursors, predicted concentrations at the sampling site are slightly lower than ob-

servations (mean fractional bias, MFB =  $-0.63$  and  $-0.77$ , respectively). ALK5 concentrations are lower than aromatic compounds and show both over- and under-predictions, with an overall MFB of  $-0.05$  and a mean fractional error (MFE) of  $0.69$ . Since the olefin species are important in affecting the oxidation capacity of the atmosphere, the general agreement between model predictions and observations suggests that the initial oxidation rate of the SOA precursors can be reasonably predicted at this location.

Predicted isoprene concentrations show distinct day-to-day variations, with higher concentrations (up to approximately 8 ppb at peak hours) in the first half of August and lower in the second half (up to 1–2 ppb in the afternoon). The predicted high concentration peaks are sharp and usually last for 1–2 h only. Observed isoprene concentrations do not show significant day-to-day variations, with peak-hour concentration of approximately 2 ppb throughout the month. As isoprene is a short-lived species under typical daytime urban atmospheric conditions, the predicted high concentrations could not be due to regional transport for the large grid cell size used in this study. Variations in the predicted day-to-day concentrations must be due to isoprene emissions estimated by the MEGAN model. A previous study indicated that the MEGAN model might have overestimated the amount of isoprene emission vegetation in the urban areas due to incorrect vegetation types and fractions and leaf area index (Kota et al., 2015). However, a few other studies suggested that a significant portion of the observed isoprene was not from live vegetation but from other sources such as vehicles in urban areas (Hellén et al., 2012; McLaren et al., 1996; Borbon et al., 2001). More detailed analyses are needed to determine what triggered the rapid increase in the predicted emissions of isoprene in the afternoon.

Oxidation products of isoprene, MACR and MVK, show better agreement between predictions and observations (MFB =  $-0.14$  and  $-0.35$  and MFE =  $0.68$  and  $0.74$ , respectively). Observed concentrations of these species are higher in the first half of the month, which can cast some doubt on the isoprene measurements. However, these species are much longer lived than isoprene; therefore, transport from high-isoprene-emission regions can contribute to the higher observed concentrations. In the second half of the month, regional emissions of isoprene decreased due to changed weather conditions, and thus reduce the amount of MACR and MVK transported to the urban grid cell where the monitor was located. In summary, good agreement of these two species with observations provides a good starting point for iSOA predictions.

For the other oxidation products (CCHO, RCHO and MEK), RCHO and MEK show excellent agreement between observed and predicted concentrations (MFE =  $-0.04$  and  $0.16$  and MFE =  $0.68$  and  $0.53$ , respectively). CCHO observations are less complete during the month and show much higher concentrations than predictions from 14 to 21 August. For all three species, a large peak in the predictions on

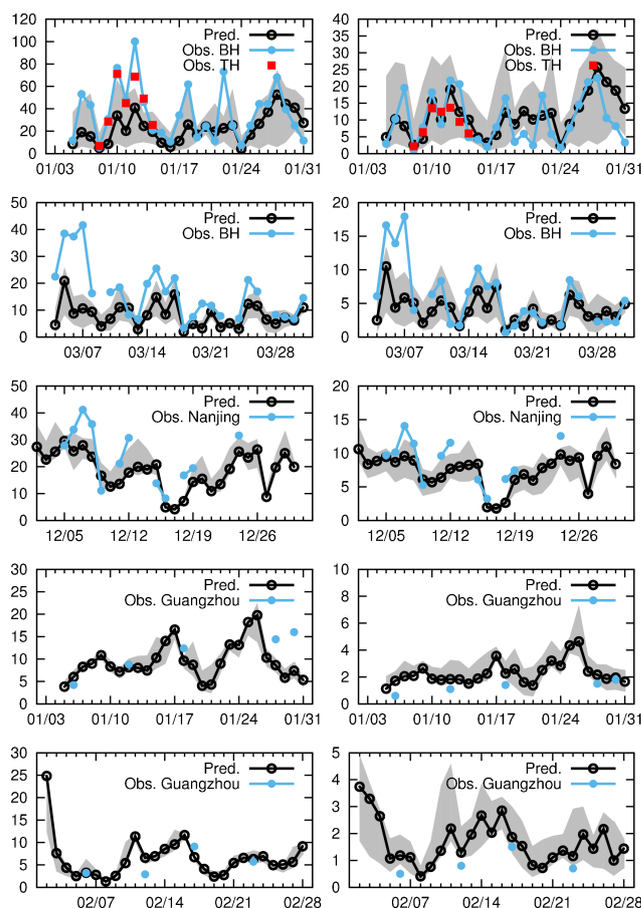
11 August was observed. However, VOC observations were not available because of an equipment problem on 10 and 11 August, making it impossible to evaluate if these predicted peaks actually occurred or not. Although this evaluation of VOC species is only for a single location in a single month due to limited observation data available during the studying period, it provides support for the VOC emissions and chemistry, at least in this region, being reasonably represented by the modeling system.

### 3.1.2 Elemental and organic carbon

Predicted EC and OC concentrations are also compared with observations, as shown in Fig. 3. EC and OC measurements are available at several locations: two locations in Beijing (on the campuses of Tsinghua University, Cao et al., 2014, and Beihang University, Wang et al., 2015; January and March 2013), one in Nanjing (Jiangsu Provincial Academy of Environmental Science, B. Li et al., 2015; December 2013), and one in Guangzhou (Tianhu, January and February 2013; Lai et al., 2016). As shown in Fig. 1, Nanjing is located in eastern China near the YRD, approximately 1000 km to the south of Beijing. Guangzhou is a megacity located in southeastern China near Hong Kong. These three cities represent very different climate and emission conditions and likely have different SOA formation pathways. Daily average concentrations were measured with different sampling intervals.

Figure 3a and b show predicted and observed OC concentrations in Beijing in January 2013, the month characterized by several high-PM pollution episodes that covered large parts of the country. The Tsinghua and Beihang campuses are 3 km apart and are located in the same model grid cell. Observation data are available at the Tsinghua site only during 8–14 January and are very similar to the observed concentrations at the Beihang site. The close agreement of the two sets of measurements during high-OC days suggests that the high concentrations (as high as  $60\text{--}80\ \mu\text{g m}^{-3}$ ) are not due to local sources very close to monitors but likely to a regional phenomenon. EC predictions agree well with observations (Fig. 2b; MFB = 0.23 and MFE = 0.48). While the model predictions of OC generally agree with the observations for day-to-day variations (MFB = -0.36, MFE = 0.53), they are significantly lower than observations on high-concentration days, except the last pollution episode in the last week of January. The good agreement of EC even on high-pollution days suggests that primary emissions and meteorological fields are accurate enough and are not the major cause of OC under-prediction. Therefore, the OC under-prediction is more likely due to under-prediction of SOA from anthropogenic sources during high-pollution days.

Figure 3c and d show that day-to-day variations in the OC and EC concentrations in Beijing during March 2013 are also well captured (MFB = -0.59 and -0.17 and MFE = 0.61 and 0.42). Spatial variations of the OC concentrations are much smaller than those in January since primary emissions



**Figure 3.** Predicted and observed concentrations of total organic carbon (OC, left column) and elemental carbon (EC, right column) in several different cities. BH: Beihang University campus; TH: Tsinghua University campus; Nanjing: Jiangsu Provincial Academy of Environmental Science; Guangzhou: Tianhu (rural site). Shaded areas represent the range of predictions in the nine grid cells with the monitor site at the center. Units are  $\mu\text{g m}^{-3}$ .

of OC from the urban areas are much reduced. OC is still more under-predicted than EC, except for the first week of March when both OC and EC concentrations are greatly under-predicted, which suggests missing primary emissions such as open burning. Figure 2e and f show that OC and EC concentrations in Nanjing during December 2013 are predicted well with slight under-prediction (MFB = -0.24 and -0.24 and MFE = 0.39 and 0.31, respectively). The day-to-day variations are also well captured, with the predicted OC and EC concentrations reaching  $30$  and  $10\ \mu\text{g m}^{-3}$  with small spatial variations. The under-prediction of OC is, at least partially, due to under-prediction in primary emissions, as both EC and OC concentrations are under-predicted. Figure 2g–j show the comparison of EC and OC in Guangzhou during January and February 2013. Although both OC and EC were in reasonable agreement with the observations on the days when the observations are available (MFB = -0.05 and 0.48

and  $MFE = 0.33$  and  $0.48$ , respectively), it is hard to conclude whether the day-to-day variations are correctly captured since observations were made every 6 days.

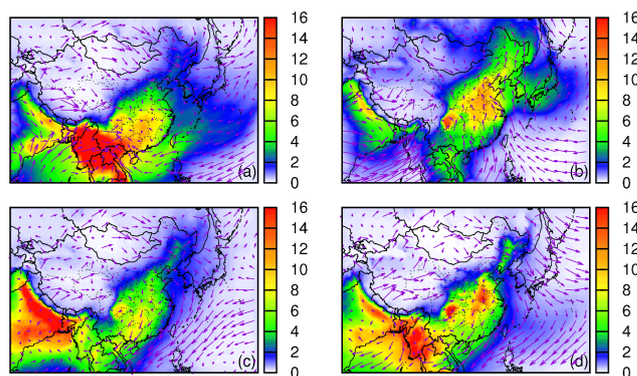
In summary, available OC and EC observations in winter and spring in three major cities suggest that emissions of EC, and thus possibly emissions of primary OC (POC), are well estimated. POC provides the medium for the partitioning of SVOCs formed from oxidation of the precursors, and correctly predicting POC is necessary for correctly predicting SOA concentrations. There are also indications that SOA is likely under-predicted, especially during the high-pollution days in winter in Beijing. While it has been suggested that VOCs with intermediate volatility (IVOCs) might also contribute to under-predictions of SOA (Zhao et al., 2016), missing other SOA precursors such as polycyclic aromatic hydrocarbons (PAHs) (Zhang and Ying, 2012) could also contribute to the OC under-prediction. Both field observations and detailed modeling studies are needed to close the gap between predicted and observed SOA concentrations.

### 3.2 Regional distribution of SOA

#### 3.2.1 Mass concentrations of SOA

Figure 4 shows predicted seasonal average SOA concentrations in (a) spring (March, April and May), (b) summer (June, July and August), (c) autumn (September, October and November), and (d) winter (December, January and February) of 2013. In spring, predicted SOA concentrations peak in Southeast Asia, which is associated with a larger amount of open burning emissions, as well as higher temperature and stronger solar radiation that enhance the photochemical production of SOA. Increase of POA could lead to additional gas-to-particle partitioning, but as shown in Sect. 3.2.2, this high SOA is mainly associated with isoprene oxidation products. Some SOA is transported to southwestern China by the southeastern monsoon, which contributes to high SOA concentrations ( $\sim 20 \mu\text{g m}^{-3}$ ) in the Yunnan and part of the Guangxi provinces. These high-SOA regions are ideal for further studies to better understand the impact of large-scale open burning on SOA and air quality. Relatively high SOA concentrations are also predicted in other southern provinces and in the Sichuan Basin, with seasonal average concentrations reaching  $\sim 10 \mu\text{g m}^{-3}$ . Predicted SOA concentrations are low in the north and northeast provinces. As indicated in Fig. S3a, SOA accounts for approximately 20–50 % of OA in the majority of the areas where SOA concentrations are high.

In summer, SOA concentrations are generally more than  $5 \mu\text{g m}^{-3}$  in a large portion of the populated areas in central and eastern China. Central provinces, such as Jiangsu, Henan, Shandong and the southern part of the Hebei in the NCP have concentrations of up to  $10\text{--}15 \mu\text{g m}^{-3}$ . This shift of high-SOA regions from southeastern to central China is accompanied by a change in the average surface wind direction from erratic wind in spring to generally southerly wind



**Figure 4.** Predicted monthly average SOA concentrations in (a) spring (March, April and May), (b) summer (June, July and August), (c) fall (September, October and November), and (d) winter (December, January and February) 2013. Units are  $\mu\text{g m}^{-3}$ . Overlain on the panels are seasonal average wind vectors.

in summer. Figure S3b shows that SOA is the major form of OA in summer, accounting for approximately 50–70 % of the OA in the densely populated areas, which is consistent with the findings in a few field measurement studies (W. Hu et al., 2016; Huang et al., 2013), and more than 80 % in most other areas.

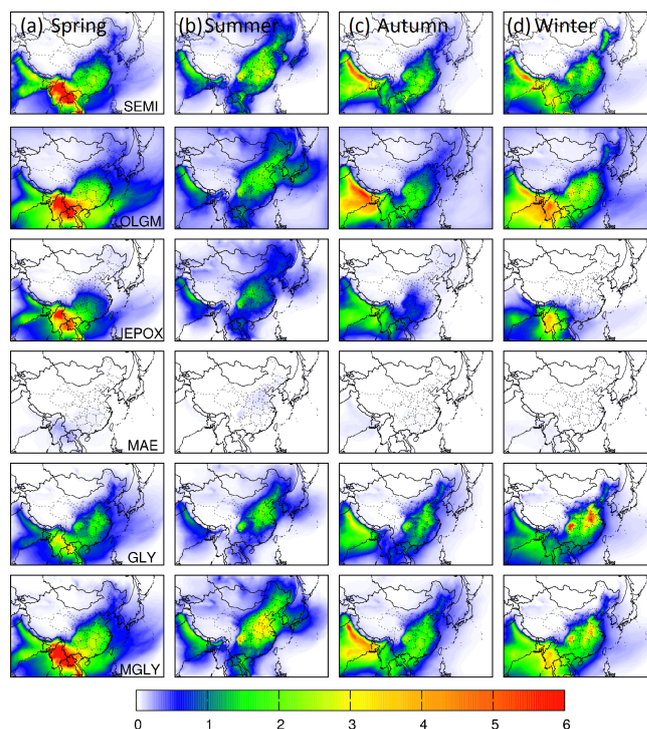
Figure 3c shows that as the wind direction changes to mostly northerly again in autumn, higher SOA concentration regions move south. The highest concentration in autumn is approximately  $8 \mu\text{g m}^{-3}$ . The spatial distribution of SOA in autumn is similar to that in spring, although lack of precursor emissions and transport of SOA from Southeast Asia leads to lower SOA concentrations in the southern border provinces. Relative contributions of SOA to total OA in autumn is approximately 30–50 %, which is also similar to that in spring, as shown in Fig. S3c. Interestingly, model predictions show that SOA concentrations in winter can still be high, with highest seasonal average concentrations reaching approximately  $20 \mu\text{g m}^{-3}$  in the Sichuan Basin and  $12\text{--}14 \mu\text{g m}^{-3}$  in Henan Province in central China. The rest of the populated areas in central and eastern China have concentrations in the range of  $5\text{--}6 \mu\text{g m}^{-3}$ . While SOA still accounts for a noticeable fraction of OA in eastern and southeastern China, its contribution to total OA is only approximately 10 % in northern and northeastern China (Fig. S3d) due to large emissions of POA from residential sources (Hu et al., 2015). A few field measurement studies also suggested that POA is more dominant in OA in winter (Huang et al., 2013; W. Hu et al., 2016; Sun et al., 2016). However, the SOA fractions of OA in these studies are higher than in this study. Although it should be noted that these field studies were taken in different years and  $\text{PM}_{10}$  were measured instead of  $\text{PM}_{2.5}$ . Predicted SOA concentrations in Xinjiang, Tibet and some other western and northwest provinces are always low due to low emissions of precursors (J. Hu et al., 2016). Averaged over the entire year of 2013, as shown in Fig. S4, it is predicted that average SOA

contribution to total OA is approximately 30 %, and the highest SOA concentration in China is in the Sichuan Basin with an annual average concentration of  $10\text{--}12\ \mu\text{g m}^{-3}$ .

### 3.2.2 SOA from traditional and reactive surface uptake pathways

Figure 5 shows the predicted seasonal average concentrations of SOA components, which represent SOA formation from different pathways: equilibrium partitioning of semivolatile SOA (SEMI), oligomerization of the condensed semivolatile SOA (OLGM), and surface irreversible uptake of IEPOX, MAE, GLY and MGLY. The SEMI and OLGM pathways are considered traditional because they are included in the default CMAQ aerosol module AERO6. The SEMI contributions to SOA were on the order of  $2\ \mu\text{g m}^{-3}$  and did not vary significantly in different seasons. The spatial distribution of the SEMI components generally follows the same distribution as the total SOA but shows more spatial heterogeneity, with higher concentrations near the source regions. This is because the primary precursors generally become oxidized relatively quickly after they are emitted. In the current SOA module, oxidation of primary precursors leads to immediate formation of two semivolatile products, which then partition to the particle phase assuming instant equilibrium. SOA aging is not considered. In contrast, the OLGM components have wider and smoother spatial distribution, as they are formed slowly from the SEMI components. In the current parameterization, the SEMI components turn into OLGM following a first-order reaction with a half-life of 20 h. The contributions of OLGM to total SOA are higher in spring and winter ( $\sim 27\%$ , see Table 2) and lower in summer and autumn (21–23 %), even though the relative contributions of the SEMI components to total SOA remain relatively constant ( $\sim 20\%$ ) in all seasons. This is likely due to more efficient removal such as wet deposition in the warmer months. Overall, the traditional pathways account for 43–47 % of the total SOA. This is higher than summer SOA simulations for the eastern US, where SOA from the traditional pathways is approximately 16 % because the contribution of isoprene to total SOA is much higher (Ying et al., 2015).

Contribution of MAE to total SOA is negligible ( $< 1\%$ ). It is known that MAE is typically formed under high- $\text{NO}_x$  conditions, but isoprene emission is generally high in rural and remote areas where  $\text{NO}_x$  emissions are low. Even though transport might occasionally mix urban plumes with sufficient isoprene, creating a high- $\text{NO}_x$  environment locally, this did not occur frequently enough to make significant contributions. SOA from IEPOX shows clear spatial and temporal distributions. IEPOX is predicted to be a major contributor to iSOA, with contributions of  $\sim 13\text{--}14\%$  in spring and lower contributions in autumn ( $\sim 9\%$ ) and winter ( $\sim 5\%$ ), averaged over the entire country. In some downwind regions, the contributions of IEPOX are much higher. GLY and MGLY have contributions from both primary emissions and sec-



**Figure 5.** Predicted seasonal average SOA component concentrations in (a) spring, (b) summer, (c) autumn and (d) winter 2011. Units are  $\mu\text{g m}^{-3}$ . SEMI: semivolatile SOA; OLGM: oligomers; IEPOX: SOA from isoprene epoxide; MAE: SOA from methacrylic acid epoxide; GLY: SOA from glyoxal; MGLY: SOA from methylglyoxal.

**Table 2.** Predicted seasonal average component contributions (%) to SOA. SEMI: semivolatile isoprene SOA; OLGM: oligomers; IEPOX: SOA from isoprene epoxide; MAE: SOA from methacrylic acid epoxide; GLY: SOA from glyoxal; MGLY: SOA from methylglyoxal.

|       | Spring | Summer | Autumn | Winter |
|-------|--------|--------|--------|--------|
| SEMI  | 19.8 % | 22.0 % | 22.1 % | 20.6 % |
| OLGM  | 26.7 % | 20.8 % | 22.7 % | 26.8 % |
| IEPOX | 13.3 % | 14.3 % | 8.6 %  | 4.5 %  |
| MAE   | 0.7 %  | 0.8 %  | 0.6 %  | 0.3 %  |
| GLY   | 14.4 % | 14.2 % | 19.4 % | 24.5 % |
| MGLY  | 25.1 % | 28.0 % | 26.7 % | 23.3 % |

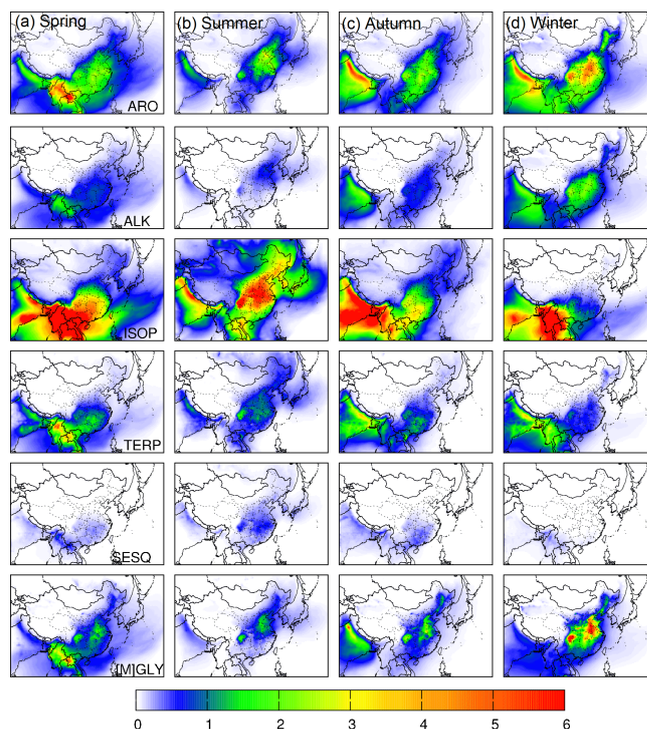
ondary formation. Spatial distributions of MGLY are wider than those of GLY, suggesting that more MGLY is due to secondary formation. Seasonal variations of GLY are also different from those of MGLY. While contributions of GLY to SOA are highest in winter ( $\sim 25\%$ ) and much lower in summer ( $\sim 14\%$ ), contributions of MGLY are relatively constant, with slightly higher contributions in summer ( $\sim 28\%$ ) and lower contributions in winter ( $\sim 23\%$ ). Emissions in winter increase significantly in China, mostly due to resi-

dential heating, suggesting the importance of anthropogenic GLY emissions in China.

### 3.2.3 SOA from different precursors

Figure 6 shows the predicted contributions to seasonal average SOA concentrations due to different types of precursor species: aromatic compounds (ARO = ARO1 + ARO2), ALK5, isoprene, TERP, SESQ, and primary glyoxal and methylglyoxal ([M]GLY = GLY + MGLY). In spring, isoprene is the largest contributor to total SOA, and it explains most of the spatial distribution of SOA shown in Fig. 4a. The peak contributions of  $4\text{--}5\ \mu\text{g m}^{-3}$  occur in the Sichuan Basin and southern China. Figure 4a indicates that some border provinces experienced cross-border transport of SOA from neighboring countries. Isoprene SOA in these near-border regions are as high as  $10\ \mu\text{g m}^{-3}$ . While it has been reported that large amounts of isoprene and other biogenic VOCs can be emitted from biomass burning (Ciccioli et al., 2014), the emission factors of isoprene vary significantly for different burning activities and fuel types (Akagi et al., 2011). Thus, the large amount of isoprene SOA predicted from open burning in this region needs to be confirmed by additional experimental studies to quantify the sources, by detailed analysis of SOA tracer compounds, for example. Averaged over the entire region in China, isoprene contributes to approximately 48 % of the total SOA in spring (see Table 3), and TERP contributes to 10 % ( $\sim 2\ \mu\text{g m}^{-3}$ ). TERP contributions are also confined in the forested regions in southern China. ARO accounts for 21 % of the total SOA in spring, and its spatial distribution is much wider, contributing a relatively uniform concentration of approximately  $2\ \mu\text{g m}^{-3}$  throughout central, eastern, southern and southeastern China.

Isoprene SOA also dominates the total SOA predicted in summer, with the highest concentrations in the Sichuan Basin. High-iSOA regions cover the NCP, the three provinces in the northeast and some provinces in east and central China. County-average contributions of iSOA account for 61 % of total SOA. Relative contribution of isoprene to total SOA is shown in Fig. S5b. Most of the areas in central China and the NCP do not have significant emissions of isoprene. With southerly summer wind, significant regional transport of iSOA from southern China contributes to the  $\text{PM}_{2.5}$  mass loading in the downwind central China and NCP areas. In contrast, SOA from TERP and SESQ does not show as much regional transport and is generally located in southeastern China where their emissions are highest (see Figs. S5c and d, S3). This is because GLY and MGLY from TERP and SESQ are much lower than from isoprene. Detailed analysis of different formation pathways to iSOA is discussed in Sect. 4.2. Together with contributions from TERP (10 %) and SESQ (4 %), the estimated contribution of biogenic SOA to total SOA in summer is approximately 75 %. Increases in the summer solar radiation and temperature facilitates the biogenic emissions. Absolute contribution of ARO to SOA in



**Figure 6.** Predicted seasonal average SOA concentrations due to contributions from different precursors in (a) spring (March, April and May), (b) summer (June, July and August), (c) autumn (September, October and November), and (d) winter (December, January and February) 2013. Units are  $\mu\text{g m}^{-3}$ . ARO: aromatics; ALK: alkane; ISOP: isoprene; TERP: monoterpenes; SESQ: sesquiterpenes; [M]GLY: primary glyoxal + methylglyoxal.

summer is approximately  $2\ \mu\text{g m}^{-3}$  in the NCP, similar to that in spring. The high fractional contribution of ARO to SOA ( $\sim 25\%$ ) is in the coastal regions in east China where the ARO emissions are high. However, the country-average relative contribution of ARO to total SOA decreases from 21 % in spring to 14 % in summer, due to significant increase in biogenic SOA,

In autumn, contributions of isoprene and monoterpene SOA decrease rapidly. The spatial distributions are similar to those in spring but without high SOA in the southern border provinces. Country-average biogenic contributions to SOA decrease to approximately 36 %. In winter, contributions of ARO increase in the Sichuan Basin and central China, reaching peak concentrations of  $4\text{--}5\ \mu\text{g m}^{-3}$ . This leads to a significant increase of ARO country-average contributions from 14 % in summer to 31 % in winter. The contribution of biogenic emissions to SOA is small in most areas, with contributions less than  $1\ \mu\text{g m}^{-3}$ . Contributions of primary GLY and MGLY increase significantly, up to  $6\ \mu\text{g m}^{-3}$  in the Sichuan Basin and in central China. The country-average contribution increases from 7 % in summer and 11–15 % in spring and autumn to 25 % in winter. The spatial distribution patterns of primary GLY and MGLY SOA do not vary much among

**Table 3.** Predicted seasonal average precursor contributions (%) to SOA. ISOP: isoprene; TERP: monoterpenes; SESQ: sesquiterpenes; ARO1 and ARO2: aromatic compounds; ALK5: long-chain alkanes; GLY: primary glyoxal; MGLY: primary methylglyoxal; and BSOA: SOA due to biogenic emissions (ISOP + TERP + SESQ; in *italic*).

|             | Spring        | Summer        | Autumn        | Winter        |
|-------------|---------------|---------------|---------------|---------------|
| ISOP        | 47.8 %        | 60.7 %        | 36.0 %        | 15.9 %        |
| TERP        | 10.0 %        | 10.3 %        | 10.8 %        | 7.4 %         |
| SESQ        | 1.6 %         | 3.6 %         | 2.6 %         | 0.5 %         |
| ARO1        | 13.2 %        | 6.8 %         | 13.3 %        | 15.2 %        |
| ARO2        | 7.8 %         | 7.6 %         | 13.4 %        | 18.7 %        |
| ALK5        | 8.2 %         | 3.9 %         | 9.5 %         | 17.5 %        |
| GLY         | 6.9 %         | 5.4 %         | 10.4 %        | 17.1 %        |
| MGLY        | 4.4 %         | 1.8 %         | 4.1 %         | 7.7 %         |
| <i>BSOA</i> | <i>59.5 %</i> | <i>74.5 %</i> | <i>49.4 %</i> | <i>23.9 %</i> |

seasons. As GLY and MGLY form SOA directly through irreversible surface uptake reactions, and there are always sufficient particles in the polluted atmosphere, this suggests that once the SOA are formed on particles, they are not transported over long distances as effectively as gas precursors and their intermediate oxidation products. The timescale of the gas-phase multistep oxidation reactions can greatly affect long-range transport and thus spatial distribution of SOA. Residential sources are the sole important contributor to total primary GLY and MGLY emissions in China (see Fig. S6). The large contributions of primary GLY and MGLY to SOA formation were not found in the previous study in the eastern US since residential emissions were not significant (Ying et al., 2015).

### 3.3 Time series in different cities

Figure 6 illustrates the time series of SOA in January and August of 2013 at four cities, i.e., Beijing in the NCP, Shanghai in the YRD, Guangzhou in the PRD and Chengdu in the Sichuan Basin. The four cities are located in the four most economically developed regions and have quite different emissions and meteorological conditions. In all cities, summer (August) SOA concentrations are higher than winter SOA concentrations (January). Chengdu has experienced the highest SOA concentrations among the four cities both in winter and summer, with the highest hourly concentrations of 35 and 50  $\mu\text{g m}^{-3}$  in January and August, respectively. Peak hourly SOA concentrations in other cities in January are approximately 20  $\mu\text{g m}^{-3}$ . In August, SOA concentrations in Shanghai and Beijing could reach approximately 40  $\mu\text{g m}^{-3}$ . Aromatic compounds are important in winter, with contributions of as much as 10  $\mu\text{g m}^{-3}$  on high-SOA days in all cities. Primary GLY and MGLY are also important in winter, especially in Chengdu. In summer, isoprene is the most important

precursor of SOA in all cities. However, the contributions of ALK, ARO and [M]GLY are not negligible.

## 4 Discussion

### 4.1 SOA from primary and secondary glyoxal and methylglyoxal

As the reversible uptake of GLY and MGLY plays an important role in SOA formation, the precursors that produce these two species are investigated using GLY and MGLY results from the precursor-tracking model. As discussed in Sect. 3.2.2, there are significant seasonal variations in the precursor contributions. Tables S1 and S2 show that isoprene is the most important secondary source of GLY and MGLY SOA in summer, on average accounting for 53 and 85 % of total GLY and MGLY SOA, respectively. This agrees well with a previous study indicating that 47 and 79 % of global gas-phase glyoxal and methylglyoxal were from isoprene oxidation (Fu et al., 2008). ARO accounts for 12 % of GLY and 11 % of MGLY SOA in summer, and its contribution increases to 31 % of GLY and 52 % of MGLY SOA in winter. Contribution of primary GLY accounts for as much as 61 % of the total GLY SOA in winter, while the contribution of secondary GLY is 65 % in summer. As described in Sect. 3.2.1 and 3.2.2, most of the MGLY is secondary, contributing from 96 % of the total MGLY SOA in summer to 74 % in winter. Liu et al. (2012) compared gas-phase GLY predictions with satellite observations and found that GLY was underestimated and an increase of aromatic emissions by 4–10 times was needed to bring the predicted GLY in line with the observations. Such a large under-estimation of aromatic compounds is not found in this study based on the observation data in Nanjing, although primary GLY is possibly underestimated. As the primary GLY emissions from residential sources have large seasonal variations, more detailed analyses of the predicted glyoxal with observations both at the surface and in space, in different seasons, are needed to further constrain the emissions of GLY and its precursors. Also note that the relative importance of MGLY in SOA formation can be affected by the uptake coefficient. While it has been demonstrated in flow tube measurements that the MGLY uptake coefficient on acidic particles is on the same order of magnitude as the uptake coefficient of glyoxal (Zhao et al., 2006), a recent study applied an uptake coefficient of MGLY that was 4 orders of magnitude lower than that of GLY, based simply on the relative magnitude of the Henry's law coefficient (Marais et al., 2016), and concluded that contributions of MGLY to SOA are negligible. More studies are needed to improve the understanding of the role of MGLY in SOA formation.

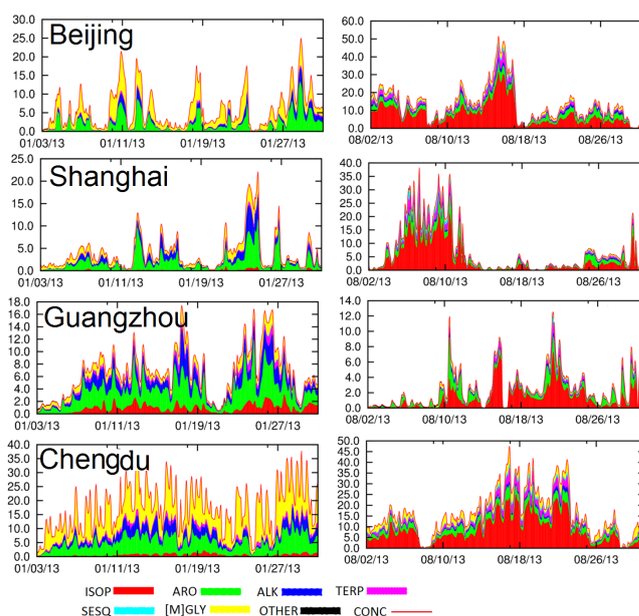
## 4.2 Isoprene SOA formation pathways

Analyses in Sect. 3 show that contribution of isoprene to total SOA is high in spring and summer. A more detailed analysis of the iSOA was conducted to reveal the contributions of components from different pathways to the predicted summer iSOA concentrations, as shown in Fig. 8. Averaged over the entire country, SEMI and OLGGM components account for only 27 % of the predicted iSOA. The rest of the iSOA is due to GLY (10 %), MGLY (35 %), IEPOX (27 %) and MAE (1 %). Spatial distributions of these components are shown in Fig. 7b–e. The SEMI component is high in the Sichuan Basin and central and eastern China, close to the forest regions of southern China with high isoprene emissions. The other components show a generally similar spatial pattern with high concentrations in the Sichuan Basin, NCP, and YRD regions.

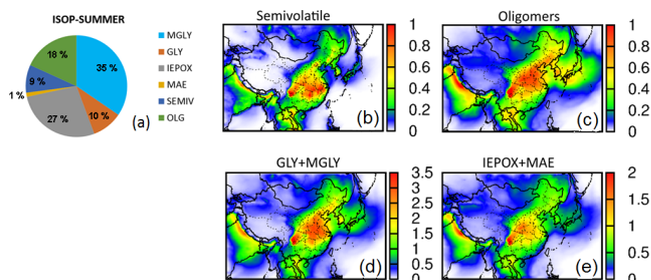
## 4.3 Sensitivity of predicted SOA to isoprene–NO<sub>x</sub>–VOC emission changes

The formation of SOA is affected by both biogenic and anthropogenic emissions. To evaluate the changes in SOA concentrations with respect to changes in anthropogenic and biogenic emissions, multiple sensitivity simulations for SOA in January and August 2013 were conducted, as summarized in Table 4. The 50 % reduction in NO<sub>x</sub> emissions and 25 % reduction in VOC emissions used in simulations 2–4 represent a rough estimation of the emission reduction target for the year 2020 based on data from Chatani et al. (2014) for a stringent emission control scenario. In simulation 5, biogenic emissions were reduced by 50 %, while anthropogenic emissions remained unchanged, as some studies showed that the MEGAN model might overestimate biogenic emissions (Carlton and Baker, 2011). This was used to evaluate the sensitivity of SOA predictions for biogenic emissions. Recently, studies of biogenic SOA in the southeastern United States suggested that SOA from isoprene could be enhanced by anthropogenic emissions; thus, reducing anthropogenic emissions might have extra benefit in reducing both biogenic and anthropogenic SOA (Carlton et al., 2010). Simulation 6 was designed to test the magnitude of this interaction in China by removing all anthropogenic emissions.

Figures 9b and 10b show that reducing NO<sub>x</sub> emissions alone by 50 % would result in both an increase and decrease in predicted SOA. In both seasons, a decrease of SOA by approximately 1 μg m<sup>-3</sup> occurs in the south and in the Sichuan Basin, while an increase in SOA of 1–2 μg m<sup>-3</sup> occurs in the Beijing–Tianjin–Hebei (BTH) and YRD regions. However, the causes of the changes are different. Analysis of the SOA components in August showed that the increase of SOA in the domain is mostly due to an increase in the SOA from the IEPOX pathway (Fig. S7). The decrease of SOA is mostly due to reduction in the SOA from GLY and MGLY, which are mainly because of reduction of GLY and MGLY from isoprene (compare Figs. S7 and S8). In January, the increase



**Figure 7.** Predicted time series of SOA concentrations (secondary y axis, units are μg m<sup>-3</sup>) and fractional contributions to SOA due to different precursors at four cities (Beijing, Shanghai, Guangzhou and Chengdu) in January and August 2013. Units are μg m<sup>-3</sup>. ARO: aromatics; ISOP: isoprene; TERP: monoterpene; SESQ: sesquiterpene; [M]GLY: glyoxal + methylglyoxal directly emitted or from other precursors; OTHER: boundary conditions.



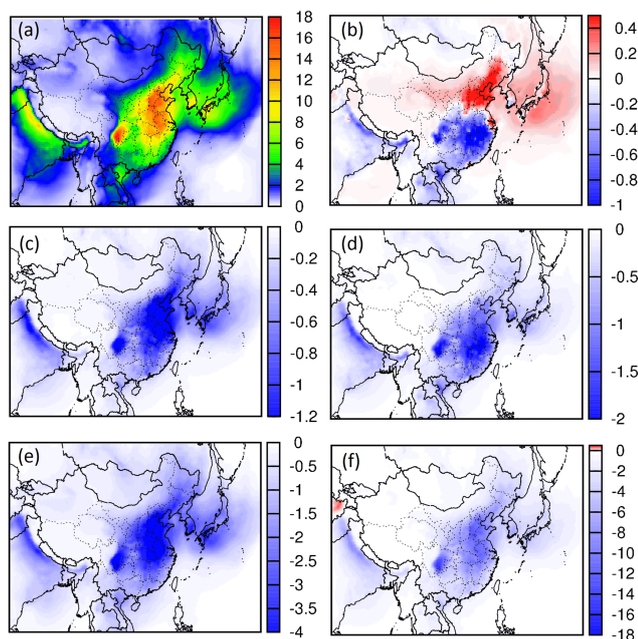
**Figure 8.** Predicted fractional component contributions to summer isoprene SOA in China (a) and spatial distributions of iSOA from the traditional (b, c) and surface uptake (d, e) pathways. Units are μg m<sup>-3</sup> for panels (b–d).

and decrease in the SOA are due to changes in the SOA from ARO and ALK5 (Fig. S9d, e). SOA from primary GLY and MGLY (Fig. S9f) also showed similar spatial patterns. In the current mechanism, ARO, ALK5 and GLY–MGLY only react with OH to form semivolatile SOA and oligomers. The results suggested that the spatial pattern change of SOA in winter due to NO<sub>x</sub> is likely due to an increase in the OH concentrations associated with NO<sub>x</sub> reduction in the high-NO<sub>x</sub> emission regions. Increase in SOA also occurs in regions downwind of the megacities due to regional transport, even though these downwind regions may not be NO<sub>x</sub>-saturated based on local emissions.

**Table 4.** Settings for sensitivity simulations.

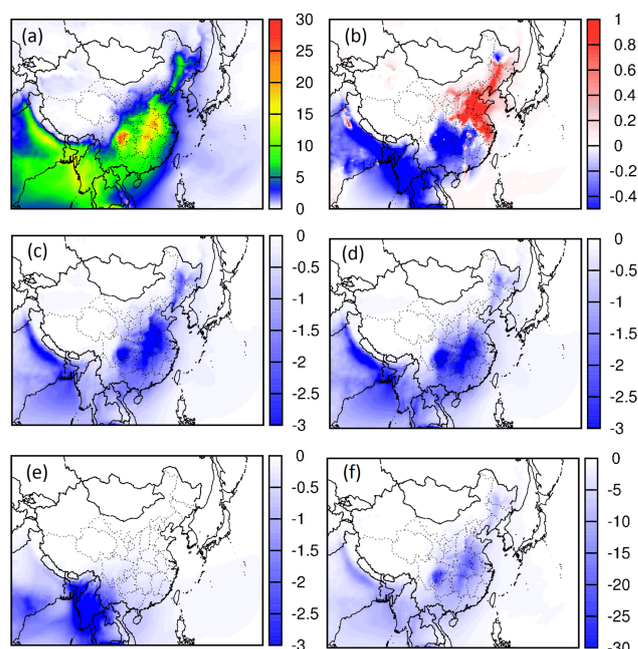
| Simulation | Abbreviation       | Notes  |
|------------|--------------------|--|
| 1          | Base               | Base case simulation, all biogenic and anthropogenic emissions at 100 %    |
| 2          | 0.5NO <sub>x</sub> | Anthropogenic NO <sub>x</sub> reduced by 50 %                              |
| 3          | 0.75VOC            | Anthropogenic VOC reduced by 25 %  |
| 4          | NV                 | Anthropogenic NO <sub>x</sub> and VOC reduced by 50 and 25 %, respectively |
| 5          | 0.5ISOP            | Biogenic isoprene emissions reduced by 50 %                                |
| 6          | BIO                | Biogenic emissions at 100 %, no anthropogenic emissions*                   |

\* All gas and particulate emissions were excluded, not only NO<sub>x</sub> and VOC emissions.



**Figure 9.** Predicted (a) spatial distribution of SOA for August 2013 and change of SOA concentrations (sensitivity case – base case) due to (b) reduction of anthropogenic NO<sub>x</sub> by 50 %, (c) reduction of anthropogenic VOC by 25 %, (d) simultaneous reduction of anthropogenic NO<sub>x</sub> and VOC by 50 and 25 %, (e) reduction of biogenic isoprene by 50 %, and (f) reduction of all anthropogenic emissions.

As shown in Fig. 9c, reducing anthropogenic VOC emissions by 25 % in August resulted in a decrease of SOA throughout the domain by 1–1.4 μg m<sup>-3</sup>, or a relative change of approximately 10 %. In January, 25 % reduction of VOCs led to an approximately 20 % reduction of the predicted SOA concentrations, or up to 3 μg m<sup>-3</sup> (Fig. 10c). This higher amount of reduction is expected because most of the SOA in winter is due to anthropogenic emissions. Figure 8d shows that simultaneous reduction of NO<sub>x</sub> and VOC in August is more effective in reducing SOA concentrations than NO<sub>x</sub> or VOC emission reduction alone. However, Fig. 9d shows that in January reducing both led to slightly less SOA reduction than reducing VOC alone. As shown in Fig. 9e, decreasing isoprene by 50 % resulted in a decrease of total SOA by



**Figure 10.** Same as Fig. 8, but for January 2013.

as much as 4 μg m<sup>-3</sup> (or 35 %) in August. Analysis of the SOA components shows that both iSOA and SOA from other precursors are decreased. This result highlights that the accuracy of the estimation of the biogenic isoprene and other biogenic VOCs can greatly affect the model predictions of iSOA. Given the high sensitivity of predicted SOA and iSOA to isoprene emissions, more studies should be carried out to evaluate, constrain and refine biogenic emission estimations in China.

Removing all anthropogenic emissions while keeping the biogenic emissions unchanged (including soil NO<sub>x</sub>) leads to very low SOA concentrations, even in the summer. Most areas are predicted to have SOA concentrations less than 2 μg m<sup>-3</sup> in summer and less than 1 μg m<sup>-3</sup> in winter. Since the reported dependence of iSOA formation on particulate nitrate and sulfate (Xu et al., 2014) was not included in the current study, the significant reduction of iSOA in the current study is due to reduced oxidation capacity of the at-

mosphere (i.e. lower OH during the day and NO<sub>3</sub> radical at night), which leads to slower formation of semivolatile components and oligomers as well as lower acidity of the aerosols that reduce the uptake coefficient of IEPOX and thus SOA. Nonetheless, the results from sensitivity simulation 6 clearly demonstrate that human-related emissions are responsible for almost all SOA. Dividing the SOA concentrations into biogenic and anthropogenic portions by the carbon source alone will erroneously underestimate the impact of manmade emissions on aerosol loadings (Setyan et al., 2012, 2014).

## 5 Conclusions

In China, predicted SOA concentrations are generally higher in summer (10–15  $\mu\text{g m}^{-3}$ ) due to large contributions of isoprene (country average, 61 %) and lower in wintertime due to emissions of alkane and aromatic compounds (51 %). Overall, 75 % of total SOA in summer, 50–60 % in autumn and spring, and approximately 24 % in winter are due to biogenic SOA. The highest SOA in mainland China occurs in the Sichuan Basin in all seasons, with hourly concentrations as high as 50  $\mu\text{g m}^{-3}$ . Regarding the SOA formation pathways, approximately half of the SOA in all seasons is due to the traditional equilibrium partitioning of semivolatile components followed by oligomerization, while the remaining is mainly due to reactive surface uptake of isoprene epoxide (IEPOX, 5–14 %), glyoxal (GLY, 14–25 %) and methylglyoxal (MGLY, 23–28 %). Manmade emissions facilitate biogenic SOA formation, and controlling anthropogenic emissions would result in reduction of both anthropogenic and biogenic SOA. It should be noted that the reported SOA concentrations in this study have not been compared with direct measurements of SOA. More detailed measurements of organic components within the aerosol phase, such as the oxidation state measurements from aerosol mass spectrometers (AMS) (W. Hu et al., 2016) and tracer species representing SOA formation from precursor species (Yang et al., 2016), are becoming available in many areas in China. Future local-scale SOA modeling studies can be conducted to better utilize these data as constraints on model parameters and input data.

## 6 Data availability

Data used in this manuscript can be provided upon request by email to the corresponding author Qi Ying (qying@civil.tamu.edu).

**The Supplement related to this article is available online at doi:10.5194/acp-17-77-2017-supplement.**

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