General comments

This paper develops a high-resolution emission inventory of IS-VOCs over China and simulates organics over the Yangtze River Delta (YRD) region using the CMAQ model. The influence of difference IS-VOC sources are investigated. The paper is well written and interesting as the influence of the different sources of organics are carefully studied. However, the novelties of this study should be better highlighted, probably by adding a section in the introduction about previous work on IS-VOC emissions in China and aboard. Also, this paper seems similar to the paper of Li et al. Environmental Pollution 2022 (see reference in specific comments below). How do the set up and findings compare to the paper of Li et al. (2022)? I suppose the IS-VOC inventories are done differently, but this should be explained.

Re: Thanks a lot for the reviewer's valuable comment. We are deeply regret that we are not able to learn from the work of Li et al. (2022) since we just submitted our work when they published the study. We have carefully read the paper of Li et al. (2022), which have done a good job simulating SOA formation and source contributions in summer China. However, we believe our study still differs from Li et al. (2022) in the following three respects. First, as the reviewer has mentioned, our study developed a high-resolution I/SVOCs emission inventory by scaling the I/SVOCs emissions from VOCs or POA emissions (depending on their dominant pollutants) and determining their volatility bins based on previous measurements of specific sources. Compared with the unified POA emission scaling method adopted by Li et al. (2022), our inventory provided more detailed I/SVOC emissions and volatility distributions of each specific source. Regrettably, the paper of Li et al. (2022) didn't provide their I/SVOCs emissions so we could not make any comparison with them. Second, in Li et al. (2022), each S/IVOC surrogate is oxidized by OH, with the saturation vapor pressure reduced by one order of magnitude. In our paper, I/SVOCs undergo OH oxidation with four products, and the mass yields are derived based on chamber experiments for I/SVOCs emitted from mobile sources (see Lu et al., 2021). Third, our modeling results refined the SOA contribution of specific sources, which is of great significance for the subsequent control of organic aerosols, but was not covered in the paper of Li et al. (2022). In addition, due to the differences in the treatment of I/SVOCs emissions inventories, the contribution of I/SVOCs emissions to SOA in our study is also very different. In their study, S/I-SOA accounted for only 8.8%, while in our study it can reach 41% in summer. Of course, we will make specific explanations and supplements on these differences in the following reply. Based on the reviewer's comments, we have added a section in the introduction about previous work on I/SVOC emissions in China and aboard.

Changes in manuscript:

1. Introduction:

I/SVOC emission inventories have been developed and applied into air quality models over the past decade. Most of them were estimated by applying different scaling factors based on their relationship with POA, volatile organic compounds (VOCs), or some proxies like naphthalene (Pye and Seinfeld, 2010; Shrivastava et al., 2011; Jathar et al., 2017; Wu et al., 2019, 2021; Li et al., 2020, 2022; Ling et al., 2022). Yet in practice, a same scaling factor was applied to most of the sources in previous studies due to the lack of measurements on I/SVOC emission factors. For example, except

biomass burning (0.75–1.5), Wu et al. (2019) utilized scaling factors of 8–30 for all of the other emission source categories, which was estimated based on the measurements of on-road mobile source. Li et al. (2020) assumed scaling factors of 1.5 for on-road mobile source, and 0.34–1.5 for the other sources, such as industrial and residential sources, which were much lower than the estimations in Wu et al. (2020). Huang et al. (2021) have tried emission factor method to quantify the I/SVOC emissions, yet the results were 60% lower than the scaling factor method, far from catching the measured amount of SOA. Obviously, roughly estimating I/SVOC emissions using one or two emission profiles as surrogates for all emission sources will create large uncertainties.

Recent studies have successively determined the volatility distribution, chemical composition, and emission factors of I/SVOCs from mobile sources, including gasoline and diesel vehicles, non-road diesel machinery, marine vessel, and aircraft (Presto et al., 2011; Cross et al., 2013; Zhao et al., 2015, 2016b; Huang et al., 2018; Qi et al., 2019; Drozd et al., 2019). I/SVOC emission profiles have been reported for nonmobile-sources as well, including coal combustion, wood-burning, cooking, fuel evaporation, and industrial and residential volatile chemical products (Huffman et al., 2009; Gentner et al., 2012; May et al., 2013; Koss et al., 2018; McDonald et al., 2018; Cai et al., 2019; Drozd et al., 2021), making the quantification of I/SVOC emissions and their involvement in air quality models possible.

In China, SOA has been emerging as an important contributor to air pollution. Field observations reveal that OA dominates (30%) the PM_{2.5} concentrations in most parts of China (Tao et al., 2017; Liu et al., 2018), among which the SOA contributes up to 80% of OA during haze pollution (Huang et al., 2014; Ming et al., 2017; Li et al., 2021). SOA formation in China has already been examined in several modeling studies. They found that by considering the POA aging and I/SVOCs oxidation in the models, which is realized by the coupling of VBS scheme, the formation and evolution of SOA can be much better simulated compared to the results of the two-product SOA modeling framework (Zhao et al., 2016a; Wu et al., 2019; Li et al., 2020; Yao et al., 2020; Huang et al., 2021). Chang et al. (2022) developed a full-volatility organic emission inventory with source-specific I/SVOC emission profiles for China, which have greatly improved the model performance on SOA concentrations. However, large gaps still exist between the observed and modeled SOA. Studies on high-resolution I/SVOC emission inventory for more specific sources are highly needed.

New references:

- Chang, X., Zhao, B., Zheng, H., Wang, S., Cai, S., Guo, F., Gui, P., Huang, G., Wu, D., Han, L., Xing, J., Man, H., Hu, R., Liang, C., Xu, Q., Qiu, X., Ding, D., Liu, K., Han, R., Robinson, A. L., and Donahue, N. M.: Full-volatility emission framework corrects missing and underestimated secondary organic aerosol sources, One Earth, 5, 403–412, 2022.
- Jathar, S. H., Woody, M., Pye, H. O. T., Baker, K. R., and Robinson, A. L.: Chemical transport model simulations of organic aerosol in southern California: model evaluation and gasoline and diesel source contributions, Atmos. Chem. Phys., 17, 4305–4318, 2017.
- Li, J., Han, Z., Wu, J., Tao, J., Li, J., Sun, Y., Liang, L., Liang, M., and Wang, Q.: Secondary organic aerosol formation and source contributions over east China in summertime, Environ. Pollut., 306, 119383, 2022.
- Ling, Z., Wu, L., Wang, Y., Shao, M., Wang, X., and Huang, W.: Roles of semivolatile and intermediate-volatility organic compounds in secondary organic aerosol formation and its implication: A review, J. Environ. Sci., 114, 259–285, 2022.

Pye, H. O. T., Seinfeld, J. H.: A global perspective on aerosol from low-volatility organic compounds, Atmos. Chem. Phys., 10, 4377–4401, 2010.

Shrivastava, M., Fast, J., Easter, R., Gustafson, W. I., Zaveri, R. A., Jimenez, J. L., Saide, P., and Hodzic, A.: Modeling organic aerosols in a megacity: comparison of simple and complex representations of the volatility basis set approach, Atmos. Chem. Phys., 11, 6639–6662, 2011.

Specific comments:

Introduction

•L77 « SOA concentration is substantially lower than that measured in the atmosphere ». This is not always the case with the VBS method, and simulated SOA tends even to be too high, see Lane et al. (2008) for example.

Lane, T. E., Donahue, N. M., and Pandis, S. N.: Simulating secondary organic aerosol formation using the volatility basis-set approach in a chemical transport model, Atmos. Environ., 42, 7439–7451, https://doi.org/10.1016/j.atmosenv.2008.06.026, 2008.

Re: Thanks for the comments. This statement is indeed inappropriate. We have rewritten this sentence to be: "the model simulated SOA concentration still has large gaps with that measured in the atmosphere"

•L78-81 Another shortcoming in the modelling of OA with the VBS 1D is the lack of representation of the hydrophilic properties of OA. VBS 1D assumes SOA condenses onto an organic phase, whereas SOA may also condense on an aqueous phase, see Kim et al (2011) for example

Kim Y., Couvidat F., Sartelet K. and Seigneur C. (2011), Comparison of different gas-phase mechanisms and aerosol modules for simulating particulate matter formation. J. Air Waste Manage. Assoc, 61, 1218-1226, doi:10.1080/10473289.2011.603999.

Re: Thanks for the comments. We have supplemented this statement in the introduction as follows. "Another shortcoming in the modelling of OA with the VBS 1D is the lack of representation of the hydrophilic properties of OA, which assumes SOA condenses onto an organic phase, whereas SOA may also condense on an aqueous phase (Kim et al., 2011)"

New references:

Kim, Y., Couvidat, F., Sartelet, K., and Seigneur, C.: Comparison of different gas-phase mechanisms and aerosol modules for simulating particulate matter formation, J. Air Waste Manage., 61, 1218–1226, 2011.

•L88: please define IVOC, SVOC. How are they quantify? Which range of volatility?

Re: Thanks for reminding. We have supplemented the definition of IVOCs and SVOCs in the revised manuscript as follows. "IVOCs refer to organic compounds with effective saturation

concentrations between 10^3 to 10^6 µg·m⁻³ at 298 K and 1 atm, while SVOCs refer to organic compounds with effective saturation concentrations between 10^{-1} to 10^3 µg·m⁻³ at 298 K and 1 atm."

•L128: « I/SVOC emission profiles have not been taken into account in previous studies. » This is not correct, they do have been taken into account in numerous studies. See the review of Ling et al. (2022) and other papers below.

Zhenhao Ling, Liqing Wu, Yonghong Wang, Min Shao, Xuemei Wang, Weiwen Huang, Roles of semivolatile and intermediate-volatility organic compounds in secondary organic aerosol formation and its implication: A review, Journal of Environmental Sciences, Volume 114, 2022, Pages 259-285, https://doi.org/10.1016/j.jes.2021.08.055.

Over the States: Jathar, S. H., Woody, M., Pye, H. O. T., Baker, K. R., and Robinson, A. L.: Chemical transport model simulations of organic aerosol in southern California: model evaluation and gasoline and diesel source contributions, Atmos. Chem. Phys., 17, 4305–4318, https://doi.org/10.5194/acp-17-4305-2017, 2017.

Over Europe: K. Sartelet, S. Zhu, S. Moukhtar, M. André, J.M. André, V. Gros, O. Favez, A. Brasseur, M. Redaelli, Emission of intermediate, semi and low volatile organic compounds from traffic and their impact on secondary organic aerosol concentrations over Greater Paris, Atmospheric Environment, Volume 180, 2018, Pages 126-137, https://doi.org/10.1016/j.atmosenv.2018.02.031.

Over China: Jie Li, Zhiwei Han, Jian Wu, Jun Tao, Jiawei Li, Yele Sun, Lin Liang, Mingjie Liang, Qin'geng Wang, Secondary organic aerosol formation and source contributions over east China in summertime, Environmental Pollution, Volume 306, 2022, 119383, https://doi.org/10.1016/j.envpol.2022.119383.

Re: Thanks for the comments. We have deleted this statement and included the reviews of previous work on I/SVOC emission profiles in the revised manuscript.

Changes in manuscript:

1. Introduction:

I/SVOC emission inventories have been developed and applied into air quality models over the past decade. Most of them were estimated by applying different scaling factors based on their relationship with POA, volatile organic compounds (VOCs), or some proxies like naphthalene (Pye and Seinfeld, 2010; Shrivastava et al., 2011; Jathar et al., 2017; Wu et al., 2019, 2021; Li et al., 2020, 2022; Ling et al., 2022). Yet in practice, a same scaling factor was applied to most of the sources in previous studies due to the lack of measurements on I/SVOC emission factors. For example, except biomass burning (0.75–1.5), Wu et al. (2019) utilized scaling factors of 8–30 for all of the other emission source categories, which was estimated based on the measurements of on-road mobile source. Li et al. (2020) assumed scaling factors of 1.5 for on-road mobile source, and 0.34–1.5 for the other sources, such as industrial and residential sources, which were much lower than the estimations in Wu et al. (2020). Huang et al. (2021) have tried emission factor method to quantify the I/SVOC emissions, yet the results were 60% lower than the scaling factor method, far from catching the measured amount of SOA. Obviously, roughly estimating I/SVOC emissions using one or two emission profiles as surrogates for all emission sources will create large uncertainties.

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In China, SOA has been emerging as an important contributor to air pollution. Field observations reveal that OA dominates (30%) the PM_{2.5} concentrations in most parts of China (Tao et al., 2017; Liu et al., 2018), among which the SOA contributes up to 80% of OA during haze pollution (Huang et al., 2014; Ming et al., 2017; Li et al., 2021). SOA formation in China has already been examined in several modeling studies. They found that by considering the POA aging and I/SVOCs oxidation in the models, which is realized by the coupling of VBS scheme, the formation and evolution of SOA can be much better simulated compared to the results of the two-product SOA modeling framework (Zhao et al., 2016a; Wu et al., 2019; Li et al., 2020; Yao et al., 2020; Huang et al., 2021). Chang et al. (2022) developed a full-volatility organic emission inventory with source-specific I/SVOC emission profiles for China, which have greatly improved the model performance on SOA concentrations. However, large gaps still exist between the observed and modeled SOA. Studies on high-resolution I/SVOC emission inventory for more specific sources are highly needed.

In this study, taking the Yangtze River Delta (YRD) region, including Jiangsu, Zhejiang, Anhui provinces and Shanghai city, as a pilot, we established a high-resolution source specific I/SVOC emission inventory. We then applied the newly established inventory into CMAQ v5.3 to evaluate the contributions of I/SVOC emissions to SOA formation by comparing the results with the observation data collected in the region. Furthermore, we also run the model in different scenarios to quantify the seasonal contributions of different sources to POA and SOA formation in the YRD region.

New references:

- Chang, X., Zhao, B., Zheng, H., Wang, S., Cai, S., Guo, F., Gui, P., Huang, G., Wu, D., Han, L., Xing, J., Man, H., Hu, R., Liang, C., Xu, Q., Qiu, X., Ding, D., Liu, K., Han, R., Robinson, A. L., and Donahue, N. M.: Full-volatility emission framework corrects missing and underestimated secondary organic aerosol sources, One Earth, 5, 403–412, 2022.
- Jathar, S. H., Woody, M., Pye, H. O. T., Baker, K. R., and Robinson, A. L.: Chemical transport model simulations of organic aerosol in southern California: model evaluation and gasoline and diesel source contributions, Atmos. Chem. Phys., 17, 4305–4318, 2017.
- Li, J., Han, Z., Wu, J., Tao, J., Li, J., Sun, Y., Liang, L., Liang, M., and Wang, Q.: Secondary organic aerosol formation and source contributions over east China in summertime, Environ. Pollut., 306, 119383, 2022.
- Ling, Z., Wu, L., Wang, Y., Shao, M., Wang, X., and Huang, W.: Roles of semivolatile and intermediate-volatility organic compounds in secondary organic aerosol formation and its implication: A review, J. Environ. Sci., 114, 259–285, 2022.
- Pye, H. O. T., Seinfeld, J. H.: A global perspective on aerosol from low-volatility organic

compounds, Atmos. Chem. Phys., 10, 4377–4401, 2010.

Shrivastava, M., Fast, J., Easter, R., Gustafson, W. I., Zaveri, R. A., Jimenez, J. L., Saide, P., and Hodzic, A.: Modeling organic aerosols in a megacity: comparison of simple and complex representations of the volatility basis set approach, Atmos. Chem. Phys., 11, 6639–6662, 2011.

Materials and methods

The authors state that they establish a new IS-VOC emission inventory. However, there is only little detail about it. More details would be welcomed.

Re: Thanks. We agree with the reviewer that "new" should be used with caution. We have removed relevant terms in the revised manuscript. Then in the "Materials and methods" section, we have added more details in I/SVOC emission estimation.

Changes in manuscript:

2. Materials and methods:

2.1 I/SVOC emission inventory

I/SVOCs commonly exist in both gas- and particle-phase in the atmosphere. Previous studies usually used POA scaling factors to estimate the I/SVOC emissions, which may lead to large uncertainties in the estimation of gas-phase organic compound-dominated sources, like oil refinery, chemical production, and industrial solvent-use. Herein, we compiled both gas-phase I/SVOCs (I/SVOCs-G) and particle-phase I/SVOCs (I/SVOCs-P) emission inventories and incorporate them into the model. Detailed process of the inventories is as follows.

- (1) Source classification: To refine the I/SVOC emissions from different sources, we divided the sources into five major categories and then further grouped them into 21 sub-categories. The major categories include industrial process sources, industrial solvent-use sources, mobile sources, residential sources, and agricultural sources. As shown in Table S1, the industrial process sources include the sectors such as oil refinery, chemical production, and pulp and paper production; Industrial solvent-use sources include textile, leather tanning, timber processing, and various industrial volatile chemical products use; Mobile sources include gasoline and diesel vehicle emissions, fuel evaporation, diesel machinery, marine vessel, and aircraft; Residential sources include coal combustion, residential solvent-use, and cooking emissions; Agricultural source is specifically referred to biomass burning in household stoves, and open burning was not included in this study.
- (2) Emission estimation: I/SVOCs-G emissions for each specific source were estimated by the ratios of total I/SVOC components to anthropogenic VOC (AVOC) components (I/SVOCs-to-VOCs). Similarly, I/SVOCs-P emissions were estimated by the ratios of total particle-phase I/SVOC components to POA (I/SVOCs-to-POA). The I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios for each source were determined according to their fractions of total I/SVOC species in VOC and POA emissions. Then we grouped different I/SVOC species into lumped I/SVOC bins based on their C* to determine the volatility distributions of each source. The I/SVOCs-G emissions were distributed into four lumped aliphatic IVOC bins across the volatility basis set from C*=10³ to 106 μg·m⁻³, two aromatic IVOC bins with the C*=10⁵ and 10⁶ μg·m⁻³, and four lumped SVOC bins with C* from 10⁻¹ and 10² μg·m⁻³. The I/SVOCs-P emissions were distributed into five bins spanning C*

from 10⁻¹ and 10³ µg·m⁻³. Source profiles of I/SVOC species for different sources were referenced from the results in previous studies. Table S1 and S2 show the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios for each specific source and their references. For industrial process, industrial solvent-use, and residential solvent-use sources, only I/SVOCs-G emissions were considered. Their I/SVOCs-G-to-VOCs ratios and emission profiles were derived from the latest version of SPECIATE 5.1 database (US EPA, 2021). For gasoline and diesel vehicles, the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios and emission profiles were referenced from a new mobilesource parameterization recommended by Lu et al. (2020). Those of diesel machinery, marine vessel, and residential coal combustion were determined by recent measurement results in China (Qi et al., 2019; Huang et al., 2018; Cai et al., 2019). The I/SVOCs-G-to-VOCs ratios and profiles of cooking and biomass burning emissions were derived from SPECIATE 5.1 database, while their particlephase ratios and profiles were referenced from two previous studies (May et al., 2013; Louvaris et al., 2017). Table S1 and S2 show the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios and their emission profiles of each specific source. The base emissions of AVOCs and POA (See Table S3) were taken from a high-resolution emission inventory for the year of 2017 developed in our previous study (An et al., 2021).

(3) Model input: Before being input into the model, the estimated I/SVOC-G and I/SVOC-P emissions were summed and then redistributed according to their phase equilibrium under their actual atmospheric state. The formula of phase equilibrium is shown in Equation (1).

$$F_{\rm p} = \frac{c_{\rm OA}}{c_{\rm OA} + c^*} \tag{1}$$

Where, F_p is the fraction of particle-phase I/SVOC emissions for each volatility bin under actual atmospheric state. C_{OA} represents the OA concentration under actual atmospheric state. We assumed it to be 10 μ g·m⁻³ in this study. C^* is the effective saturation concentration of each volatility bin. After redistribution, the I/SVOC emissions for each source category were allocated into 4 km \times 4 km grids and hourly temporal profiles using the same method as the criteria pollutants.

New references:

Louvaris, E. E., Florou, K., Karnezi, E., Papanastasiou, D. K., Gkatzelis, G. I., and Pandis, S. N.: Volatility of source apportioned wintertime organic aerosol in the city of Athens, Atmos. Environ., 158, 138–147, 2017.

L153-165: it is mentioned that the gas-phase IS-VOCs are obtained from ratios of IS-VOCs to VOCs and particle phase IS-VOCs are obtained from ratios of IS-VOCs to POA). Then IS-VOCs are distributed into volatility bins. The ratios are detailed in the supplementary material S1 (which should be referred to in the paper, but is not in the current version). How are the ratio defined? Only a few references are given in Supp S1. Please add a reference for each activity sector. How are the distributions into volatility bins defined?

By setting up differently the IS-VOCs in the gas phase and the IS-VOCs in the particle phase, how can we ensure that the gas and particle phases are consistent?

Re: Thanks for the comments. We agree with the reviewer that the methodology of I/SVOC emission inventory compilation needs more elaboration. Therefore, in the revised manuscript, we

provide a more detailed description of the estimation for I/SVOCs-G and I/SVOCs-P emissions and their volatility distributions. After the I/SVOC emission estimation, we then summarize them and redistributed their gas- and particle-phase emissions in each bin according to their phase equilibrium under actual atmospheric state to ensure the consistency of gas- and particle-phase emission before input into the model. We have made more detailed additions and revisions in the revised manuscript.

Changes in manuscript:

2.1 I/SVOC emission inventory

I/SVOCs commonly exist in both gas- and particle-phase in the atmosphere. Previous studies usually used POA scaling factors to estimate the I/SVOC emissions, which may lead to large uncertainties in the estimation of gas-phase organic compound-dominated sources, like oil refinery, chemical production, and industrial solvent-use. Herein, we compiled both gas-phase I/SVOCs (I/SVOCs-G) and particle-phase I/SVOCs (I/SVOCs-P) emission inventories and incorporate them into the model. Detailed process of the inventories is as follows.

- (1) Source classification: To refine the I/SVOC emissions from different sources, we divided the sources into five major categories and then further grouped them into 21 sub-categories. The major categories include industrial process sources, industrial solvent-use sources, mobile sources, residential sources, and agricultural sources. As shown in Table S1, the industrial process sources include the sectors such as oil refinery, chemical production, and pulp and paper production; Industrial solvent-use sources include textile, leather tanning, timber processing, and various industrial volatile chemical products use; Mobile sources include gasoline and diesel vehicle emissions, fuel evaporation, diesel machinery, marine vessel, and aircraft; Residential sources include coal combustion, residential solvent-use, and cooking emissions; Agricultural source is specifically referred to biomass burning in household stoves, and open burning was not included in this study.
- (2) Emission estimation: I/SVOCs-G emissions for each specific source were estimated by the ratios of total I/SVOC components to anthropogenic VOC (AVOC) components (I/SVOCs-to-VOCs). Similarly, I/SVOCs-P emissions were estimated by the ratios of total particle-phase I/SVOC components to POA (I/SVOCs-to-POA). The I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios for each source were determined according to their fractions of total I/SVOC species in VOC and POA emissions. Then we grouped different I/SVOC species into lumped I/SVOC bins based on their C* to determine the volatility distributions of each source. The I/SVOCs-G emissions were distributed into four lumped aliphatic IVOC bins across the volatility basis set from C*=10³ to 10⁶ $\mu g \cdot m^{-3}$, two aromatic IVOC bins with the C*=10⁵ and 10⁶ $\mu g \cdot m^{-3}$, and four lumped SVOC bins with C* from 10⁻¹ and 10² μg·m⁻³. The I/SVOCs-P emissions were distributed into five bins spanning C* from 10⁻¹ and 10³ μg·m⁻³. Source profiles of I/SVOC species for different sources were referenced from the results in previous studies. Table S1 and S2 show the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios for each specific source and their references. For industrial process, industrial solvent-use, and residential solvent-use sources, only I/SVOCs-G emissions were considered. Their I/SVOCs-G-to-VOCs ratios and emission profiles were derived from the latest version of SPECIATE 5.1 database (US EPA, 2021). For gasoline and diesel vehicles, the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios and emission profiles were referenced from a new mobilesource parameterization recommended by Lu et al. (2020). Those of diesel machinery, marine vessel, and residential coal combustion were determined by recent measurement results in China (Qi et al.,

2019; Huang et al., 2018; Cai et al., 2019). The I/SVOCs-G-to-VOCs ratios and profiles of cooking and biomass burning emissions were derived from SPECIATE 5.1 database, while their particle-phase ratios and profiles were referenced from two previous studies (May et al., 2013; Louvaris et al., 2017). Table S1 and S2 show the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios and their emission profiles of each specific source. The base emissions of AVOCs and POA (See Table S3) were taken from a high-resolution emission inventory for the year of 2017 developed in our previous study (An et al., 2021).

(3) Model input: Before being input into the model, the estimated I/SVOC-G and I/SVOC-P emissions were summed and then redistributed according to their phase equilibrium under their actual atmospheric state. The formula of phase equilibrium is shown in Equation (1).

$$F_{\rm p} = \frac{c_{\rm OA}}{c_{\rm OA} + c^*} \tag{1}$$

Where, F_p is the fraction of particle-phase I/SVOC emissions for each volatility bin under actual atmospheric state. C_{OA} represents the OA concentration under actual atmospheric state. We assumed it to be 10 μ g·m⁻³ in this study. C^* is the effective saturation concentration of each volatility bin. After redistribution, the I/SVOC emissions for each source category were allocated into 4 km \times 4 km grids and hourly temporal profiles using the same method as the criteria pollutants.

Changes in Supplementary information:

Table S1. Gas-phase I/SVOCs-to-VOCs ratios for specific sources and emission profiles used in CMAQ simulations. The characters in brackets are the source codes in the SPECIATE 5.1 database.

Source		I/SVOCs-G to VOCs	Volatility (C* at 298 K, μg·m ⁻³)									<u> </u>	
			IVOCP6	IVOCP5	IVOCP4	IVOCP3	SVOCP2	SVOCP1	SVOCP0	SVOCN1	IVOCP6ARO	IVOCP5ARO	References
			106	105	10 ⁴	10 ³	10 ²	10	1	10-1	10^{6}	10 ⁵	•
Industrial process	Oil refinery	0.039	0.759	0.123	0.004	0.110	0.003	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Chemical production	0.282	0.430	0.230	0.025	0.116	0.199	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Pulp and paper	0.140	0.571	0.393	0.028	0.006	0.001	0.001	0.000	0.000	0.000	0.000	SPECIATE 5.1
Industrial solvent-use	Textile	2.473	0.041	0.448	0.182	0.268	0.040	0.002	0.019	0.000	0.000	0.000	SPECIATE 5.1
	Leather tanning	0.231	1.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Timber processing	0.119	0.584	0.416	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Furniture coating	0.021	0.888	0.112	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Solvent-based coating	0.177	0.948	0.044	0.008	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Water-based coating	0.504	0.096	0.893	0.011	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Dry cleaning	0.004	0.885	0.115	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Paint remover	0.072	0.987	0.010	0.004	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
Mobile sources	Gasoline vehicle	0.265	0.206	0.056	0.113	0.098	0.000	0.000	0.000	0.000	0.406	0.121	Lu et al., 2020
	Diesel vehicle	1.358	0.331	0.318	0.244	0.095	0.000	0.000	0.000	0.000	0.004	0.007	Lu et al., 2020
	Fuel evaporation	0.002	0.841	0.159	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
	Diesel machinery	0.400	0.282	0.279	0.264	0.102	0.057	0.012	0.003	0.000	0.000	0.000	Qi et al., 2019
	Marine vessel	0.300	0.230	0.375	0.193	0.097	0.029	0.000	0.000	0.000	0.077	0.000	Huang et al., 2018
	Aircraft	0.482	0.761	0.148	0.063	0.028	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1
Residential sources	Coal combustion	0.180	0.439	0.439	0.088	0.035	0.000	0.000	0.000	0.000	0.000	0.000	Cai et al., 2019
	Residential solvent-use	0.240	0.938	0.047	0.003	0.007	0.000	0.003	0.000	0.001	0.000	0.000	SPECIATE 5.1
	Cooking	0.036	0.554	0.374	0.052	0.015	0.003	0.001	0.000	0.000	0.000	0.000	SPECIATE 5.1
Agriculture sources	Biomass burning	0.064	0.337	0.330	0.215	0.118	0.000	0.000	0.000	0.000	0.000	0.000	SPECIATE 5.1

Table S2. Particle-phase I/SVOCs-to-POA ratios for specific sources and emission profiles used in CMAQ simulations.

Source		I/SVOCs-P		Volatilit					
		to POA	IVOCP3	SVOCP2	SVOCP1	SVOCP0	SVOCN1	References	
		to POA	10 ³	10^{2}	10	1	10-1	_	
Mobile sources	Gasoline vehicle	0.901	0.000	0.323	0.406	0.073	0.197	Lu et al., 2020	
	Diesel vehicle	0.867	0.000	0.419	0.420	0.099	0.063	Lu et al., 2020	
	Diesel machinery	0.420	0.455	0.204	0.123	0.131	0.087	Qi et al., 2019	
	Marine vessel	0.469	0.305	0.140	0.185	0.166	0.204	Huang et al., 2018	
Residential sources	idential sources Cooking		0.670	0.157	0.003	0.000	0.000	Louvaris et al., 2017	
Agriculture sources Biomass burning		0.150	0.500	0.250	0.125	0.125	0.000	May et al., 2013	

Results and discussion

L293-296. The estimation of IS-VOC in the gas phase computed in this study is compared to what was found in other studies. However, there is not much details and it is hard to understand what differs between the studies. A section should be added to detail what was done in previous emission inventory of IS-VOC in China and aboard. This should be probably added in the introduction. I guess that the factors used to estimate IS-VOC are different in this study than in other studies over China, because IS-VOC are estimated from AVOCs. This should be clearly stated in the introduction. Also, note that IS-VOCs have been estimated from AVOCs in other previous studies over US and Europe.

Re: Thanks for the comments. We have supplemented the details of I/SVOC emission estimation in previous studies in the introduction section of the revised manuscript.

Changes in manuscript:

1. Introduction:

I/SVOC emission inventories have been developed and applied into air quality models over the past decade. Most of them were estimated by applying different scaling factors based on their relationship with POA, volatile organic compounds (VOCs), or some proxies like naphthalene (Pye and Seinfeld, 2010; Shrivastava et al., 2011; Jathar et al., 2017; Wu et al., 2019, 2021; Li et al., 2020, 2022; Ling et al., 2022). Yet in practice, a same scaling factor was applied to most of the sources in previous studies due to the lack of measurements on I/SVOC emission factors. For example, except biomass burning (0.75–1.5), Wu et al. (2019) utilized scaling factors of 8–30 for all of the other emission source categories, which was estimated based on the measurements of on-road mobile source. Li et al. (2020) assumed scaling factors of 1.5 for on-road mobile source, and 0.34–1.5 for the other sources, such as industrial and residential sources, which were much lower than the estimations in Wu et al. (2020). Huang et al. (2021) have tried emission factor method to quantify the I/SVOC emissions, yet the results were 60% lower than the scaling factor method, far from catching the measured amount of SOA. Obviously, roughly estimating I/SVOC emissions using one or two emission profiles as surrogates for all emission sources will create large uncertainties.

New references:

- Jathar, S. H., Woody, M., Pye, H. O. T., Baker, K. R., and Robinson, A. L.: Chemical transport model simulations of organic aerosol in southern California: model evaluation and gasoline and diesel source contributions, Atmos. Chem. Phys., 17, 4305–4318, 2017.
- Li, J., Han, Z., Wu, J., Tao, J., Li, J., Sun, Y., Liang, L., Liang, M., and Wang, Q.: Secondary organic aerosol formation and source contributions over east China in summertime, Environ. Pollut., 306, 119383, 2022.
- Ling, Z., Wu, L., Wang, Y., Shao, M., Wang, X., and Huang, W.: Roles of semivolatile and intermediate-volatility organic compounds in secondary organic aerosol formation and its implication: A review, J. Environ. Sci., 114, 259–285, 2022.
- Pye, H. O. T., Seinfeld, J. H.: A global perspective on aerosol from low-volatility organic compounds, Atmos. Chem. Phys., 10, 4377–4401, 2010.
- Shrivastava, M., Fast, J., Easter, R., Gustafson, W. I., Zaveri, R. A., Jimenez, J. L., Saide, P., and Hodzic, A.: Modeling organic aerosols in a megacity: comparison of simple and complex representations of the volatility basis set approach, Atmos. Chem. Phys., 11, 6639–6662, 2011.

L352. Please summarize the method of Lu et al. (2020) in a few sentences.

Re: Thanks for the comments. We have supplemented the summarization of the methods of Lu et al. (2020) in the revised manuscript.

Changes in manuscript:

3.1.2 Volatility distributions of I/SVOCs

Note that IVOCs in vehicle exhaust are dominated by aromatics, which have faster OH reaction rates and higher SOA yields compared to aliphatics in the same volatility bin (Zhao et al., 2016b; Drozd et al., 2019). Lu et al. (2020) therefore defined two additional lumped IVOC species with logC* bins at 5 and 6 to account for the aromatic IVOCs in vehicle exhaust according to the measurements in previous studies (Zhao et al., 2015; Zhao et al., 2016b). Here in this study, we also split the aromatic IVOC emissions from mobile sources and found that aromatic IVOCs accounted for 23% of the total I/SVOC emissions from the mobile source.

L449. Is there an increase of AVSOA in IMPROVE compared to BASE? It is surprising if there isn't, as organic concentrations are higher in the IMPROVE simulation, leading to higher absorbing mass.

Re: Thanks. The AVSOA concentration in IMPROVE simulation case increased compared to the BASE. The increasing rate was about 30%. We have supplemented some explanation in the revised manuscript.

Changes in manuscript:

3.2.1 Simulation results of OA concentrations

The seasonal average concentration of AVSOA in the BASE case was only 0.25 μg·m⁻³. The average AVSOA concentration in the IMPROVE case increased by 30.1% compared with the BASE

case due to higher OA loading. Nonetheless, AVSOA still exhibited very limited contribution to the regional OA concentration, whereas average concentration of BVOC derived SOA (BVSOA, 1.7 µg·m⁻³) was much higher.

L451. What was expected?

Re: Sorry for the mistake. We have deleted this statement in the revised manuscript.

Changes in manuscript:

The seasonal average concentration of AVSOA in the BASE case was only $0.25~\mu g\cdot m^{-3}$. The average AVSOA concentration in the IMPROVE case increased by 30.1% compared with the BASE case due to higher OA loading. Nonetheless, AVSOA still exhibited very limited contribution to the regional OA concentration, whereas average concentration of BVOC derived SOA (BVSOA, $1.7~\mu g\cdot m^{-3}$) was much higher.

Conclusions

L655. « SOA increased by 1.2 times in IMPROVE simulation ». However, L431 stated that OA are 38% higher in IMPROVE simulation. Why are these numbers different?

Re: OA is composed of POA and SOA. On L655, we only compared SOA in IMPROVE and BASE cases, while on L431, total OA concentrations were compared.

General Comments:

An and coauthors have assembled an emission inventory of organic compounds spanning likely particle-phase, semivolatile, intermediate volatility, and highly volatile pollutants for a comprehensive list of key sources in China. After a brief summary of the changes incurred by this new inventory relative to an existing Base, the authors apply both inventories to model ambient air concentrations of primary and secondary OA using a state-of-the-art chemical transport model, CMAQ. The methods used in this study appear generally sound (with an exception discussed below) and the analysis is complete. The information provided by the study in terms of source contributions for OA should be valuable, I expect, to policy-makers in China, and to the larger Earth system modeling community. I was particularly impressed at the model performance improvement that the authors were able to document. I have some concerns about details of the implementation, some questions about surprising results, and several minor suggestions, which I have included below.

Major Concerns:

1. I am missing something fundamental from the presentation in the paper. The authors have documented both I/SVOC-G emissions and I/SVOC-P emissions, but are they being added together in the model? I think it is very likely that there is overlap in this region and that if both are used, then they will be double-counting some emissions, especially the SVOCs, but also the IVOCs. Can the authors be much more specific about how they fit these pieces together? In lines 241-242, the authors mention subtracting the semivolatile portion from the total POA, but there needs to be more detailed description in one place of what is going on here.

Re: Thanks for the reviewer's comments. The I/SVOCs-G and I/SVOCs-P emissions in this study were estimated independently. The reason for this consideration is that the I/SVOC emissions of most sources include both gas- and particle-phase. Therefore, we determined the I/SVOCs-G based on the fractions of I/SVOC components in VOCs in the gas-phase profiles, while the I/SVOCs-P were determined based on the fractions of I/SVOC components in POA in the particle-phase profiles. The fractions in the gas- and particle-phase profiles were usually obtained from the gas- (eg Tenax tube) and particle-phase (eg filters) sampling system, respectively. We believe there was no doublecounting for the I/SVOC emissions. After estimating I/SVOCs-G and I/SVOCs-P emissions, we added them together and redistributed into the model according to their phase equilibrium under the actual atmosphere state. In the revised manuscript, we have supplemented the description of I/SVOC emission estimation. The SOA formed from I/SVOCs-G was estimated using the parameterization within the VBS framework in Lu et al. (2020). POA was treated as semivolatile to account for its gas-particle partitioning and ageing process and segregated to several particle species, which varied in their volatility that quantified with the metric $C^* = 10^{-1}$ to $10^3 \,\mu\text{g}\cdot\text{m}^{-3}$ (Donahue et al., 2006). I/SVOCs-P emissions from different sources were then speciated and input as semivolatile accordingly. The remaining POA emissions excluding I/SVOCs-P were treated as nonvolatile POC (primary organic carbon) and PNCOM (primary non-carbon organic matter). We have supplemented

the description of POA modeling scheme in the revised manuscript.

Changes in manuscript:

2.1 I/SVOC emission inventory

I/SVOCs commonly exist in both gas- and particle-phase in the atmosphere. Previous studies usually used POA scaling factors to estimate the I/SVOC emissions, which may lead to large uncertainties in the estimation of gas-phase organic compound-dominated sources, like oil refinery, chemical production, and industrial solvent-use. Herein, we compiled both gas-phase I/SVOCs (I/SVOCs-G) and particle-phase I/SVOCs (I/SVOCs-P) emission inventories and incorporate them into the model. Detailed process of the inventories is as follows.

- (1) Source classification: To refine the I/SVOC emissions from different sources, we divided the sources into five major categories and then further grouped them into 21 sub-categories. The major categories include industrial process sources, industrial solvent-use sources, mobile sources, residential sources, and agricultural sources. As shown in Table S1, the industrial process sources include the sectors such as oil refinery, chemical production, and pulp and paper production; Industrial solvent-use sources include textile, leather tanning, timber processing, and various industrial volatile chemical products use; Mobile sources include gasoline and diesel vehicle emissions, fuel evaporation, diesel machinery, marine vessel, and aircraft; Residential sources include coal combustion, residential solvent-use, and cooking emissions; Agricultural source is specifically referred to biomass burning in household stoves, and open burning was not included in this study.
- (2) Emission estimation: I/SVOCs-G emissions for each specific source were estimated by the ratios of total I/SVOC components to anthropogenic VOC (AVOC) components (I/SVOCs-to-VOCs). Similarly, I/SVOCs-P emissions were estimated by the ratios of total particle-phase I/SVOC components to POA (I/SVOCs-to-POA). The I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios for each source were determined according to their fractions of total I/SVOC species in VOC and POA emissions. Then we grouped different I/SVOC species into lumped I/SVOC bins based on their C* to determine the volatility distributions of each source. The I/SVOCs-G emissions were distributed into four lumped aliphatic IVOC bins across the volatility basis set from C*=10³ to 10⁶ $\mu g \cdot m^{-3}$, two aromatic IVOC bins with the C*=10⁵ and 10⁶ $\mu g \cdot m^{-3}$, and four lumped SVOC bins with C* from 10⁻¹ and 10² µg·m⁻³. The I/SVOCs-P emissions were distributed into five bins spanning C* from 10⁻¹ and 10³ µg·m⁻³. Source profiles of I/SVOC species for different sources were referenced from the results in previous studies. Table S1 and S2 show the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios for each specific source and their references. For industrial process, industrial solvent-use, and residential solvent-use sources, only I/SVOCs-G emissions were considered. Their I/SVOCs-G-to-VOCs ratios and emission profiles were derived from the latest version of SPECIATE 5.1 database (US EPA, 2021). For gasoline and diesel vehicles, the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios and emission profiles were referenced from a new mobilesource parameterization recommended by Lu et al. (2020). Those of diesel machinery, marine vessel, and residential coal combustion were determined by recent measurement results in China (Qi et al., 2019; Huang et al., 2018; Cai et al., 2019). The I/SVOCs-G-to-VOCs ratios and profiles of cooking and biomass burning emissions were derived from SPECIATE 5.1 database, while their particlephase ratios and profiles were referenced from two previous studies (May et al., 2013; Louvaris et al., 2017). Table S1 and S2 show the I/SVOCs-G-to-VOCs and I/SVOCs-P-to-POA ratios and their

emission profiles of each specific source. The base emissions of AVOCs and POA (See Table S3) were taken from a high-resolution emission inventory for the year of 2017 developed in our previous study (An et al., 2021).

(3) Model input: Before being input into the model, the estimated I/SVOC-G and I/SVOC-P emissions were summed and then redistributed according to their phase equilibrium under their actual atmospheric state. The formula of phase equilibrium is shown in Equation (1).

$$F_{\rm p} = \frac{c_{\rm OA}}{c_{\rm OA} + c^*} \tag{1}$$

Where, F_p is the fraction of particle-phase I/SVOC emissions for each volatility bin under actual atmospheric state. C_{OA} represents the OA concentration under actual atmospheric state. We assumed it to be 10 μ g·m⁻³ in this study. C^* is the effective saturation concentration of each volatility bin. After redistribution, the I/SVOC emissions for each source category were allocated into 4 km \times 4 km grids and hourly temporal profiles using the same method as the criteria pollutants.

New references:

Louvaris, E. E., Florou, K., Karnezi, E., Papanastasiou, D. K., Gkatzelis, G. I., and Pandis, S. N.: Volatility of source apportioned wintertime organic aerosol in the city of Athens, Atmos. Environ., 158, 138–147, 2017.

2.2 Model configuration

POA was treated as semivolatile to account for its gas-particle partitioning and ageing process and segregated to several particle species, which varied in their volatility that quantified with the metric $C^* = 10^{-1}$ to $10^3 \, \mu g \cdot m^{-3}$ (Donahue et al., 2006). I/SVOCs-P emissions from different sources were then speciated and input as semivolatile accordingly. The remaining POA emissions excluding I/SVOCs-P were treated as nonvolatile POC (primary organic carbon) and PNCOM (primary noncarbon organic matter).

2. It is surprising in Table 2 that there are no I/SVOC-P emissions from the industrial processes, aircraft, and coal combustion emissions. Do the authors consider this realistic, or is this an area where more data are needed?

Re: Thanks. We agree with the reviewer's comments. There should be some I/SVOCs-P emissions from the industrial processes, aircraft, and coal combustion like power plants and boilers. However, we didn't consider the I/SVOCs-P emissions from the above source for the following two reasons. One is that POA emissions from these sources were limited, which will not affect the results in this study. According to our base emissions inventory, their POA emissions accounted for 1.6%, 0.1% and 4.0% of total emissions, respectively. Second, the profiles of I/SVOCs-P components are difficult to obtain, which also indicates that more measurements of I/SVOCs emission sources are needed in the future. We have addressed the absence of these emission sources in the revised manuscript.

Changes in manuscript:

3.1.1 Source-specific I/SVOC emissions

I/SVOCs-P emissions were 82.96 Gg. The largest contributor of I/SVOCs-P emissions came from biomass burning and diesel vehicle, accounting for 53.24% and 11.88% of the total, followed by gasoline vehicle (5.23%), marine vessel (2.66%), diesel machinery (2.54%), and biomass burning (1.75%). Note that the I/SVOCs-P emissions from coal combustion (e.g. power plants, boilers, etc.), other industrial processes, and aircraft were not included in this study. On the one hand, the POA emissions (see Table S3) from these sources were limited, accounting for less than 5%, which could be expected that their I/SVOCs-P emissions were also relatively low. On the other hand, the profiles of I/SVOCs-P components of these sources were still difficult to obtain. More measurements of the I/SVOC emissions from these sources is very necessary in the future.

3. Can the authors explain some of the interesting trends shown in Fig. 2. Specifically, I find it curious that there is a substantial fraction of I/SVOC-G in the C*=0.1 ug m-3 bin for agricultural sources. Also, the residential source has an odd volatility trend, with very little emissions in the lower IVOC range. Do the authors think this is realistic? I recommend replacing gas-phase and particle-phase in Fig. 2 with 'I/SVOC-G' and 'I/SVOC-P' or something similarly precise. These are not necessarily gas-phase or particle-phase emissions, but are instead emissions based on VOC and filter OC measurements (I assume; it's not really discussed in the description of the emission inventory).

Re: Thanks. We agree with the reviewer's comments. The I/SVOCs-G and I/SVOCs-P profiles in Fig. 2 still cannot capture the actual emission characters in real-world. This is mainly due to the limited available I/SVOC emission measurements. Regarding the problem that the agricultural source has a substantial fraction of I/SVOCs-G in the C*=0.1 μg·m⁻³, we find there are some mistakes in the profile mapping after inspection. Therefore, we have made the following changes in the revised manuscript according to the comments. First, we have checked and corrected the profile data in Fig. 2 again. Then we summed the "Gas-phase" and "Particle-phase" emissions in each bin in Fig. 2(b-g) since the fractions of gas- and particle-phase emissions were determined by real atmospheric conditions, which was meaningless to be exhibited in the figures.

Changes in manuscript:

3.1.2 Volatility distributions of I/SVOCs

Figure 2 shows the volatility distribution of I/SVOC emissions from different sources as well as their gas-particle distributions under actual atmospheric state. The I/SVOC emissions generally showed an increasing trend with the increase of volatility. As shown in Figure 2(a), IVOC emissions (logC* bins at 3–6) accounted for 86% of the total I/SVOCs emissions, overwhelmingly dominated by industrial process and mobile sources. SVOCs (logC* bins at 0–2) and low-volatile organic compounds (LVOCs, logC* bins at -1) contributed to 11% and 3% of the total I/SVOCs emissions. In terms of the contributing sectors, mobile sources, industrial process, and solvent-use dominated the total I/SVOC emissions. While the IVOCs were equally contributed by above-listed three sources, residential and mobile sources dominated the SVOCs and LVOCs emissions.

We further investigated the contributions of different volatility bins to each source category. The mobile source was dominated by IVOC emission (88%). Note that IVOCs in vehicle exhaust are dominated by aromatics, which have faster OH reaction rates and higher SOA yields compared

to aliphatics in the same volatility bin (Zhao et al., 2016b; Drozd et al., 2019). Lu et al. (2020) therefore defined two additional lumped IVOC species with logC* bins at 5 and 6 to account for the aromatic IVOCs in vehicle exhaust according to the measurements in previous studies (Zhao et al., 2015; Zhao et al., 2016b). Here in this study, we also split the aromatic IVOC emissions from mobile sources and found that aromatic IVOCs accounted for 23% of the total I/SVOC emissions from the mobile source. The industrial process and solvent-use sources were also dominated by IVOC emissions, accounting for 81% and 97%, respectively. The volatility distribution of residential sources was relatively uniform, with IVOCs, SVOCs and LVOCs accounting for 40%, 30%, and 30%. Agricultural (i.e., biomass burning) sources were more concentrated in IVOCs, accounting for 76%, while SVOCs accounted for 24%. It should be noted that other than mobile sources, the emission profiles of the other sources were mainly derived from SPECIATE 5.1 database (US EPA, 2021) in this study, which may be inconsistent with real-world emissions in China. To further reduce the uncertainty in the I/SVOC emission inventory, measurements of I/SVOC emissions from different local sources are therefore important and urgently needed in the future.

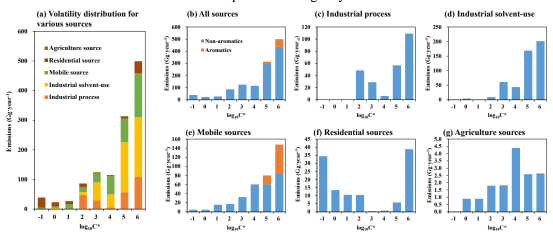


Figure 2. Volatility distributions of I/SVOCs emitted from different sources in the YRD region.

4. I was surprised to not see a multi-panel figure showing the spatial emissions of LVOC, SVOC, IVOC and VOC. Can this be added? This would aid comparison to contemporary studies like Chang et al. (2022; https://www.sciencedirect.com/science/article/pii/ S2590332222001488)

Re: Thanks for the reminding. We have supplemented the figures and associated descriptions of spatial emissions of VOCs, IVOCs, SVOCs, and LVOCs in the revised manuscript.

Changes in manuscript:

3.1.3 Spatial distributions of I/SVOC emissions in YRD region

Figure 3 compares the spatial distributions of AVOC, IVOC, SVOC, and LVOC emissions in the YRD region. The IVOC, SVOC, and LVOC emissions were largely concentrated in city clusters in eastern YRD, and hotspots can also be observed in the northern agglomerations. The distributions of I/S/LVOC emissions were generally consistent with that of the AVOC emissions in the region. Compared to the spatial distributions of I/S/LVOC emissions in Chang et al. (2022), our emissions had similar spatial distributions but at a higher resolution. Emission hotspots in urban areas can be captured more clearly in this study, which will help improve the simulation in urban areas.

Figure 4 shows the spatial distributions of source-specific I/SVOC emissions in the YRD region. There were considerable differences in the spatial distributions of I/SVOC emissions from different sources. The I/SVOC emissions from industrial sources (including industrial process and industrial solvent-use) were mainly concentrated in the eastern urban agglomeration, which was related to the developed industrial activities in the region. The I/SVOC emissions from mobile and residential sources clustered into multiple hotspots in urban areas, while emissions from agricultural sources were mainly distributed in northern YRD, where frequent agricultural activities exist.

We also compare the spatial distributions of I/SVOC emissions with those of POA and BVOCs. We found that POA emissions were more concentrated in urban centers associated with mobile and residential sources (See Figure S1). BVOC emissions in the YRD region were mainly distributed in the southern area, where AVOC and IVOC emissions were relatively low. The difference in the spatial distributions of I/SVOC, AVOC, BVOC, and POA emissions implies that the sources of organic components in different areas of the region are quite different, which will be discussed in the following sections.

New references:

Chang, X., Zhao, B., Zheng, H., Wang, S., Cai, S., Guo, F., Gui, P., Huang, G., Wu, D., Han, L., Xing, J., Man, H., Hu, R., Liang, C., Xu, Q., Qiu, X., Ding, D., Liu, K., Han, R., Robinson, A. L., and Donahue, N. M.: Full-volatility emission framework corrects missing and underestimated secondary organic aerosol sources, One Earth, 5, 403–412, 2022.

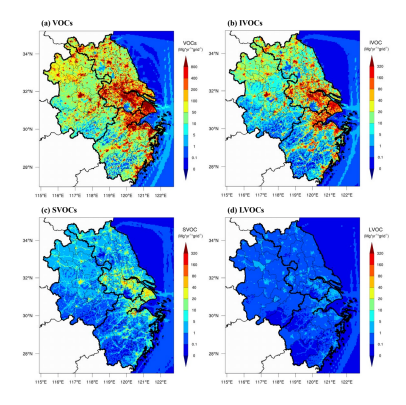


Figure 3. Spatial distributions of VOC, IVOC, SVOC, and LVOC emissions in the YRD region for the year 2017.

5. In Fig. 6, it is surprising that the POA trend is relatively unchanged between the BASE and IMPROVE cases, when perhaps 60% of the POA is being evaporated (line 329-330). Can the authors shed some light here? Is most of the I/SVOC-P going into the LVOC bins? Judging by Table S2, it doesn't seem like it.

Re: Thanks for the comments. We re-examined the simulations of POA concentrations for the IMPROVE simulation case after re-distributing the gas- and particle-phase I/SVOC emissions according to their phase equilibrium under the actual atmosphere state. The results indicated POA concentrations in the IMPROVE case decreased by 12%–20% compared with the BASE case. We have made the corrections in the revised manuscript.

Changes in manuscript:

3.2.1 Simulation results of OA concentrations

The seasonal average concentration of POA was 5.5 μg·m⁻³ in the BASE case, with the lowest in summer (3.8 μg m⁻³) and the highest in winter (6.9 μg m⁻³). High POA concentrations in winter was mainly induced by the stagnant meteorological conditions such as low wind speed and boundary layer height, and vice versa in summer. For the spatial distributions as presented in Figure 6, POA concentrations in northern YRD were high and mainly concentrated in urban areas, which was consistent with the distributions of POA emissions (Figure S1). The POA concentrations in the IMPROVE simulation decreased by 12%–20% compared with the BASE case. In the IMPROVE simulation, the POA was treated as semi-volatile, where gas–particle partitioning and multigeneration oxidation were considered (Murphy et al., 2017). Entering into the atmosphere, more semi-volatile compounds evaporated into gas-phase and then generated SOA through multigeneration oxidation, which reduced the POA concentrations relatively.

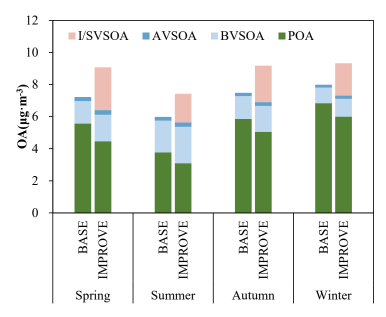


Figure 5. Comparisons of the regional average concentrations of POA and SOA formed from AVOCs, BVOCs, and I/SVOCs in different seasons from the BASE and IMPROVE simulations.

6. Although the authors have done a commendable job in the writing, there are more than a

few grammatically incorrect and awkward phrases throughout. I recommend the authors have a technical writer give feedback, or someone from the journal staff.

Re: Thanks. Grammar and writing have been checked by some technical writers in the revised manuscript.

Minor Suggestions/Typos:

1. Line 53: Jathara --> Jathar.

Re: Thanks. It has been revised.

2. Line 53-54: Recommend rephrasing to: "It is challenging to constrain the abundance of OA precursors and to identify key sources."

Re: Thanks. We have rephrased the sentence as the reviewer recommended.

3. Line 57: Please add references for studies using the three source apportionment techniques. Also add a reference for the AMS to the next sentence.

Re: Please see the following changes.

Changes in manuscript:

1. Introduction

Great efforts have been made in the identification of OA sources through source apportionment of the measured OA components, such as positive matrix factorization (PMF) (Zhang et al., 2011), chemical mass balance (CMB) model (Zheng et al., 2002) or multilinear engine (ME-2) (Canonaco et al., 2013). The Aerodyne high-resolution time-of-flight aerosol mass spectrometer (AMS), has been proven to be a powerful tool in quantification and chemical characterization of different OA components in real-time (Canagaratna et al., 2007).

New references:

Zhang, Q., Jimenez, J. L., Canagaratna, M. R., Ulbrich, I. M., Ng, N. L., Worsnop, D. R., and Sun, Y.: Understanding atmospheric organic aerosols via factor analysis of aerosol mass spectrometry: a review, Anal. Bioanal. Chem., 401, 3045–3067, 2011.

Zheng, M., Cass, G. R., Schauer, J. J., and Edgerton, E. S.: Source Apportionment of PM_{2.5} in the Southeastern United States Using Solvent-Extractable Organic Compounds as Tracers, Environ. Sci. Technol., 36, 2361–2371, 2002.

4. Line 74: model --> modeling

Re: Thanks. It has been revised.

5. Line 75: which advantages in --> which has advantages for

Re: Thanks. It has been revised.

6. Line 77-78: Please be careful here. Some models do include multi-generational oxidation of vapors. Are you referring to a specific model here? A specific CMAQ version? Note that CMAQ has included aging via oligomerization of anthropogenic and biogenic vapors for

some time.

Re: Thanks for the reminding. We have deleted this statement in the revised manuscript.

7. Line 87: Recommend rephrasing 'due to the missing of' to 'due to the underestimation of'

Re: Thanks. It has been revised.

8. Line 102-104: Many of these profiles are not in SPECIATE5.1, and SPECIATE5.1 generally does not support lumped species by volatility. Recommend removing this sentence, or change 'most' to 'some'.

Re: Thanks. We have removed this statement in the revised manuscript.

9. Line 106: Is 30% really dominating? Maybe replace 'dominates' with 'contributes significantly to'?

Re: Thanks. It has been revised.

10. Line 116-127: Recommend moving to Methods section

Re: Thanks for the comments. According to another reviewer's comments, we have rewritten this section to highlight the novelty of this study in I/SVOC emission estimation through a survey of established methodologies in previous studies. The specific modifications are as follows:

Changes in manuscript:

1. Introduction:

I/SVOC emission inventories have been developed and applied into air quality models over the past decade. Most of them were estimated by applying different scaling factors based on their relationship with POA, volatile organic compounds (VOCs), or some proxies like naphthalene (Pye and Seinfeld, 2010; Shrivastava et al., 2011; Jathar et al., 2017; Wu et al., 2019, 2021; Li et al., 2020, 2022; Ling et al., 2022). Yet in practice, a same scaling factor was applied to most of the sources in previous studies due to the lack of measurements on I/SVOC emission factors. For example, except biomass burning (0.75–1.5), Wu et al. (2019) utilized scaling factors of 8–30 for all of the other emission source categories, which was estimated based on the measurements of on-road mobile source. Li et al. (2020) assumed scaling factors of 1.5 for on-road mobile source, and 0.34–1.5 for the other sources, such as industrial and residential sources, which were much lower than the estimations in Wu et al. (2020). Huang et al. (2021) have tried emission factor method to quantify the I/SVOC emissions, yet the results were 60% lower than the scaling factor method, far from catching the measured amount of SOA. Obviously, roughly estimating I/SVOC emissions using one or two emission profiles as surrogates for all emission sources will create large uncertainties.

New references:

- Jathar, S. H., Woody, M., Pye, H. O. T., Baker, K. R., and Robinson, A. L.: Chemical transport model simulations of organic aerosol in southern California: model evaluation and gasoline and diesel source contributions, Atmos. Chem. Phys., 17, 4305–4318, 2017.
- Li, J., Han, Z., Wu, J., Tao, J., Li, J., Sun, Y., Liang, L., Liang, M., and Wang, Q.: Secondary organic aerosol formation and source contributions over east China in summertime, Environ. Pollut., 306, 119383, 2022.
- Ling, Z., Wu, L., Wang, Y., Shao, M., Wang, X., and Huang, W.: Roles of semivolatile and intermediate-volatility organic compounds in secondary organic aerosol formation and its implication: A review, J. Environ. Sci., 114, 259–285, 2022.
- Pye, H. O. T., Seinfeld, J. H.: A global perspective on aerosol from low-volatility organic compounds, Atmos. Chem. Phys., 10, 4377–4401, 2010.
- Shrivastava, M., Fast, J., Easter, R., Gustafson, W. I., Zaveri, R. A., Jimenez, J. L., Saide, P., and Hodzic, A.: Modeling organic aerosols in a megacity: comparison of simple and complex representations of the volatility basis set approach, Atmos. Chem. Phys., 11, 6639–6662, 2011.

11. Line 199: What was used for biogenic emissions? MEGAN?

Re: Yes. Sorry forgot to explain in the text. We have supplemented the statement of biogenic emission estimation in the revised manuscript.

Changes in manuscript:

2.2 Model configuration

Biogenic volatile organic compounds (BVOCs) emissions were estimated based on MEGAN (the Model of Emissions of Gases and Aerosols from Nature) version 2.10 driving by inputs of the leaf area index (LAI) from MODIS product, plant functional types (PFT) base on remote sensing data, inline coupled emission factors and meteorology simulated by the WRF model. Detail configurations of MEGAN can be obtained from our previous study (Liu et al., 2018).

12. Line 206-207: The AERO7 CMAQ aerosol scheme does not include IVOCs with C* between 104 and 106 ug m-3. It also doesn't include aromatic IVOCs. Did the authors add these?

Re: Yes. We included a new-developed of VBS scheme recommended by Lu et al. (2020) in this study. The scheme included the OH oxidation of IVOCs with C* between 10^4 and 10^6 $\mu g \cdot m^{-3}$ and aromatic IVOCs. The description of this scheme is shown in the fifth paragraph in Section "2.2 Model configuration".

13. Line 228-230: Is the multigenerational oxidation scheme equivalent to that used in Lu et al and Murphy et al, or did the authors derive their own?

Re: The scheme was equivalent to that used in Lu et al. and Murphy et al. We have mentioned in the manuscript.

14. Table 1: Recommend reformatting to get rid of the repeated phrase "VOC and I/SVOC emissions in the region". Maybe change the column 'Emission settings' to something like 'Sources with added I/SVOC emissions'. Then the first row would be 'none', the second row 'all', and each latter row would be something like 'all except XX'. I don't think you need the Notes column.

Re: Thanks for the reviewer's recommendation. We have reformatted the table in the revised manuscript.

Changes in manuscript:

2.3 Model simulations

Table 1. Settings of simulation cases.

Name	Sources with added I/SVOC emissions					
BASE	none					
IMPROVE	all					
CASE1	all except industrial process					
CASE2	all except industrial solvent-use					
CASE3	all except mobile sources					
CASE4	all except residential sources					
CASE5	all except biomass burning					
CASE6	all except biogenic sources					
CASE7	without VOCs and I/SVOC emissions					
CASE8	all except gasoline vehicle					
CASE9	all except diesel vehicle					
CASE10	all except diesel machinery					
CASE11	all except marine vessel					
CASE12	all except cooking					

15. Line 296-300: I don't understand these sentences. Please rephrase.

Re: Thanks for the reminding. We also find these sentences doesn't seem to make much sense. We have removed these statements in the revised manuscript.

16. Line 358-359: Please be careful here when referring to species as 'gas-phase'. You have chosen to plot these data as particle-phase or gas-phase, which I don't think is the best way to go, since in reality, they are just scaled off the particle-phase and gas-phase emissions, respectively. They could be in either phase when emitted.

Re: Thanks for the reminding. We understand the reviewer's suggestion that the gas- and particle-phase of the I/SVOCs components are dynamically changing and should not be simply divided. Therefore, we have merged the gas- and particle-phase data of I/SVOCs in each volatility bin in Figure 2 in the revised manuscript.

Changes in manuscript:

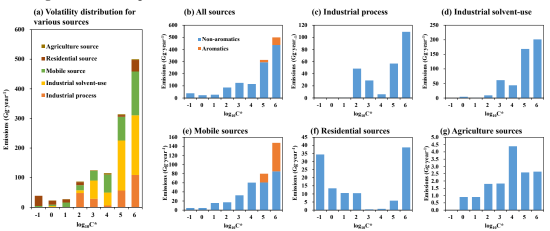


Figure 2. Volatility distributions of I/SVOCs emitted from different sources in the YRD region.

17. There are two tables labelled Table S6.

Re: Thanks. It has been revised.

18. 5: Color axes are difficult to read. Please enlarge them.

Re: Thanks. We have enlarged the color axes of the figures.

Changes in manuscript:

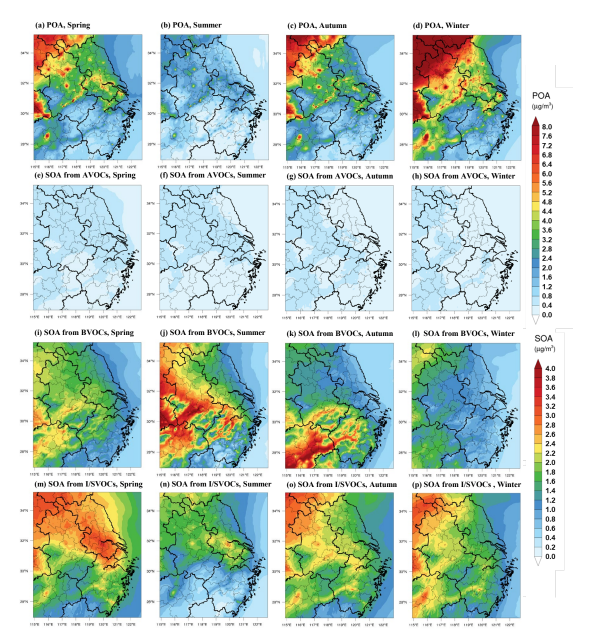


Figure 6. Spatial distributions of modeled POA and SOA formed from AVOCs, BVOCs, and I/SVOCs in different seasons in the IMPROVE simulation.

19. Section 3.3: I recommend adding a few sentences comparing to the YRD-specific source contributions reported by Chang et al. (2022) since the goals of these studies are highly related.

Re: Thanks for the comments. We have supplemented some statements on the comparison with the results reported by Chang et al. (2022) in the revised manuscript.

Changes in manuscript:

3.3.1 POA and SOA sources in the region

Our results were generally similar with those of Chang et al. (2022) for YRD region. We both

found the domestic combustion mainly engaged in cooking emissions had a major contribution to OA. Next was volatile chemical products (VCPs), especially the use of solvents, paints, and adhesives in industrial sector, also made a high contribution. Note that industrial process also took up a high fraction in our OA simulation, while it was lower in Chang et al. (2022)'s study. The difference in I/SVOC emission estimates was the main reason for this divergence. Mobile sources in both studies had similar contributions, which accounted for about 10% to total OA. Comparatively, our source classification was more specific, which will help identify more specific OA sources to design more refined regional control countermeasures.

20. Line 610: supe-regional --> super-regional

Re: Thanks. It has been revised.