

# Changes of air pollutant emissions in China during two clean air action periods derived from the newly developed Inversed Emission Inventory for Chinese Air Quality (CAQIEI)

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## Abstract

A new long-term emission inventory called the Inversed Emission Inventory for Chinese Air Quality (CAQIEI) was developed in this study by assimilating surface observations from the China National Environmental Monitoring Centre (CNEMC) using the ensemble Kalman filter and the Nested Air Quality Prediction Modeling System. This inventory contains the constrained monthly emissions of NO<sub>x</sub>, SO<sub>2</sub>, CO, primary PM<sub>2.5</sub>, primary PM<sub>10</sub>, and NMVOCs in China from 2013 to 2020, with a horizontal resolution of 15 km × 15 km. This paper documents detailed descriptions of the assimilation system and the evaluation results for the emission inventory. The results suggest that CAQIEI can effectively reduce the biases in the *a priori* emission inventory, with the normalized mean biases ranging from -9.1% to 9.5% in the *a posteriori* simulation, which are significantly reduced from the biases in the *a priori* simulations (-45.6% to 93.8%). The calculated RMSEs (0.3 mg/m<sup>3</sup> for CO and 9.4–21.1 µg/m<sup>3</sup> for other species, on the monthly scale) and correlation coefficients (0.76–0.94) were also improved from the *a priori* simulations, demonstrating the good performance of the data assimilation system. Based on CAQIEI, we estimated China's total emissions (including both natural and anthropogenic emissions) of the 6 species in 2015 to be as follows: 25.2 Tg of NO<sub>x</sub>, 17.8 Tg of SO<sub>2</sub>, 465.4 Tg of CO, 15.0 Tg of PM<sub>2.5</sub>, 40.1 Tg of PM<sub>10</sub>, and 46.0 Tg of NMVOCs. From 2015 to 2020, the total emissions reduced by 54.1% for SO<sub>2</sub>, 44.4% for PM<sub>2.5</sub>, 33.6% for PM<sub>10</sub>, 35.7% for CO, and 15.1% for NO<sub>x</sub>, but increased by 21.0% for NMVOCs. It is also estimated that the emission reductions were larger during 2018–2020 (from -26.6% to -4.5%) than during 2015–2017 (from -23.8% to 27.6%) for most species. Particularly, the total Chinese NO<sub>x</sub> and NMVOC emissions were shown to increase during 2015–2017, especially over the Fenwei Plain area (FW) where the emissions of particulate matter (PM) also increased. The situation changed during 2018–2020 when the upward trends were contained and reversed to a downward trends for both the total emissions of NO<sub>x</sub> and NMVOC, and the PM emissions over FW. This suggests that the emission control policies may be improved in the 2018–2020 action plan. We also compared the CAQIEI with other air pollutant emission inventories in China, which verified our inversion results in terms of total emissions of NO<sub>x</sub>, SO<sub>2</sub> and NMVOCs, and more importantly identified the potential uncertainties in current emission inventories. Firstly, the CAQIEI suggested higher CO emissions in China, with CO emissions estimated by CAQIEI (426.8 Tg) being more than

45 twice the amount in previous inventories (120.7–237.7 Tg). Significantly higher emissions were also suggested over the  
46 western and northeastern China for other air pollutants. Secondly, the CAQIEI suggested higher NMVOC emissions than  
47 previous emission inventories by about 30.4–81.4% over the North China Plain (NCP)-region but suggested lower NMVOC  
48 emissions by about 27.6–0.0% over the Southeast China (SE)-region. Thirdly, the CAQIEI suggested smaller emission  
49 reduction rates during 2015–2018 than previous emission inventories for most species except of CO. Particularly, China’s  
50 NMVOC emissions were shown to have increased by 26.6% from 2015 to 2018, especially over the NCP (by 38.0%), northeast  
51 China (by 38.3%), and central China (60.0%). These results provide us with new insight into the complex variations of the air  
52 pollutant emissions in China during ~~its~~ two recent clean air actions, which has the potential to improve our understanding of  
53 air pollutant emissions in China and their impacts on air quality. The whole datasets are available at  
54 <https://doi.org/10.57760/sciencedb.13151> (Kong et al., 2023).

## 55 **1 Introduction**

56 Air pollution is a serious environmental issue owing to its substantial impacts on human health, ecosystems, and climate  
57 change (Von Schneidmesser et al., 2015; Cohen et al., 2017; Bobbink et al., 1998). According to the World Health  
58 Organization, air pollution-induced strokes, lung cancer, and heart disease are causing millions of premature deaths worldwide  
59 every year (WHO, 2016). The fine particulate matter (PM<sub>2.5</sub>) in the atmosphere not only degrades visibility but also affects the  
60 radiative forcing of the climate, both directly and indirectly (Martin et al., 2004). After removal from the atmosphere through  
61 dry and wet deposition, air pollutants such as sulfur, nitrate, and ammonium contribute significantly to soil acidification,  
62 eutrophication, and even biodiversity reduction (Krupa, 2003; Hernández et al., 2016).

63 China has experienced severe PM<sub>2.5</sub> pollution in recent decades, due to its large emissions of air pollutants associated  
64 with rapid urbanization and high consumption of fossil fuels (Kan et al., 2012; Song et al., 2017). The annual concentrations  
65 of PM<sub>2.5</sub> in 2013 reached 106, 67 and 47  $\mu\text{g}/\text{m}^3$  over the Beijing–Tianjin–Heibei, Yangtze River Delta, and Peral River Delta  
66 region, respectively, which were all higher than China’s national standard (35  $\mu\text{g}/\text{m}^3$ ), and 5–10 times higher than that of the  
67 World Health Organization (10  $\mu\text{g}/\text{m}^3$ ). To tackle this problem, strict emission control policies (so-called “clean air action  
68 plans”) have been proposed by China’s government, including the “Action Plan on the Prevention and Control of Air Pollution”  
69 from 2013 to 2017 (hereinafter called the “2013–2017 action plan”), and the “Three-year Action Plan for Winning the Blue  
70 Sky War” from 2018–2020 (hereinafter called the “2018–2020 action plan”). With the successful implementation of these two  
71 action plans, the air quality was substantially improved in China, as evidenced in both observational and reanalysis datasets  
72 (Li et al., 2020b; Zheng et al., 2017; Krotkov et al., 2016; Zhong et al., 2021; Li et al., 2017a; Kong et al., 2021). However,  
73 with the deepening of air pollution control, unexpected changes have occurred in China, bringing about new challenges for the  
74 mitigation of air pollution in the future. On the one hand, despite a significant decline in PM<sub>2.5</sub> concentrations in China, severe  
75 haze still occasionally occurs during the wintertime (Zhou et al., 2022b; Li et al., 2017c). In addition, field measurements in  
76 cities over different regions of China consistently show different responses of aerosol chemical compositions to the emission  
77 control policies (Tang et al., 2021; Zhou et al., 2019; Wang et al., 2022; Zhang et al., 2020; Li et al., 2019a; Xu et al., 2019b;  
78 Lei et al., 2021; Zhou et al., 2022a). Compared with other aerosol species that showed substantial decreases during the clean  
79 air action plans, nitrate has shown a weaker response to the control measures, remaining at high levels and in some cases  
80 having even increased slightly. As a result, nitrate is playing an increasingly important role in heavy haze episodes in winter,  
81 and dominates the chemical composition of PM<sub>2.5</sub> (Fu et al., 2020; Xu et al., 2019a), leading to a rapid transition from sulphate-  
82 to nitrate-driven aerosol pollution (Li et al., 2019a; Wang et al., 2019b). On the other hand, photochemical pollution has  
83 deteriorated in China, with ozone (O<sub>3</sub>) concentrations having increased substantially in eastern China during 2013–2017 (Li et  
84 al., 2019b; Lu et al., 2018; Lu et al., 2020; Wang et al., 2020b).

85 These unexpected changes have raised considerable concern among the scientific community and policymakers regarding  
86 the overall effects of the clean air action plans, and how to coordinate the control of PM<sub>2.5</sub> and O<sub>3</sub> pollution. Addressing this  
87 problem requires a comprehensive understanding of the effects of the clean air action plans on the emissions of different air  
88 pollutantschemical species. In this respect, previous studies have compiled several long-term air pollutant emission inventories  
89 in China using the bottom-up approach – for example, the Multi-resolution Emission Inventory for China (MEIC) developed  
90 by Tsinghua University for 2010–2020 (Zheng et al., 2018); the Air Benefit and Cost and Attainment Assessment System-  
91 Emission Inventory version 2.0 (ABaCAS-EI v2.0) developed by Tsinghua University for 2005–2021 (Li et al., 2023); the  
92 Regional Emission Inventory in Asia (REAS) for 1950–2015 developed by Kurokawa and Ohara (2020); the Emissions  
93 Database for Global Atmospheric Research (EDGAR) for 1970–2018 developed by Jalkanen et al. (2012); the Hemispheric  
94 Transport of Air Pollution (HTAP) Inventory for 2000–2018 developed by Crippa et al. (2023); and the Community Emissions  
95 Data System (CEDS) Inventory for 1970–2019 developed by McDuffie et al. (2020). These emission inventories have provided  
96 the community with important insights into the long-term changes in the air pollutant emissions ~~of different air pollutants~~ in  
97 China, thus playing an indispensable role in our understanding of the effects of the country’s clean air action plans on emissions  
98 and air quality. However, due to the lack of accurate activity data and emission factors, bottom-up emission inventories are  
99 subject to large uncertainties, particularly during the clean air action periods when the activity data and emission factors  
100 changed considerably and were difficult to track. Consequently, the estimated emission rates from different bottom-up  
101 emission inventories could differ by more than a factor of 2 (Elguindi et al., 2020). For example, the estimated emissions for  
102 the year 2010 from different bottom-up inventories were 104.9–194.5 Tg for carbon monoxide (CO), 15.6–25.4 Tg for nitrogen  
103 oxides (NO<sub>x</sub>), 22.9–27.0 Tg for non-methane volatile organic compounds (NMVOCs), 15.7–35.5 Tg for sulfur dioxide (SO<sub>2</sub>),  
104 1.28–2.34 Tg for black carbon (BC), and 2.78–4.66 Tg for organic carbon (OC), reflecting the large uncertainty in current  
105 bottom-up estimates of air pollutant emissions in China, which hinders the proper assessment of the effects of the clean air  
106 action plans.

107 Inverse modeling of multiple air pollutant emissions (i.e., a top-down approach) provides an attractive way to constrain  
108 bottom-up emissions by reducing the discrepancy between the model and observation through the use of data assimilation.  
109 Numerous studies have confirmed the effectiveness of such a top-down method in verifying bottom-up emission estimates and  
110 reducing their uncertainties (e.g., Elbern et al., 2007; Henze et al., 2009; Miyazaki and Eskes, 2013; Tang et al., 2013; Koohkan  
111 et al., 2013; Koukouli et al., 2018; Jiang et al., 2017; Muller et al., 2018; Paulot et al., 2014; Qu et al., 2017). Based on long-  
112 term satellite observations, the top-down method has also been used to track the long-term variations of emissions. For example,  
113 Zheng et al. (2019) estimated the global emissions of CO for the period 2000–2017 based on a multi-species atmospheric  
114 Bayesian inversion approach; Qu et al. (2019) constrained global SO<sub>2</sub> emissions for the period 2005–2017 by assimilating  
115 satellite retrievals of SO<sub>2</sub> columns using a hybrid 4DVar/mass balance emission inversion method; by assimilating satellite  
116 observations of multiple species, Miyazaki et al. (2020b) simultaneously estimated global emissions of CO, NO<sub>x</sub>, and SO<sub>2</sub> for  
117 the period 2005–2018; and, most recently, a regional top-down estimation of PM<sub>2.5</sub> emissions in China during 2016–2020 was  
118 carried out by Peng et al. (2023) by assimilating surface observations. These studies provide us with valuable clues for  
119 evaluating bottom-up emissions and improving our knowledge on the changes in emissions of different species in China during  
120 the clean air action plans. However, most of these studies focused on emission trends at the global scale, which involved the  
121 use of coarse model resolutions (>1°) that may be insufficient to capture the spatial variability of emission variations at the  
122 regional scale. Meanwhile, current long-term, top-down estimates mainly focus on single species and do not fully cover the  
123 two clean air action periods in China. Indeed, to date, there are still no long-term, top-down estimates of major air pollutant  
124 emissions in China that fully cover the two clean air action periods.

125 In a previous study performed by our group, we developed a high-resolution air quality reanalysis dataset over China  
126 (CAQRA) for the period 2013–2020 to track the air quality trends in China during the clean air action periods (Kong et al.,  
127 2021). In the present study, as a follow up to this work, we constrained the long-term emission trends of major air pollutants

128 in China for 2013–2020 (which will be extended in the future on a yearly basis) by assimilating surface observations of air  
129 pollutants from the China National Environmental Monitoring Centre (CNEMC) using an ensemble Kalman filter and the  
130 Nested Air Quality Prediction and Forecasting System (NAQPMS). In the following sections, we present detailed descriptions  
131 of the chemical data assimilation, the evaluation results of the inversed emission inventory, and the estimated emission trends  
132 of different air pollutants in China during the clean air action periods.

## 133 2 The chemical data assimilation system

134 We used the chemical data assimilation system (ChemDAS) developed by the Institute of Atmospheric Physics, Chinese  
135 Academy of Sciences, to constrain the long-term emission changetrends of different air pollutants in China, which was used  
136 in the development of CAQRA in our previous work (Kong et al., 2021). Since the chemical transport model (CTM) and the  
137 observations used in the top-down estimation were the same as those used in CAQRA, we only briefly describe these two  
138 components in the following two subsections, instead concentrating on providing a fuller description (in the third subsection)  
139 of the inversion scheme in ChemDAS.

### 140 2.1 Chemical transport model

141 The NAQPMS model was used as the forecast model to represent the atmospheric chemistry in this study, and the Weather  
142 Research and Forecasting (WRF) model was used as the meteorological model to provide the meteorological input data.  
143 NAQPMS contains comprehensive modules for the emission, diffusion, transportation, deposition, and chemistry processes in  
144 the atmosphere, and has been used in previous inversion studies (Tang et al., 2013; Kong et al., 2019; Wu et al., 2020a; Kong  
145 et al., 2023). Detailed configurations of the different modules used in NAQPMS are available in these publications.

146 Figure 1 shows the domain of the inverse model, which is the same as that used in CAQRA, with a fine-scale horizontal  
147 resolution of 15 km. The HTAPv2.2 emission inventory was used as the a priori estimate of anthropogenic emissions in China,  
148 which includes emissions from the energy, industry, transport, residential, agriculture, air and ship sectors with a base year of  
149 2010 (Janssens-Maenhout et al., 2015). It is a harmonized global emission inventory that comprises of different regional  
150 gridded inventories. Within the region of China, the air pollutant emissions were mainly provided by the MEIC emission  
151 inventory (Janssens-Maenhout et al., 2015). The a priori estimates of emissions from other sourcesinventory includes the  
152 anthropogenic emissions obtained from the HTAP v2.2 emission inventory that provides the emissions from energy, industry,  
153 transport, residential, agriculture, air and ship sectors with a base year of 2010 (Janssens Maenhout et al., 2015); biogenic  
154 emissions obtained from the Monitoring Atmospheric Composition and Climate (MACC) project (Sindelarova et al., 2014);  
155 biomass burning emissions obtained from the Global Fire Emissions Database (GFED), version 4 (Van Der Werf et al., 2010;  
156 Randerson et al., 2017); soil and lightning NO<sub>x</sub> emissions obtained from Yan et al. (2003) and Price et al. (1997); and marine  
157 volatile organic compound emissions obtained from the POET database (Granier et al., 2005). The dust emissions were  
158 calculated online in NAQPMS as a function of the relative humidity, frictional velocity, mineral particle size distribution, and  
159 the surface roughness (Li et al., 2012), while the sea salt emissions were calculated using the scheme of Athanasopoulou et al.  
160 (2008). Note that since we aimed to estimate the air pollutant emissions and their changes from the surface observation, we  
161 did not consider the temporal variation in the *a priori* emission inventory. This would ensure that–so–that the top-down  
162 estimated emission trends were only derived from the surface observations, without being influenced by the trends in the prior  
163 emission inventory. In this way, our top-down estimation can serve as an independent estimation of the air pollutant emission  
164 changes in China. MeanwhileIn addition, we used the constant diurnal variation of the emissions in this study due to the lack  
165 of information on the diurnal variation of the emissions from different sectors, which is a potential limitation in our current  
166 work. However, since the emission inversion was performed on the daily basis (Sect. 2.3.3), the diurnal variations of the  
167 emission may not significantly influence the simulation results of the daily mean concentrations of air pollutants (less than 1

168 ppbv for SO<sub>2</sub>, NO<sub>2</sub> and O<sub>3</sub>) according to the sensitivity experiments conducted by Wang et al. (2010). The initial condition  
169 was treated as clean air in NAQPMS, with a 2-week spin-up time. Top and boundary conditions were provided by the Model  
170 for Ozone and Related Chemical Tracers (MOZART) (Brasseur et al., 1998; Hauglustaine et al., 1998) data products provided  
171 by National Center for Atmospheric Research (NCAR). Note that since the MOZART data products were not available for  
172 years after 2018, the multi-year average results from 2013 to 2017 were used for the simulations after 2018. Because most of  
173 the model boundaries were set in the clean areas and are located at distance from China, we assumed that the differences in  
174 boundary conditions would not significantly affect the modeling results over the China. To improve the performance of  
175 meteorological simulation, a 36-h free run of the WRF model was conducted for each day by using the NCAR/NCEP 1°×1°  
176 reanalysis data. The simulation results of the first 12 h were treated as the spin-up run, and the remaining 24 h were used to  
177 provide the meteorological inputs for the NAQPMS model. The evaluation results for the WRF simulation are available in  
178 Text S1 in the Supplement, which suggests acceptable performance of the WRF simulation for the inversion estimates (Table  
179 S1).

## 180 2.2 Assimilated observations

181 The assimilated observational dataset in this study was the same as that used in CAQRA, which includes surface  
182 concentrations of PM<sub>2.5</sub>, PM<sub>10</sub> (coarse particulate matter), SO<sub>2</sub>, NO<sub>2</sub> (nitrogen dioxide), CO, and O<sub>3</sub>, from 2013 to 2020,  
183 obtained from CNEMC (Fig. 1). Before the assimilation, outliers of the observations were filtered out by using an automatic  
184 quality control method developed by Wu et al. (2018). Four types of outliers characterized by temporal and spatial  
185 inconsistencies, instrument-induced low variances, periodic calibration exceptions, and lower PM<sub>10</sub> concentrations than those  
186 of PM<sub>2.5</sub>, were filtered out to prevent adverse impacts on the inversion process. As estimated in Kong et al. (2021), about 1.5%  
187 of observational data were filtered out after quality control, but further assessment showed that it had few effects on the average  
188 concentrations of different species, which were estimated to be less than 1 µg/m<sup>3</sup> for the gaseous air pollutants and less than  
189 5 µg/m<sup>3</sup> for the particulate matter. Estimation of observation error is also important ~~for~~ the inversion of emissions since the  
190 observational error and background errors determine the degree of adjustment to the emissions. The observational error  
191 comprises the measurement error and the representativeness error induced by the different spatial scales that the model and  
192 observations represent. The estimations of these two components of observational error were the same as those used in CAQRA,  
193 detailed descriptions of which are available in Kong et al. (2021).

194 It should be noted that the number of observation sites were not constant throughout the whole inversion period, being  
195 approximately 510 in 2013 and then increasing to 1436 in 2015. According to Fig. S1, the observation sites were mainly  
196 concentrated in the megacity clusters ~~in China~~ (e.g., North China Plain, Yangtze River Delta and Pearl River Delta) and the  
197 capital cities of each province in 2013. The number of observation sites continued to increase across the China in 2014 and  
198 2015. In particular, many areas that were previously unobserved ~~in 2013~~ have added monitoring stations in 2014 and 2015,  
199 which significantly increased the observation coverage ~~of~~ China and could lead to spurious trends in the top-down estimated  
200 emissions. Figure 2 shows the changes in the observational coverage over different regions of China from 2013 to 2020  
201 indicated by the ratio of areas that were influenced by observations to the total area of each region. To investigate the potential  
202 impacts of this on the top-down estimations, the changes in the coverage of observations over different regions of China from  
203 2013 to 2020 were firstly calculated by the ratio of areas that were influenced by observations to the total area of each region  
204 (Fig. 2). It can be clearly seen that the observational coverage increased from 2013 to 2015 with the expansion of the air quality  
205 monitoring network in China, and became stable after 2015. However, the influence of the variation in the number of  
206 observation sites varied among different regions. Over the North China Plain (NCP) region, the observational coverage was  
207 approximately 90% in 2013, and reached 100% in 2014, suggesting that the variation in the observation sites may have little  
208 influence on the estimated emission changes ~~in emissions~~ there. A similar conclusion can be drawn for the Southeast China  
209 (SE) region, where the observational coverage was about 75% in 2013 and reached 100% in 2015. Elsewhere, in the other four

210 regions, the influence of the variation in observation sites is expected to be larger because of the low observational coverage  
 211 in both 2013 and 2014. For example, the observational coverage over the Northwest China (NW) region was less than 10% in  
 212 2013, but increased to about 60% in 2015. To better illustrate the impact of changes in observational coverage on the inversions,  
 213 a sensitivity analysis of the emission increments with the fixed observation sites or varying observation sites is performed  
 214 in this study (Text S2† and Fig. S2). It shows that the additional emission increments caused by the increases of observation  
 215 sites would weaken the decreasing trends estimated in the fixed-site scenario for the emissions of PM<sub>2.5</sub>, NO<sub>x</sub> and NMVOC  
 216 and even lead to increasing trends for the emissions of PM<sub>10</sub> and CO. In contrast, the increases of observation sites would  
 217 enhance the decreasing trends of SO<sub>2</sub> estimated in the fixed-site scenario. Such different behaviors are mainly related to the  
 218 different sign of the emission increment of different species as we illustrated in Text S2†. These results highlighted the  
 219 significant influences of the site differences on the estimated emissions and their trends, which should be noted by the potential  
 220 users. Therefore, in order to reduce this influence on the estimated emission trends, in our following analysis we mainly  
 221 analyze the emission trends after 2015, when the observational coverage had stabilized in all regions.

## 222 2.3 Data assimilation algorithm

223 We used the modified EnKF coupled with state augmentation method to constrain the long-term emissions of different  
 224 air pollutants. EnKF is an advanced data assimilation method originally proposed by Evensen (1994) that features representing  
 225 the background error covariance matrix with a stochastic ensemble of model realizations. Through the use of ensemble  
 226 simulations, it has the ability to consider the indirect relationship between the emissions and chemical concentrations caused  
 227 by the complex physical and chemical processes in the atmosphere. It also allows for the estimation of flow-dependent  
 228 emission–concentration relationships that vary in time and space depending on the atmospheric conditions. The modified  
 229 EnKF is an offline application of the EnKF method that works by decoupling the analysis step from the ensemble simulation,  
 230 which has benefits in the reuse of costly ensemble simulations and makes high-resolution long-term inversion affordable (Wu  
 231 et al., 2020a). In this method, the ensemble simulation was performed firstly with the perturbed emissions, and then the  
 232 observations were assimilated to constrain the emissions (Wu et al., 2020a). The state augmentation method is a commonly  
 233 used parameter estimation method (Tandeo et al., 2020) in which the air pollutant emissions are taken as the state variable and  
 234 are updated according to the error covariance between the emissions and the concentrations of related species.

### 235 2.3.1 State variable and ensemble generations

236 The state variable used in this study was chosen following our previous multi-species inversion study (Kong et al., 2023),  
 237 which included the scaling factors for the emissions of fine-mode unspiciated aerosol (PMF), coarse-mode unspiciated aerosol  
 238 (PMC), BC, OC, NO<sub>x</sub>, SO<sub>2</sub>, CO, and NMVOC, as well as the chemical concentrations of PM<sub>2.5</sub>, PM<sub>10-2.5</sub> (PM<sub>10</sub> minus PM<sub>2.5</sub>),  
 239 NO<sub>2</sub>, SO<sub>2</sub>, CO, and daily maximum 8-h O<sub>3</sub> (MDA8h O<sub>3</sub>), which are formulated as follows:

$$240 \mathbf{x} = [\mathbf{c}, \boldsymbol{\beta}]^T, \quad (1)$$

$$241 \mathbf{c} = [\text{PM}_{2.5}, \text{PM}_{10-2.5}, \text{NO}_2, \text{SO}_2, \text{CO}, \text{MDA8h O}_3], \quad (2)$$

$$242 \boldsymbol{\beta} = [\boldsymbol{\beta}_{\text{PMF}}, \boldsymbol{\beta}_{\text{PMC}}, \boldsymbol{\beta}_{\text{BC}}, \boldsymbol{\beta}_{\text{OC}}, \boldsymbol{\beta}_{\text{NO}_x}, \boldsymbol{\beta}_{\text{SO}_2}, \boldsymbol{\beta}_{\text{CO}}, \boldsymbol{\beta}_{\text{NMVOC}}], \quad (3)$$

243 where  $\mathbf{x}$  denotes the vector of the state variable,  $\mathbf{c}$  denotes the vector of the chemical concentrations of different species, and  
 244  $\boldsymbol{\beta}$  denotes the vector of the scaling factors for the emissions of different species. Note that although the chemical concentration  
 245 variables are included in the state variable, they are not optimized simultaneously with the emission in the analysis step and  
 246 are only used to estimate the covariance between the emission and concentrations. Detailed descriptions of the state variables  
 247 are available in Table 1.

248 The ensemble of the scaling factors for different species was generated independently using the same method of Kong et  
 249 al. (2021), which has a medium size of 50 and considers the uncertainties of major air pollutant emissions in China, including  
 250 SO<sub>2</sub>, NO<sub>x</sub>, CO, NMVOCs, ammonia, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, and OC. The uncertainties of these species were considered to be 12%,

251 31%, 70%, 68%, 53%, 132%, 130%, 208% and 258%, respectively according to the estimates of Li et al. (2017b) and Streets  
 252 et al. (2003). Note that in this study we did not perturb the emissions of different sectors to reduce the degrees of freedom in  
 253 the ill-posed inverse estimation problem. Instead, we only perturbed the total emissions of different species. Therefore, only  
 254 the total emissions of different species were constrained in this study. The ensemble of the chemical concentrations was then  
 255 generated through an ensemble simulation based on NAQPMS and the perturbed emissions calculated by multiplying the *a*  
 256 *priori* emissions by the ensemble of the scaling factors. This treatment implicitly assumes that the uncertainty in the chemical  
 257 concentration is mainly caused by the emission uncertainty. This makes sense on a monthly or yearly basis, considering that  
 258 substantial changes in emissions are expected to have taken place during the clean air action plans, which are subject to large  
 259 uncertainty. However, the lack of consideration of other error sources, such as those of the meteorological simulation and the  
 260 model itself, may lead to underestimation of the background error covariance and ~~overcorrection of the emissions adjustment~~,  
 261 which is a potential limitation of this study. In addition, the dust and sea salt emissions were not perturbed and constrained in  
 262 this study, and thus the errors in the simulated fine and coarse dust emissions would influence the inversion of PM<sub>2.5</sub> and PM<sub>10</sub>  
 263 emissions. As a result, the top-down estimated PM<sub>2.5</sub> and PM<sub>10</sub> emissions will contain errors in the simulated dust and sea salt  
 264 emissions. Particularly, we did not consider the emissions of coarse dust during the inversion process since there is large  
 265 uncertainty in the simulated coarse dust emissions by current dust emission schemes (Zeng et al., 2020; Kang et al., 2011).  
 266 The we found large errors in the simulated coarse dust concentration ~~that could have significantly influenced~~ the inversion  
 267 results of PM<sub>10</sub> emissions. For example, the simulated coarse dust concentration could sometimes be several orders of  
 268 magnitude higher than the observed PM<sub>10</sub> concentration, leading to too low values of the inverse PM<sub>10</sub> emissions  
 269 (approximately 0) over the regions that were not the typical dust source regions but were influenced by the transportation of  
 270 coarse dust. Therefore, we only used simulated PM<sub>10</sub> concentrations from other sources in the inversion of PM<sub>10</sub> emissions to  
 271 avoid the influences of the too large errors in simulated. Consequently This is also similar to assume that the coarse dust  
 272 emission is equal to zero during the assimilation. However, in this way, the top-down estimated PM<sub>10</sub> emissions in this study  
 273 would comprise all coarse dust emissions which should be noted by potential users. A detailed description of the ensemble  
 274 generation is available in Kong et al. (2021).

### 275 2.3.2 Inversion algorithm

276 We used a deterministic form of EnKF (DEnKF) proposed by Sakov and Oke (2008) to update the scaling factors of the  
 277 emissions of different species, which is formulated as follows:

$$278 \bar{\mathbf{x}}^a = \bar{\mathbf{x}}^b + \mathbf{K}(\mathbf{y}^o - \mathbf{H}\bar{\mathbf{x}}^b), \quad (4)$$

$$279 \mathbf{X}^a = \mathbf{X}^b - \frac{1}{2} \mathbf{K} \mathbf{H} \mathbf{X}^b \quad (5)$$

$$280 \mathbf{K} = \lambda \mathbf{B}_e^b \mathbf{H}^T (\mathbf{H} \lambda \mathbf{B}_e^b \mathbf{H}^T + \mathbf{R})^{-1}, \quad (6)$$

$$281 \mathbf{B}_e^b = \frac{1}{N-1} \sum_{i=1}^N \mathbf{X}_i^b (\mathbf{X}_i^b)^T, \quad (7)$$

$$282 \bar{\mathbf{x}}^b = \frac{1}{N} \sum_{i=1}^N \mathbf{x}_i^b; \mathbf{X}_i^b = \mathbf{x}_i^b - \bar{\mathbf{x}}^b, \quad (8)$$

283 where  $\bar{\mathbf{x}}$  denotes the ensemble mean of the state variable; the superscript **b** and **a** respectively denote the *a priori* and *a*  
 284 *posteriori* estimate;  $\mathbf{X}^a$  is the analysed anomalies that can be used to calculate the uncertainty of the *a posteriori* emissions.  $\mathbf{K}$   
 285 is the Kalman gain matrix;  $\mathbf{B}_e^b$  is the background error covariance matrix calculated by the background perturbation  $\mathbf{X}^b$ ;  $\mathbf{y}^o$  is  
 286 the vector of the observation and  $\mathbf{R}$  is the observation error covariance matrix;  $\mathbf{H}$  is the linear observation operator, which  
 287 maps the model space to the observation space;  $\lambda$  is the inflation factor used to compensate for the underestimation of the  
 288 background error caused by the limited ensemble size and unaccounted error sources, which is calculated using the method of  
 289 Wang and Bishop (2003),

$$290 \lambda = \frac{(\mathbf{R}^{-1/2} \mathbf{d})^T \mathbf{R}^{-1/2} \mathbf{d} - p}{\text{trace}\{\mathbf{R}^{-1/2} \mathbf{H} \mathbf{B}_e^b (\mathbf{R}^{-1/2} \mathbf{H})^T\}} \quad (9)$$

$$\mathbf{d} = \mathbf{y}^o - \mathbf{H}\mathbf{x}^b \quad (10)$$

where  $\mathbf{d}$  is the observation innovation and  $\mathbf{p}$  is the number of observations. Table S21 summarized the calculated average value (standard deviation) of the used inflation factor for different species. It shows that the inflation factor over the east China (including NCP and SE region) was generally round 1.0, suggesting that the original ensemble can well represent the simulation errors of the different air pollutants over these regions. The inflation factor is larger over the western China (including SW, NW and Central regions), especially for PM<sub>10</sub> (36.0–78.1) and SO<sub>2</sub> (7.8–176.1), suggesting that the original ensemble may underestimate the simulation errors of the air pollutants. This is associated with the large biases in the simulated air pollutant concentrations over there and reflect that the emission uncertainties assumed in our studies may be underestimated over these regions. This also highlighted the importance of the use of inflation method during the inversion, otherwise it would lead to filter divergency caused by the underestimations of the background error covariance.

In order to reduce the influence of the spurious correlations on the performance of data assimilation, the EnKF was performed locally in this study in that the analysis was calculated grid by grid with the assumption that only measurements located within a certain distance (cutoff radius) from a grid point would influence the analysis results of this grid. The use of this local analysis method also allowed the inflation factor to be calculated locally and to vary in time and space, which can help characterize the spatiotemporal variations of errors as we illustrated above. Similar to in Kong et al. (2021) and Kong et al. (2023), the cutoff radius was chosen as 180 km for each species based on the wind speed and the lifespan of the species (Feng et al., 2020). The same local scheme with a buffer area was also employed during the inversion to alleviate the discontinuities in the updated state caused by the cut-off radius. A detailed description of the local analysis scheme is available in Kong et al. (2021).

Table 1 summarizes the corresponding relationships between the emissions and chemical concentrations. Similar to Ma et al. (2019) and Miyazaki et al. (2012), we did not consider the inter-species correlation during the assimilation, to prevent the spurious correlations between non- or weakly related variables. In most cases, observations of one particular species were only allowed to adjust the emissions of the same species. The assimilation of PM<sub>2.5</sub> mass observation was more complicated as there are multiple error sources in the simulated mass concentrations of PM<sub>2.5</sub>, not only from primary emission, but also from secondary production. In this study, the PM<sub>2.5</sub> mass observation was used to constrain the emissions of PMF, BC and OC but not used to constrain the emissions of its precursors to avoid the spurious correlations and nonlinear chemistry effects, which is similar to the scheme used in Ma et al. (2019). This is feasible since the emissions of primary PM<sub>2.5</sub> (i.e., PMF, BC and OC) and the emissions of PM<sub>2.5</sub> precursors (e.g., SO<sub>2</sub>, NO<sub>2</sub>) were perturbed independently in our method, thus the contributions of primary PM<sub>2.5</sub> emission and the secondary PM<sub>2.5</sub> productions to the PM<sub>2.5</sub> mass could be isolated through the use of ensemble simulations. Meanwhile, the use of iteration inversion method (which will be introduced later) can further reduce the influence of the errors in the precursors' emissions on the inversion of primary PM<sub>2.5</sub> emission, because the errors of its precursors' emission would be constrained by their own observations during the iterations. However, the lack of assimilation of speciated PM<sub>2.5</sub> observations may lead to uncertainties in the estimated emissions of PMF, BC and OC, which is a potential limitation in current work. For example, if the a priori simulated PM<sub>2.5</sub> equals the observations, the emissions of PMF, BC and OC would not be adjusted by using the current method. However, in such cases, there may still be discrepancies in the proportions of the emissions of different PM<sub>2.5</sub> components. To adjust the emissions of PMF, we used the observations of PM<sub>10-2.5</sub> to avoid the potential cross-correlations between PM<sub>2.5</sub> and PM<sub>10</sub> (Peng et al., 2018; Ma et al., 2019). For the NO<sub>x</sub> emissions, although the O<sub>3</sub> concentration are chemically related to the NO<sub>x</sub> emissions, we did not use the O<sub>3</sub> concentrations to constrain the NO<sub>x</sub> emission in this study since there is nonlinear relationship between the O<sub>3</sub> concentration and NO<sub>x</sub> emission which would lead to wrong adjustment of NO<sub>x</sub> emissions (Tang et al., 2016).

The inversion of NMVOC emission is more difficult than other species due to the lack of long-term nationwide NMVOC observations and the strong chemical activity. Previous studies usually assimilated the satellite observations of formaldehyde and glyoxal to constrain the NMVOC emissions, such as Cao et al. (2018) and Stavrou et al. (2015). However, these



334 inversion studies are hindered by the  $\text{NO}_x$ -VOC- $\text{O}_3$  chemistry and the inherent uncertainty in the satellite observations of  
335 formaldehyde and glyoxal. Considering the strong chemical relationship between the  $\text{O}_3$  and NMVOC, some pioneer studies  
336 have also explored the method of assimilating ground-level  $\text{O}_3$  concentrations to constrain the NMVOC emissions (Ma et al.,  
337 2019; Xing et al., 2020), and demonstrated the effectiveness of this approach. For example, Ma et al. (2019) found that the  
338 assimilation of  $\text{O}_3$  concentration could adjust the NMVOC emissions in the direction resembling the bottom-up inventories,  
339 and the forecast skill of  $\text{O}_3$  concentrations were also improved, indicating that the constrained NMVOC emissions are improved  
340 relative to their a priori. Inspired by these studies, we have made an attempt to constrain the NMVOC emissions based on the  
341 MDA8h  $\text{O}_3$ . The use of MDA8h  $\text{O}_3$  rather than the daily mean  $\text{O}_3$  concentration is to avoid the effects of the nighttime  $\text{O}_3$   
342 chemistry. For example, the simulation errors in the titration effects of  $\text{NO}_x$  may influence the simulated  $\text{O}_3$  concentrations  
343 during nighttime and affect the inversion results of NMVOC. An important issue that should be noted when using the MDA8h  
344  $\text{O}_3$  to constrain the NMVOC emission is the nonlinear interactions among  $\text{NO}_x$ , NMVOC and  $\text{O}_3$ . On the one hand, the  $\text{O}_3$   
345 concentrations are dependent not only on the NMVOC emissions but also on the  $\text{NO}_x$  emissions. The errors in the a priori  
346 emissions of  $\text{NO}_x$  would also contribute to the simulation errors of  $\text{O}_3$ , and deteriorate the inversion of NMVOC. The iteration  
347 inversion scheme could help deal with this issue as the errors in the  $\text{NO}_x$  emissions will be constrained by the  $\text{NO}_2$  observations  
348 in the next iteration, which can reduce the influences of errors in the  $\text{NO}_x$  emission on the inversion of NMVOC emission  
349 based on MDA8h  $\text{O}_3$  concentrations. This is in fact similar to the approach used by Xing et al. (2020) who firstly constrained  
350 the  $\text{NO}_x$  emissions based on observations of  $\text{NO}_2$ , and then constrained the NMVOC emissions based on  $\text{O}_3$  concentrations.  
351 Also, in Feng et al. (2024), the  $\text{NO}_2$  observations were simultaneously assimilated to constrain the  $\text{NO}_x$  emissions to account  
352 for the influences of errors in  $\text{NO}_x$  emissions on the NMVOC emissions, suggesting that the iteratively nonlinear joint inversion  
353 of  $\text{NO}_x$  and NMVOCs is an effective way to address the intricate relationship among VOC- $\text{NO}_x$ - $\text{O}_3$  (Feng et al., 2024).  
354 Similarly, the errors in the CO emissions which may be significant according to our following analysis are also constrained in  
355 a similar way to reduce the potential influences on the inversion of NMVOC emission. On the other hand, the emission  
356 adjustments of NMVOC may exhibit bidirectionality dependent on the VOC-limited or  $\text{NO}_x$ -limited regimes. According to  
357 Fig. 3, the NMVOC emissions were adjusted in alignment with the direction of the  $\text{O}_3$  errors, suggesting a VOC-limited regime  
358 over urban areas in China, given that the  $\text{O}_3$  observation sites are predominantly situated in the urban areas. This agrees with  
359 Ren et al. (2022) who diagnosed the  $\text{NO}_x$ -VOC- $\text{O}_3$  sensitivity based on the satellite retrievals and found that the VOC-limited  
360 regimes are mainly located in the urban areas in China. This suggests that the relationship between the  $\text{O}_3$  concentrations and  
361 VOC emissions could be reasonably reflected by our inversion system, providing the feasibility in utilizing the  $\text{O}_3$  observations  
362 to constrain the VOC emissions. Note that due to the lack observations of the VOC components, we only optimize the gross  
363 emissions of the VOC during the assimilation.

364 As we illustrated before, there exists nonlinear effects in the atmospheric chemistry which could influence the inversion  
365 results of different species. In addition, since we did not consider the temporal variations in the a priori emissions, it was  
366 expected that there would be significant biases in the a priori emissions for the years after 2013, as substantial changes in  
367 emissions were expected owing to the implementation of strict emission control measures. Such bias in the a priori emissions  
368 does not conform to the unbiased hypothesis assumption of the EnKF ~~that the a priori estimate is unbiased~~, which could ~~thus~~  
369 lead to incomplete adjustments of the a priori emissions and degrade the performance of the data assimilation (Dee and Da  
370 Silva, 1998). To address these issues, an iteration inversion scheme was employed in this study, which has been used previously  
371 in Kong et al. (2023). The main idea of the iteration inversion scheme is to preserve the background perturbation  $\mathbf{X}^b$  but to  
372 update the ensemble mean of the state variable  $\bar{\mathbf{x}}^b$  based on ~~the inversion results of the kth iteration and corresponding the~~  
373 model simulations driven by the inversion results of the kth iteration. ~~Therefore According to this~~, a new single model  
374 simulation is required to be conducted by using the a posteriori emission from the previous iteration as the input to update the  
375 ensemble mean of the original ensemble. This enables the observational information and the adjusted emissions to be promptly  
376 incorporated into the model, thereby providing feedback for the adjustments of emission in the next iteration. However, we

377 did not reassemble the ensemble simulation for each iteration due to the expensive computational cost of the ensemble  
378 simulation. Therefore, in each iteration calculation, the ensemble perturbation that were used to calculate the background error  
379 covariance matrix remains the same with only the ensemble mean being updated based on the inversion results of the previous  
380 iteration. The state variable used in the  $(k + 1)$ th inversions is then formulated as follows:

$$381 \mathbf{x}_i^{b,k+1} = [\mathbf{c}^k + \mathbf{c}_i^e - \bar{\mathbf{c}}^e, \boldsymbol{\beta}^k + \boldsymbol{\beta}_i^e - \bar{\boldsymbol{\beta}}^e]^T, \quad (11)$$

382 where  $\mathbf{c}^k$  represents the model simulations driven by using the inversed emissions of the  $k$ th iteration,  $\mathbf{c}_i^e$  represents the  $i$ th  
383 member of ensemble simulations with an ensemble mean of  $\bar{\mathbf{c}}^e$ ,  $\boldsymbol{\beta}^k$  represents the updated scaling factors at the  $k$ th iteration,  
384 and  $\boldsymbol{\beta}_i^e$  represents the  $i$ th member of the ensemble of scaling factors with a mean value of  $\bar{\boldsymbol{\beta}}^e$ . In each iteration, all emissions  
385 are updated simultaneously and tTwo rounds of iteration were conducted in this study based on our previous inversion study  
386 to maintain a balance between the inversion performance and the computational cost of the long-term inversions (Kong et al.,  
387 2023).

### 388 2.3.3 Setup of inversion estimation

389 Based on this inversion scheme, we firstly constrained the daily emissions of PMF, PMC, BC, OC, NO<sub>x</sub>, SO<sub>2</sub>, CO, and  
390 NMVOCs, from 2013 to 2020, based on the daily averaged observations of PM<sub>2.5</sub>, PM<sub>10-2.5</sub>, NO<sub>2</sub>, CO, and MDA8h O<sub>3</sub>.  
391 However, due the lack of enough speciated PM<sub>2.5</sub> observation, the model performance driven by the inverse emission for the  
392 BC, OC and primary unspciated PM<sub>2.5</sub> have not been thoroughly evaluated. It is thus currently unclear for the quality of the  
393 inverse emissions of BC, OC and primary unspciated PM<sub>2.5</sub>. Also, the lack of speciated PM<sub>2.5</sub> observations could lead to  
394 uncertainties in the estimated emissions of PMF, BC, and OC as we mentioned before. Considering this, similar to in Kong et  
395 al. (2023), although we made attempt to estimate the emissions of BC, OC and primary unspciated PM<sub>2.5</sub>, we have reservations  
396 about their inversion results and only provide the emissions of PM<sub>2.5</sub> (PMC+BC+OC) and PM10 (PM<sub>2.5</sub> + PMC) in current  
397 stage. In future, we will collect more speciated PM<sub>2.5</sub> observations to comprehensively quantify the accuracy of their inversion  
398 results, after which the emissions of these species would be released. Meanwhile, the speciated PM<sub>2.5</sub> observations could be  
399 assimilated under the current inversion framework. This could provide us with further constrains on the emissions of BC, OC  
400 and primary PM<sub>2.5</sub>. similar to in Kong et al. (2023), we only provide the emissions of PM<sub>2.5</sub> (PMF+BC+OC) and PM<sub>10</sub>  
401 (PM<sub>2.5</sub>+PMF) for the aerosol species since the lack of speciated PM<sub>2.5</sub> observations would lead to uncertainties in the estimated  
402 emissions of PMF, BC, and OC as we mentioned before. Meanwhile, as mentioned in subsection 2.3.1, the meteorological and  
403 model uncertainty were not considered in the ensemble simulation. Thus, the errors in the meteorological simulation would  
404 cause fluctuations in the daily emissions that could contaminate the inversion results and are difficult to isolate from the  
405 inherent variations of emissions (Tang et al., 2013). Considering this, the daily emissions were averaged to monthly values to  
406 reduce the influences of random model errors after the assimilation.

## 407 3 Performance of the chemical data assimilation system

### 408 3.1 Analysis of OmF and emission increment

409 The observation-minus-forecast (OmF) and emission increment (*a posteriori* emission minus *a priori* emission) were  
410 firstly analyzed to demonstrate the performance of the data assimilation. As shown in Fig. 3, the *a priori* simulation generally  
411 underestimated the PM<sub>2.5</sub> concentrations over the NCP, SE and SW regions (positive OmF values) during 2013–2014, but  
412 overestimated the PM<sub>2.5</sub> concentrations from 2016, reflecting the effects of the emission control measures during these years.  
413 In the NE, NW and central China (hereafter, “Central”) regions, obvious underestimation of the PM<sub>2.5</sub> concentration was found  
414 (positive OmF values) throughout almost the entire assimilation period. Similarly, the OmF values of PM<sub>10</sub> were positive  
415 throughout the whole assimilation period over all regions of China. In contrast, the OmF values for SO<sub>2</sub> were negative for most

416 regions, and the negative OmF values over the NCP region became larger as the years progressed, which reflects the effects  
417 of the emission control measures. The OmF for NO<sub>2</sub> reveals a seasonal variation over the NCP and SE regions, with  
418 ~~negative~~ values during summer and ~~positive~~ values during winter, while there were obvious positive OmF  
419 values over the NE, SW, NW and Central regions. In terms of CO, large positive OmF values were found over all regions of  
420 China, and there were decreasing trends in the OmF values of CO over different regions of China associated with the emission  
421 control policies during these years. The OmF values for O<sub>3</sub> were positive over most regions of China, except the NW region.  
422 These results provide us with valuable information on the potential deficiencies in the *a priori* emissions. However, since our  
423 inversion method did not differentiate between anthropogenic and natural emissions, the biases in the model simulation may  
424 also be attributable to the errors in natural emissions such as dust, especially over the major dust-source areas of China (e.g.,  
425 the NW and Central regions). In addition, the effects of emission control were not considered in the *a priori* emissions, which  
426 is another important contributor to the errors in the model simulation for the later years. Thus, the emission increments  
427 calculated by the assimilation should reflect the combined effects of errors in the anthropogenic and natural emissions, as well  
428 as the emission control.

429 The calculated emission increments were consistent with the OmF values for all species, which indicates that the data  
430 assimilation method can probably constrain the emissions based on the observations. According to Fig. 3, the emission  
431 increments were positive for PM<sub>2.5</sub> over the NE, NW and Central regions, for NO<sub>2</sub> over the NE, SW, NW and Central regions,  
432 and for PM<sub>10</sub>, CO and NMVOC over almost all regions throughout the assimilation period. In contrast, the emission increments  
433 were negative for the SO<sub>2</sub> emissions for most cases. Consistent with the OmF values, the emission increments were positive  
434 for PM<sub>2.5</sub> over the NCP, SE and SW regions during 2013–2014, but became negative from 2016 owing to the implementation  
435 of strict emission control measures. The emission increments for NO<sub>x</sub> also showed significant seasonal variation over the NCP  
436 and SE regions, being positive during winter and negative during summer. The *a posteriori* biases for the model simulations  
437 of different species were also plotted to assess the performance of the data assimilation. It can be clearly seen that the biases  
438 were substantially reduced for all species, and the calculated root-mean-square error (RMSE) reduced by 23.2–52.8% for PM<sub>2.5</sub>,  
439 19.9–37.8% for PM<sub>10</sub>, 36.4–77.3% for SO<sub>2</sub>, 18.3–25.2% for NO<sub>2</sub>, 29.9–40.5% for CO, and 4.4–26.1% for O<sub>3</sub> over the different  
440 regions of China, suggesting a good performance of the data assimilation system.

### 441 3.2 Evaluation of the inversion results

442 Table 2 shows the calculated evaluation statistics for the inversion at different temporal scales. It can be clearly seen that  
443 the model simulation with the *a posteriori* emission inventory reproduced well the magnitude and temporal variations of the  
444 different air pollutants in China, with calculated correlation coefficients of approximately 0.77, 0.72, 0.64, 0.67, 0.69 and 0.71,  
445 and normalized mean biases of approximately 4.5%, -4.6%, -9.0%, -3.9%, -8.8% and 9.5%, for the hourly concentrations of  
446 PM<sub>2.5</sub>, PM<sub>10</sub>, SO<sub>2</sub>, NO<sub>2</sub>, CO and O<sub>3</sub>, respectively. Moreover, the *a posteriori* model simulation achieved comparable accuracy  
447 with the air quality reanalysis data we developed in Kong et al. (2021) in terms of the RMSE, which was 32.4 μg·m<sup>-3</sup>, 53.1  
448 μg·m<sup>-3</sup>, 24.9 μg·m<sup>-3</sup>, 19.9 μg·m<sup>-3</sup>, 0.56 mg·m<sup>-3</sup> and 34.9 μg·m<sup>-3</sup>, respectively, for these species at the hourly scale. At the daily,  
449 monthly and yearly scales, the constrained model simulation performed better, with RMSEs of about 9.1–20.0 μg·m<sup>-3</sup> (PM<sub>2.5</sub>),  
450 18.5–31.6 μg·m<sup>-3</sup> (PM<sub>10</sub>), 11.5–16.0 μg·m<sup>-3</sup> (SO<sub>2</sub>), 8.1–12.8 μg·m<sup>-3</sup> (NO<sub>2</sub>), 0.28–0.39 mg·m<sup>-3</sup> (CO), and 14.2–26.1 μg·m<sup>-3</sup> (O<sub>3</sub>),  
451 which were respectively reduced by 56.7–67.3%, 49.2–52.1%, 68.8–72.8%, 36.3–39.8%, 47.0–58.0%, and 22.9–30.5%  
452 compared to the RMSEs of the *a priori* simulations. We also compared the model performance driven by the inverse inventory  
453 with that driven by more recent bottom-up inventories (MEIC and HTAPv3) by taking the simulation results of year 2020 as  
454 an example to give us a more objective understanding of the accuracy of the inverse emission inventory. It shows that the  
455 inverse emission generally achieves better performance in simulating the air pollutant concentrations in China than the MEIC  
456 and HTAPv3 (Table S3). It is also encouraging to find that the model performance driven by CAQIEI and MEIC-HTAPv3 is  
457 similar for PM<sub>2.5</sub>, PM<sub>10</sub>, and SO<sub>2</sub> over the NCP, NE, SE and SW regions, both significantly improved from the a priori emission

458 [inventory. This suggest that both the top-down and recent bottom-up emission inventories have good performance in capturing](#)  
459 [the emission changes of these species over these regions and they yield consistent estimations. Detailed information on the](#)  
460 [configurations of the model simulation results driven by MEIC-HTAPv3 and the comparisons results are available in Text S3.](#)  
461 [All t](#)These validation results ~~confirm~~ confirm the good performance of the data assimilation method and suggest that the  
462 inversed emissions inventory has the capability to reasonably represent the magnitude and long-term trends of the air pollutant  
463 emissions in China during 2013–2020.

## 464 **4 Results**

465 Based on the top-down estimation, the gridded emissions for PM<sub>2.5</sub>, PM<sub>10</sub>, SO<sub>2</sub>, CO, NO<sub>x</sub> and NMVOCs over China from  
466 2013 to 2020 were developed into what we have called the Inversed Emissions Inventory for Chinese Air Quality (CAQIEI).  
467 In the following sections, we first analyze the magnitude and seasonality of the air pollutant emissions in China by taking 2015  
468 as a reference year when the number of observation sites became stable. After that, the changes in emissions of different air  
469 pollutants from 2015 to 2020 are analyzed and compared between the two clean air action plans in China. Note that due to the  
470 impacts of the changes in observation coverage, it is difficult to estimate the overall emission reduction rates during the 2013–  
471 2017 action plan by using our inversion results. The emission change rates during 2015–2017 were then sampled in this study  
472 to assess the mitigation effects during the 2013–2017 action plan and to be compared with the emission change rates during  
473 2018–2020. Finally, CAQIEI is compared to the previous bottom-up and top-down emission inventories to validate our top-  
474 down estimation and identify the potential uncertainties in the current understanding of China’s air pollutant emissions.

### 475 **4.1 Top-down estimated Chinese air pollutant emissions in 2015**

476 The top-down estimated emissions of different species in 2015 are as follows: 25.2 Tg of NO<sub>x</sub>, 17.8 Tg of SO<sub>2</sub>, 465.4 Tg  
477 of CO, 15.0 Tg of PM<sub>2.5</sub>, 40.1 Tg of PM<sub>10</sub>, and 46.0 Tg of NMVOCs. Note that these values not only contain anthropogenic  
478 emissions but also natural (e.g., dust and biogenic NMVOC) emissions. Thus, the top-down estimated emissions of PM and  
479 NMVOCs were higher than those estimated by previous studies, as we mention in ~~the~~ following sections. Emission maps of  
480 all species in 2015 are shown in Fig. 4, and the calculated emissions of different species over different regions are presented  
481 in Table 3. According to Fig. 4, higher air pollutant emissions are widely distributed in the megacity clusters (e.g., NCP,  
482 Yangtze River Delta and Pearl River Delta) and the developed cities in China, reflecting the influences of human activities.  
483 NCP was the region with the largest emission intensity of air pollutants in China, contributing 5.1 Tg of NO<sub>x</sub>, 3.5 Tg of SO<sub>2</sub>,  
484 82.2 Tg of CO, 2.7 Tg of PM<sub>2.5</sub>, 8.7 Tg of PM<sub>10</sub> and 9.0 Tg of NMVOCs to the total emissions in China. The inversion results  
485 also demonstrate the contribution of natural sources to the air pollutant emissions, such as the soil NO<sub>x</sub> emissions and the  
486 biogenic NMVOC emission distributed in the Tibet Plateau region. In general, the majority of air pollutant emissions were  
487 located in eastern China (including the NCP, NE and SE regions), where the economy is relatively well developed, which in  
488 total accounted for 66.0% of NO<sub>x</sub>, 60.9% of SO<sub>2</sub>, 57.5% of CO, 60.4% of PM<sub>2.5</sub>, 60.5% of PM<sub>10</sub>, and 67.8% of NMVOC  
489 emissions in China. However, although the GDP of western China (including the SW, NW and Central regions) is less than  
490 one third that of eastern China, the top-down estimation indicates that the air pollutant emissions in western China could have  
491 accounted for about 32.2–42.5% of the total emissions, which reflects the low emission control levels over these regions.

492 Figure 5 shows the monthly variations of air pollutant emissions in China for year 2015. The monthly profile of NO<sub>x</sub>  
493 emissions was relatively flat among the six species. SO<sub>2</sub> and CO showed higher emissions during wintertime because of the  
494 enhanced residential emissions associated with higher coal consumption for heating during that time of year. Meanwhile, the  
495 emission factor for CO from vehicles in winter was also higher than in other seasons, due to additional emissions from the  
496 cold-start process (Kurokawa et al., 2013; Li et al., 2017b). PM<sub>2.5</sub> and PM<sub>10</sub> had higher emissions during winter and spring,  
497 which, on the one hand was due to the enhanced emissions from the residential and industrial sectors during wintertime (Li et

498 al., 2017b), whilst on the other hand was due to the enhanced dust emissions during the spring season (Fan et al., 2021).  
499 Emissions of NMVOCs exhibited strong monthly variations, with higher emissions mainly in summer because of the enhanced  
500 NMVOC emissions from biogenic sources.

## 501 4.2 Top-down estimated emission changes of different air pollutants

### 502 4.2.1 Emission changes of particulate matter

503 Figure 6 shows the top-down estimated emission changes of  $PM_{2.5}$  and  $PM_{10}$  over China during two clean air action  
504 periods. Both  $PM_{2.5}$  and  $PM_{10}$  emissions decreased substantially, by 44.3% and 21.2% respectively, from 2013 to 2020. On  
505 the contrary, the top-down estimates showed increases of  $PM_{2.5}$  and  $PM_{10}$  emissions in 2014 and 2015, but this would be a  
506 spurious trend caused by the changes of observation sites as we discussed in Text S24. Therefore, the emissions in 2013 and  
507 2014 were discarded to prevent the spurious trends. According to Fig. 6, the  $PM_{2.5}$  emissions decreased by 14.5% from 2015  
508 (15.0 Tg) to 2017 (12.8 Tg), and the reduction in emissions was roughly uniform throughout the period, which was about 8%  
509 compared to previous years. The  $PM_{10}$  emissions showed a smaller reduction rate (-7.2%) than that of  $PM_{2.5}$ , decreasing from  
510 40.1 Tg in 2015 to 37.2 Tg in 2017. Compared with the emission reduction rate during 2015–2017, both  $PM_{2.5}$  and  $PM_{10}$   
511 showed larger emission reduction rates during 2018–2020, estimated to be 27.2% and 25.5%, respectively. The emission  
512 reductions in each year were also larger, especially for  $PM_{10}$ . For example,  $PM_{2.5}$  and  $PM_{10}$  emissions reduced by about 19.3%  
513 and 14.0% in 2019 compared to 2018. This may have been due to that in addition to the strict controls imposed on the industrial  
514 and power sectors during the 2013–2017 action period, the residential emissions have been strengthened during the 2018–  
515 2020 action period. In particular, “coal-to-electricity” and “coal-to-gas” strategies were vigorously implemented in northern  
516 China during the 2018–2020 action to reduce coal consumption and related air pollutant emissions (Liu et al., 2016; Wang et  
517 al., 2020a). Thus, our inversion results confirm the effectiveness of the controls on residential emissions in terms of reducing  
518 the emissions of  $PM_{2.5}$  and  $PM_{10}$ . In addition, the control of non-point sources, such as blowing-dust emissions, was also  
519 strengthened during the 2018–2020 action period, which is consistent with the faster reduction of  $PM_{10}$  emissions during 2018–  
520 2020. The annual trends of  $PM_{2.5}$  and  $PM_{10}$  emissions were also calculated in China using the Mann–Kendall trend test and  
521 the Theil–Sen trend estimation method, the results of which are summarized in Table 4. The calculation of emission trends can  
522 help extend the existing emission datasets forward in time to produce up-to-date products. The top-down estimated trends of  
523  $PM_{2.5}$  and  $PM_{10}$  emissions were -1.4 and -2.6 Tg/year during 2015–2020, attributable to the strict emission control measures  
524 imposed during the two clean air action plans. As mentioned, the decreasing trends were larger during 2018–2020 (-1.5 and  
525 -4.6 Tg/year) than during 2015–2017 (-1.1 and -1.5 Tg/year).

526 On the regional scale (Fig. S3), it can be clearly seen that the  $PM_{2.5}$  emissions decreased consistently over all regions, by  
527 59.8% in NCP, 49.6% in SE, 39.5% in NE, 35.8% in SW, 33.2% in NW, and 41.0% in Central, from 2015 to 2020. The NCP  
528 region showed the largest reduction in emissions among the six regions, with its emission reduction rate being almost larger  
529 than 10% in each year. This is consistent with the strictest emission control policies having been imposed over the NCP region.  
530 The SE region showed a similar ~~emission reductions in emissions~~ to the NCP region, with its emission reduction rate being  
531 larger than 10% in most years. Obvious increases of  $PM_{2.5}$  emissions could be found over the NW region from 2013 to 2015  
532 owing to the increase in the number of observation sites in those years. After 2015,  $PM_{2.5}$  emissions generally decreased over  
533 the NW region, while there was a slight rebound in  $PM_{2.5}$  emissions in 2016 and 2018, possibly due to the influences of the  
534 errors in fine dust emission. The Central region showed different characteristics of emission changes to the other regions  
535 insofar as it showed little change in  $PM_{2.5}$  emissions during 2015–2018 but large reductions in 2019. This may be consistent  
536 with the control of emissions over the Fenwei Plain area (the part of the Central region where the emission intensity is largest)  
537 being weak during the 2013–2017 action plan but strengthened during the 2018–2020 action plan. In terms of the  $PM_{2.5}$   
538 emission trends over the different regions, the calculated  $PM_{2.5}$  emission trends were about -0.32 Tg/year in NCP, -0.32

539 Tg/year in SE, -0.24 Tg/year in NE, -0.21 Tg/year in SW, -0.09 Tg/year in NW, and -0.15 Tg/year in Central, from 2015 to  
540 2020.

541 The changes of PM<sub>10</sub> emissions were generally similar to those of PM<sub>2.5</sub>, i.e., with decreases in all regions from 2015 to  
542 2020 (Fig. S4). The top-down estimated PM<sub>10</sub> emission reductions from 2015 to 2020 were about 3.5 Tg (40.0%) in NCP, 2.6  
543 Tg (35.5%) in SE, 3.0 Tg (36.6%) in NE, 2.0 Tg (35.9%) in SW, 1.0 Tg (25.3%) in NW, and 1.3 Tg (21.6%) in Central; and  
544 the calculated trends were about -0.64 Tg/yr, -0.52 Tg/yr, -0.51 Tg/yr, -0.40 Tg/yr, -0.20 Tg/yr, and -0.27 Tg/yr,  
545 respectively. However, due to the influences of the changes in the number of observation sites, the PM<sub>10</sub> emissions over the  
546 NE, SW and NW regions increased substantially from 2013 to 2015, while they decreased in almost all years after 2015.  
547 Different from the other regions, the Central region showed increases in PM<sub>10</sub> emissions from 2015 to 2018, by about 0.92 Tg  
548 (14.9%), but substantial decreases in 2019 and 2020. The result also shows that most PM<sub>10</sub> emission reductions were achieved  
549 during the 2018–2020 action plan. According to CAQIEI, the PM<sub>10</sub> emissions decreased by 0.64–2.3 Tg (17.4–31.8%) from  
550 2018 to 2020, which accounted for 48.4–169.0% of the total reduction in emissions from 2015 to 2020. This again emphasizes  
551 the effectiveness of the control of blowing-dust emissions during the 2018–2020 action plan.

## 552 4.2.2 Emission changes of gaseous air pollutants

### 553 4.2.2.1 SO<sub>2</sub> and CO

554 Figure 7 shows the emission changes of different gaseous air pollutants in China from 2013 to 2020. Similar to the PM  
555 emissions, SO<sub>2</sub> and CO emissions decreased continuously during the two action plan periods, with top-down estimated  
556 emission reductions of about 9.6 Tg (54.1%) and 166.3 Tg (35.7%) for SO<sub>2</sub> and CO from 2015 to 2020, respectively.  
557 Meanwhile, both SO<sub>2</sub> and CO showed a significant decreasing trend from 2015 to 2020, with estimated trends of approximately  
558 -2.1 Tg/yr and -36.0 Tg/yr, respectively (Table 5). The reductions in SO<sub>2</sub> and CO emissions are closely consistent with the  
559 strict emission control measures imposed during the action plan periods, such as the phasing out of outdated industrial capacity  
560 and high-emitting factories, the strengthening of emission standards for industry and the power sector, the elimination of small  
561 coal-fired industrial boilers, and the replacement of coal with cleaner energies, which reflects the effectiveness of the emission  
562 control measures during the two action plan periods. Reductions of SO<sub>2</sub> emission were generally steady during the two action  
563 plan periods, which were approximately 4.2 Tg (23.8%) from 2015 to 2017 and 2.5 Tg (23.5%) from 2018 to 2020. However,  
564 CO showed a different emission reduction rate during the two action plan periods, with its emission reductions (67.1 Tg, 18.3%)  
565 during 2018–2020 being larger than those (45.6 Tg, 9.8%) during 2015–2017. This contrast may reflect the different emission  
566 control policies during the two clean air action periods, as well as the different emission distributions among the sectors  
567 between SO<sub>2</sub> and CO. According to the estimates of Zheng et al. (2018), the share of emissions from the industrial and power  
568 sector for SO<sub>2</sub> (77%) is nearly double that for CO (39%). Thus, the smaller reduction of CO emissions -than that of SO<sub>2</sub> during  
569 2015–2017 provides evidence that the 2013–2017 action plan mainly focused on controlling the emissions from the industrial  
570 and power sectors. During the 2018–2020 action plan, strict control measures targeted on the residential and transportation  
571 sectors were also implemented, which together account for 61% of CO emissions but only 23% of SO<sub>2</sub> emissions. As a result,  
572 CO showed a larger emission reduction rate during 2018–2020, while the emission reduction rate for SO<sub>2</sub> was similar to that  
573 during 2015–2017. The calculated trends of SO<sub>2</sub> and CO emissions during the two action plans are presented in Table 4, which  
574 are -2.1 Tg/yr and -1.3 Tg/yr for SO<sub>2</sub>, and -22.8 Tg/yr and -33.5 Tg/yr for CO, respectively.

575 The reduction of SO<sub>2</sub> and CO emissions was also evident on the regional scale (Fig. S5 and S6). According to the top-  
576 down estimation, the reduction of SO<sub>2</sub> emissions ranged from 0.44 to 2.42 Tg (41.7–69.9%) from 2015 to 2020, with the NCP  
577 region exhibiting the largest reductions. The calculated decreasing trend of SO<sub>2</sub> emissions was also significant over all regions,  
578 ranging from -0.08 Tg/yr over the NW region to -0.57 Tg/yr over the NCP region (Table 5). With regards to the emission  
579 reduction rate during the different action plans, the results suggest that the emission reduction rate of SO<sub>2</sub> was higher during  
580 2015 – 2017 (by 20.8–39.8%) than that during 2018–2020 (16.6–29.0%) over the NCP, SE, NE and SW regions. This may

581 have been because, after the strict emission controls imposed upon industry and power plants during the 2013–2017 action  
582 plan, the room for further reductions in SO<sub>2</sub> emissions become smaller during the 2018–2020 action plan over these regions.  
583 Although residential and vehicle emissions were controlled more strictly during the 2018–2020 action plan, in total they  
584 account for ~20% of anthropogenic SO<sub>2</sub> emissions in China (Zheng et al., 2018). Thus, the enhanced reductions in SO<sub>2</sub>  
585 emissions from the residential and transportation sectors may not have been able to fully compensate for the weakened  
586 reductions from the industrial and power sectors, leading to a smaller SO<sub>2</sub> emission reduction rate over these regions. In  
587 contrast, the SO<sub>2</sub> emission reduction rate during 2018–2020 (31.1–34.8%) was higher than that during 2015–2017 (14.1–  
588 20.4%) over the NW and Central regions. This may have been due to the fact that the emission controls over the NW and  
589 Central regions were relatively weak during the 2013–2017 action plan (as also evidenced by the emission reduction rates of  
590 other species) owing to its less-developed economy. During the 2018–2020 action plan, the emission controls over these two  
591 regions were strengthened, which led to their higher emission reduction rates. Accordingly, the enhanced SO<sub>2</sub> emission  
592 reduction rates over the NW and Central regions compensated for the weakened reduction rates over the other regions, leading  
593 to a steady SO<sub>2</sub> emission reduction rate on the national scale.

594 The reductions of CO emissions from 2015 to 2020 were approximately 14.9–42.3 Tg (21.6–51.4%) over the different  
595 regions of China, with significant decreasing trends ranging from –3.0 to –8.7 Tg/yr (Fig. S6 and Table 5). Consistent with  
596 the comparisons of national CO emission reduction rates between the two action plans, the emission reduction rates during  
597 2015–2017 (4.4–24.6%) were estimated to be smaller than those during 2018–2020 (12.2–24.6%) over all the different regions  
598 except the Central region, where the CO emission reduction rate was similar during the two action plans (Fig. S6).

#### 599 4.2.2.2 NO<sub>x</sub> and NMVOCs

600 The top-down estimated NO<sub>x</sub> and NMVOC emissions showed different changes to the other four species, by increasing  
601 during 2015–2017 but declining during 2018–2020. Specifically, NO<sub>x</sub> emissions increased slightly by 5.9% from 2015 (25.2  
602 Tg) to 2017 (26.6 Tg), with a non-significant increasing trend of 0.74 Tg/yr. Then, NO<sub>x</sub> emissions began to decrease in 2018,  
603 with a top-down estimated emission reduction and calculated trend of approximately 3.1 Tg (12.7%) and –1.6 Tg/yr,  
604 respectively, from 2018 to 2020. NMVOCs showed stronger emission increases than did NO<sub>x</sub>, with top-down estimated  
605 emission increases of approximately 12.7 Tg (27.6%) and a calculated emission trend of about 6.3 Tg/yr from 2015 to 2017.  
606 Similar to NO<sub>x</sub>, NMVOC emissions began to decrease after 2018, with a top-down estimated reduction of approximately 2.6  
607 Tg (–4.4%) from 2018 to 2020, and a calculated trend of about –1.3 Tg/yr.

608 The increases of NO<sub>x</sub> and NMVOC emissions during 2015–2017 suggest that the 2013–2017 action plan may not have  
609 achieved desirable mitigation effects on these two species. For NO<sub>x</sub> emissions, the upward trend may have been associated  
610 with the following factors. On the one hand, vehicle exhaust is one of the most important sources of NO<sub>x</sub> in China, accounting  
611 for 31% of all NO<sub>x</sub> emissions nationally (Zheng et al., 2018). From 2013 to 2017, the number of vehicles in China continued  
612 to increase and reached 310 million in 2017, approximately 33.5% higher than in 2013 (MEE, 2017), which led to increases  
613 of NO<sub>x</sub> emissions from vehicles in China. On the other hand, although the 2013–2017 action plan was effective in reducing  
614 the NO<sub>x</sub> emissions from coal-fired power plants by promoting denitrification facilities and an ultra-low emission standard, the  
615 mitigation impacts on industrial NO<sub>x</sub> emissions may have been relatively small. For example, Wang et al. (2019a) compiled a  
616 unit-based emissions inventory for China’s iron and steel industry from 2010 to 2015, based on detailed survey results of  
617 approximately 4900 production facilities in mainland China. They found that there were almost no NO<sub>x</sub> control measures in  
618 China’s iron and steel industry during 2010–2015, resulting in a 12.4% increase in China’s NO<sub>x</sub> emissions from the iron and  
619 steel industry in 2015 compared to 2010. In addition, although the penetration rate of denitrification facilities in China’s cement  
620 industry reached 92% in 2015, the actual operating rate of denitrification facilities in the cement industry was not desirable,  
621 due to the lack of online emission monitoring systems. According to the research results of the Ministry of Ecology and  
622 Environment, 800, 1300, and 1400 cement production kilns were equipped with selective non-catalytic denitrification facilities

623 from 2013 to 2015, but the actual operating rates were only 51%, 54% and 73%, respectively (Liu et al., 2021). In addition,  
624 the new precalciner kilns used in the cement industry have a higher NO<sub>x</sub> emission factor, such that the shift from traditional  
625 vertical kilns to precalciner kilns has to some extent increased the cement industry's emissions of NO<sub>x</sub> (Liu et al., 2021). Thus,  
626 there is evidence that the mitigation effects of the industrial control measures on NO<sub>x</sub> emissions may not be as significant as  
627 expected. Overall, the increased number of vehicles may have offset the emission mitigation effects brought about by the  
628 control of power plants, and the mitigation effects of controlling industrial NO<sub>x</sub> emissions were also undesirable. Consequently,  
629 NO<sub>x</sub> emissions in China may not have decreased, and even increased slightly, during the 2013–2017 action plan. Figure S7  
630 further shows the changes in NO<sub>x</sub> emissions over different regions of China, revealing that NO<sub>x</sub> emissions over the NCP, SE,  
631 NE and SW regions were roughly unchanged (by less than 5%) from 2015 to 2017, while they increased over NW (18.6%)  
632 and Central (17.5%). This is consistent with previous results and indicates that NO<sub>x</sub> emissions may have increased over the  
633 NW and Central regions, possibly due to their increased human activities and weak emission controls.

634 In terms of NMVOC emissions, since the inversion results did not differentiate between anthropogenic and biogenic  
635 sources, the changes in NMVOC emissions may have been related to both anthropogenic and biogenic emissions. With respect  
636 to anthropogenic emissions, previous bottom-up studies have suggested that China's NMVOC emissions did not decline during  
637 the 2013–2017 action plan, due to the lack of effective control measures on the chemical industry and solvent use (Zheng et  
638 al., 2018; Li et al., 2019c). According to the estimates of Li et al. (2019c), China's NMVOC emissions from solvent use  
639 increased by 11.1% in 2017 compared to those in 2015. Meanwhile, the increase in the number of vehicles in China may also  
640 have led to an increase in NMVOC emissions from transportation. Thus, the increases of NMVOC emission during 2015–  
641 2017 estimated by our inversion inventory may be related to the increases in anthropogenic NMVOC emissions from the  
642 chemical industry, solvent use, and vehicles. For the trends of biogenic NMVOC emissions, the CAMS global emission  
643 inventory shows that there were only little changes in the biogenic NMVOC emissions in China from 2013 to 2018 (Sect.  
644 4.3.3), suggesting little contributions of the biogenic sources to the increased NMVOC emission in China. Figure S8 further  
645 shows the changes in NMVOC emissions over different regions of China, which suggests consistent increases in NMVOC  
646 emissions from 2015 to 2017 over different regions. According to the top-down estimations, NMVOC emissions increased by  
647 30.5%, 25.2%, 18.5%, 10.9%, 50.5% and 63.1% over the NCP, SE, NE, SW, NW and Central regions, respectively. Again,  
648 the NW and Central regions exhibited the largest emission increases among the six regions, which is consistent with their  
649 elevated levels of human activity and weak emission controls.

650 The decrease in NO<sub>x</sub> and NMVOC emissions after 2018 suggests that the emission control strategy of the Chinese  
651 government had reached a point of optimization. The 2018–2020 action plan not only strengthened the controls over the  
652 industrial and power sectors, but also the transportation sector, especially for diesel vehicles with high NO<sub>x</sub> emissions. For  
653 example, the Chinese government released the “Action Plan for the Control of Diesel Trucks”, and vigorously promoted an  
654 adjustment of the transportation structure of China by gradually improving the availability of rail transport. As a result, there  
655 was a downward trend in NO<sub>x</sub> emissions in China. The top-down estimated reductions of NO<sub>x</sub> emissions were approximately  
656 0.81 Tg (17.2%) over NCP, 0.98 Tg (14.0%) over SE, 0.37 Tg (9.4%) over NE, 0.51 Tg (12.2%) over SW, 0.13 Tg (11.0%)  
657 over NW, and 0.32 Tg (9.2%) over Central (Fig. S7). The decrease in NMVOC emissions after 2018 may on the one hand  
658 have been related to the strengthening of vehicle controls during the 2018–2020 action plan, whilst on the other hand it may  
659 have been related to the promotion of clean heating plans in the northern region, which reduced the emissions of NMVOCs  
660 from residential sources. However, the decreases in NMVOC emissions were smaller than those in NO<sub>x</sub>, which were estimated  
661 to be 0.84 Tg (6.9%) over NCP, 0.47 Tg (2.8%) over SE, 0.98 Tg (10.1%) over NE, and 0.53 Tg (14.1%) over NW (Fig. S6).  
662 Different from other regions, the NMVOC emissions over the SW and Central regions remained almost unchanged during the  
663 2018–2020 action plan (Fig. S8).



### 664 4.2.3 Changes in the distribution pattern of emissions in China

665 Due to the different emission control intensities over the different regions of China, the emission distribution patterns of  
666 the different species may also have been altered, which could have influenced the distributions of air pollution in China. Based  
667 on CAQIEI, we further investigated the emission distribution patterns, as well as their changes, during the two action plans.  
668 Maps of the emission changes of different species during 2015–2017 and 2018–2020 are presented in Fig. 8. The shares of  
669 emissions in 2015, 2017 and 2020 by each subregion of China are also presented (Fig. 9). It can be seen that the emission  
670 changes during the 2015–2017 were more heterogeneous than those during 2018–2020. The air pollutant emissions after the  
671 2018–2020 action plan showed consistent reductions over most regions of China, while there were obvious emission increases  
672 detected from 2015 to 2017. This is consistent with the different emission control effects during the two clean air action plans  
673 as mentioned in previous sections. Due to its strictest emission control policies, the NCP region showed consistent emission  
674 reductions of SO<sub>2</sub>, NO<sub>x</sub>, CO, PM<sub>2.5</sub> and PM<sub>10</sub> during the two clean air action plans. Accordingly, the shares of emissions in the  
675 NCP region continued to decrease during the two action plan periods (Fig. 9). For example, the share of SO<sub>2</sub> emissions in the  
676 NCP region decreased from 19.4% to 15.4% during the period of 2015–2017, and from 15.4% to 12.7% during the 2018–2020  
677 action plan. In contrast, NMVOC emissions increased obviously over the NCP region from 2015 to 2017, and decreased during  
678 2018–2020. However, its share did not change significantly, being roughly 20% throughout both periods. As for other regions,  
679 increases of SO<sub>2</sub>, NO<sub>x</sub>, PM<sub>2.5</sub>, PM<sub>10</sub> and NMVOC emissions during 2015–2017 could be found over the Central region. More  
680 specifically, the emission increases were mainly located in the Fenwei Plain area of the Central region, which was due to the  
681 fact that this area was not included as a key region of emission controls during the 2013–2017 action plan. However, the  
682 Fenwei Plain area was added as a key emission control region during the 2018–2020 action plan, which is consistent with the  
683 emission reductions for these species over the Central region (Fig. 8). As a result, the shares of SO<sub>2</sub> and PM<sub>2.5</sub> emissions in the  
684 Central region increased during 2015–2017 but decreased during 2018–2020 (Fig. 9). However, the shares of NO<sub>x</sub>, PM<sub>10</sub> and  
685 NMVOC emissions continued to increase over the Central region during the two clean air action plans, which suggests larger  
686 roles of air pollutant emissions in that region. In contrast, the share of CO emissions in the Central region continued to decrease  
687 in the two action plans, from 17.7% in 2015 to 13.4% in 2020.

688 In terms of the shares of emissions in eastern and western China, the top-down estimation suggests an increased share of  
689 NO<sub>x</sub>, PM<sub>2.5</sub>, PM<sub>10</sub> and NMVOC emissions in western China after the two clean air action plans (Fig. 9), which indicates slower  
690 emission reductions for these species in western China. However, the share of CO emissions in western China was reduced  
691 after the two clean air action plans. Although the share of SO<sub>2</sub> emissions in western China increased during 2015–2017, it  
692 turned to a decrease during 2018–2020.

### 693 4.3 Comparisons with different emission inventories

694 In this section, the CAQIEI is compared with the previous long-term bottom-up and top-down emission inventories in  
695 China to validate our inversion results and provide the clues for the potential uncertainty in the current air pollutant emission  
696 inventories. The bottom-up emission inventories used in the comparison include MEIC (Zheng et al., 2018), ABaCAS (Li et  
697 al., 2023), HTAPv3 (Crippa et al., 2023), EDGARv6 (Jalkanen et al., 2012) and CEDS (McDuffie et al., 2020), while the top-  
698 down emission inventory is obtained from the updated Tropospheric Chemistry Reanalysis (TCR-2) (Miyazaki et al., 2020b).  
699 However, it is difficult to directly compare our inversion results with these emission inventories considering that the inversion  
700 emission includes both anthropogenic and natural emissions. To better compare our inversion results with previous inventories,  
701 the natural emission sources, including soil NO<sub>x</sub> emissions and biogenic emissions obtained from the CAMS global emission  
702 inventory (<https://ads.atmosphere.copernicus.eu/cdsapp#!/dataset/cams-global-emission-inventories?tab=overview>; last  
703 accessed 26 July 2023) and the biomass burning emissions obtained from the Global Fire Assimilation System (GFAS) (Kaiser  
704 et al., 2012) are taken as a reference to account for the influences of natural sources. The CAMS and GFAS emission inventory

705 are used because they are state-of-art natural emission inventories and can provide us with [independent](#) long-term estimations  
706 of natural emissions. Since the latest year of most emission inventories is 2018, the comparisons were conducted between 2015  
707 and 2018. Note that due to the complexity in the estimations of natural sources, significant uncertainty exists in the estimated  
708 natural emissions. As a result, the comparison results would be sensitive to the used natural emission inventories, especially  
709 for the species with large amount of natural emission, such as the NMVOC and particulate matter. Therefore, it should be  
710 aware of that the comparison conducted here and the derived implications are on the basis of the natural emissions estimated  
711 by CAMS and GFAS. In addition, the natural dust emissions are not considered in the comparisons, which would influence  
712 the comparisons of the PM emissions.

### 713 4.3.1 Magnitude

#### 714 4.3.1.1 NO<sub>x</sub>

715 Figure 10 shows the average emissions of different air pollutants in China during 2015–2018 obtained from CAQIEI and  
716 the previous emission inventories plus the natural sources we considered. Comparisons of the emission estimations on the  
717 regional scale and gridded scale are also presented (Fig. 11 and Fig. S9). The results show that the CAQIEI has slightly higher  
718 NO<sub>x</sub> emissions in China than the other inventories. Considering that CAQIEI includes both anthropogenic and natural sources,  
719 this discrepancy could be explained by the natural NO<sub>x</sub> sources. According to the estimations of CAMS and GFAS, the soil  
720 and biomass-burning NO<sub>x</sub> emissions are approximately 1.9 and 0.08 Tg/yr, which explains well the higher NO<sub>x</sub> emissions  
721 given by CAQIEI. After consideration of the natural sources, MEIC, HTAPv3 and EDGARv6 agree well with our inversion  
722 results on the national scale, with their differences within 1.0–7.4%. The NO<sub>x</sub> emission estimated by ABaCAS, CEDS and  
723 TCR-2 are slightly lower than CAQIEI and other emission inventories. However, the differences between CAQIEI and these  
724 inventories were found to range from 15.9% to 21.3%, which is within the previous estimated uncertainties of NO<sub>x</sub> emissions  
725 in China (Kurokawa and Ohara, 2020; Li et al., 2017b; Li et al., 2023). These results suggest that the total NO<sub>x</sub> emissions in  
726 CAQIEI are generally consistent with the current estimations of the anthropogenic and natural NO<sub>x</sub> emissions in China. On  
727 the regional scale, the top-down estimated NO<sub>x</sub> emissions show good agreement with the previous emission inventories over  
728 the NCP and SE regions, with their differences ranging from 1.0%–26.8%, suggesting good consistency in the estimations of  
729 NO<sub>x</sub> emissions over these two regions. This makes sense because NCP and SE are the two most developed regions in China,  
730 and where surveys and research on emissions are most sufficient. The differences are larger over the other regions. In the NE  
731 region, CAQIEI has higher NO<sub>x</sub> emissions than the other inventories by 5–70%, suggesting higher anthropogenic or biomass-  
732 burning emissions over there. The estimations made by MEIC, CEDS and TRC-2 are closer to our estimates, with their  
733 differences being approximately 5.4–23.3%, while the differences are larger for ABaCAS, HTAPv3 and EDGARv6 (36.7–  
734 70.0%). Over the SW and Central regions, there are large diversity in the previous emission inventories with estimations by  
735 HTAPv3 and EDGARv6 almost double those of MEIC, ABaCAS, CEDS and TCR-2. The CAQIEI suggests a midst estimation  
736 which is within the range of previous emission inventories. In the NW region, CAQIEI is consistently higher than other  
737 inventories, by 22.7–64.2%, which suggests a potential missing source of the NO<sub>x</sub> emissions over this region.

#### 738 4.3.1.2 SO<sub>2</sub>

739 For SO<sub>2</sub> emissions, since natural sources contribute little (only about 0.02 Tg/yr) to them in China, the discrepancies  
740 between CAQIEI and previous emission inventories are mainly attributable to the differences in anthropogenic emissions. As  
741 shown in Fig. 10, CAQIEI agrees well with HTAPv3 and CEDS on the national scale, with their differences being  
742 approximately ±2%, but is higher than MEIC, ABaCAS and TCR-2 by 17.4–32.9%. In contrast, EDGARv6 may have a  
743 positive bias in its estimated SO<sub>2</sub> emissions, which are roughly double those of CAQIEI and other inventories. On the regional  
744 scale, our results agree well with MEIC, ABaCAS, HTAPv3, CEDS and TCR-2 over the NCP region, with their differences  
745 ranging from 1.0 to 18.1%. In the SE region, CAQIEI suggest lower SO<sub>2</sub> emissions than previous emission inventories, except

746 TCR-2. The differences are relatively smaller for the MEIC and ABaCAS inventories by around -15%, but larger for HTAPv3,  
747 EDGARv6 and CEDS (ranging from -47.3% to -113.2%). In contrast, CAQIEI suggests higher SO<sub>2</sub> emissions than all  
748 previous emission inventories over the NE region by about 14.8–132.0%, indicating a possible missing sources over there.  
749 Similarly, the CAQIEI and HTAPv3 suggests higher SO<sub>2</sub> emissions than the MEIC, ABaCAS, CEDS and TCR-2 by 27.0–  
750 75.6% in the NW region, and by 44.3–77.7% in the Central region.

### 751 4.3.1.3 CO

752 For CO emissions, CAQIEI is substantially higher than the previous emission inventories, with the estimated CO  
753 emissions of CAQIEI being about three times higher than the bottom-up inventories and more than double those of the top-  
754 down estimates made by TCR-2. According to GFAS, the average rate of CO biomass-burning emissions in China from 2015  
755 to 2018 was about 3.4 Tg/yr. Yin et al. (2019), based on MODIS fire radiative energy data, also estimated China's CO biomass-  
756 burning emissions to be about 5.0 (2.3–7.8) Tg/yr. The biogenic CO emissions obtained from the CAMS global emission  
757 inventory were approximately 2.3 Tg/yr. According to these estimates, natural CO emissions in China have a magnitude of  
758 about 10<sup>1</sup>, which is rather small compared with anthropogenic sources, and cannot explain the large discrepancies between  
759 CAQIEI and other inventories. Thus, the CAQIEI suggest much higher anthropogenic CO emissions in China than the existing  
760 emission inventories. In fact, the potential underestimation of CO anthropogenic emissions has been investigated in previous  
761 studies and is regarded as the main reason for the negative bias in global or hemispheric CO simulations (Stein et al., 2014;  
762 Gaubert et al., 2020). Regionally, Kong et al. (2020) compared a suite of 13 modeling results from six different CTMs—  
763 namely, NAQPMS, CMAQ, WRF-Chem, NU-WRF, NHM-Chem and GEOS-Chem—with observations over the NCP and  
764 Pearl River Delta regions under the framework of the Model Inter-Comparison Study for Asia III (MICS-Asia III), and found  
765 consistent negative biases in the CO simulations of all models, pointing toward potential underestimations of CO emissions in  
766 China. Previous inversion studies have also reported higher a posteriori CO emissions than their *a priori* emission inventories  
767 (Bergamaschi et al., 2000; Miyazaki et al., 2012; Petron et al., 2002; Petron et al., 2004; Tang et al., 2013; Gaubert et al., 2020).  
768 For example, the constrained CO emissions reported by Gaubert et al. (2020) are 80% higher than the CEDS over the northern  
769 China. Our inversion results are consistent with these inversion studies, suggesting higher anthropogenic CO emissions in  
770 China. However, direct evidence in support of such high CO emissions in China reported by our study is still limited currently.  
771 Thus, we compiled more inversion results within the period of 2013–2020 from previous studies to further validate our  
772 inversion results, which are summarized in Table 6. It can be clearly seen that there are large differences in the estimated CO  
773 emissions between the inversion results based on surface observations and those based on satellite data. Our inversion results  
774 are consistent with the results of Feng et al. (2020), with China's CO emissions in December 2017 estimated at approximately  
775 1500.0 kt/day and 1388.1 kt/day, respectively. In addition, Feng et al. (2020) used the CMAQ model to constrain CO emissions,  
776 which is different from the model we used. This may indicate that the model uncertainty would not significantly influence the  
777 inversion results of CO emissions. However, the top-down estimated CO emissions based on satellite data (163.6–553.4 kt/day)  
778 are much lower than those based on surface observations, although they are all higher than their *a priori* emissions. The lower  
779 CO emission estimations based on satellite data assimilation may be attributable to the lower sensitivities of satellite data to  
780 surface concentrations, suggesting that the assimilation of satellite data alone may not be adequate to correct the negative  
781 biases in the *a priori* emissions. This deficiency has also been revealed by Miyazaki et al. (2020b), who found undercorrected  
782 surface CO emissions in the extratropics of the Northern Hemisphere in TCR-2. However, the assimilation of surface  
783 observations can be influenced by the uncertainties in the modeled vertical mixing, which could lead to the uncertainties in the  
784 inverted CO emissions based on surface observations. Therefore, the inverted CO emissions in CAQIEI could be partly  
785 supported by previous inversion studies based on surface observations, but more evidence is still needed to justify the  
786 magnitude of the inverted CO emissions. Besides anthropogenic sources, the chemical production of CO via oxidation of  
787 methane (CH<sub>4</sub>) and NMVOCs, as well as the CO sinks via the hydroxyl radical (OH) reaction, also influence the simulation

788 of CO (Stein et al., 2014; Gaubert et al., 2020; Müller et al., 2018). Due to the important role of OH in the chemical production  
789 and sinks of CO, the inversion of CO emissions is sensitive to the modeled OH abundance and the emissions of CH<sub>4</sub> and  
790 NMVOCs. According to the estimation of Müller et al. (2018), the magnitude of inversed CO emissions in China could differ  
791 by more than 40% when different levels of OH concentrations are used in the model. Thus, the much higher estimations of  
792 CO emissions in our inversion results may also be partly explained by the underestimation of CO chemical production or the  
793 overestimation of the CO sink.

#### 794 4.3.1.4 PM<sub>2.5</sub>

795 In terms of PM<sub>2.5</sub>, the CAQIEI suggests higher emissions than ABaCAS, HTAPv3 and EDGARv6 by about 20%, and by  
796 47.7% than MEIC on the national scale. Larger discrepancies mainly occur in the NE and NW regions, where CAQIEI is about  
797 27.2–114.9% and 83.2–143.2% higher than the previous inventories. The differences in the estimated PM<sub>2.5</sub> emissions may be  
798 related to the uncertainties in the biomass-burning or anthropogenic sources in the NE region (Wu et al., 2020b), while in the  
799 NW region, the errors in the fine-dust emissions may also contribute to the differences in the estimated PM<sub>2.5</sub> emissions there.  
800 The differences in the estimated PM<sub>2.5</sub> emissions are relatively smaller in the NCP and SE regions, ranging from –18.9% to  
801 20.4%, suggesting better agreement in the estimated PM<sub>2.5</sub> emissions over these two regions. In the SW region, CAQIEI is  
802 closer to HTAPv3 and EDGARv6, with their differences being about 6.3% and –9.5% respectively, and is higher than MEIC  
803 and ABaCAS by 54.2% and 28.6%, suggesting higher uncertainty in the estimated PM<sub>2.5</sub> emissions over there.

#### 804 4.3.1.5 PM<sub>10</sub>

805 For PM<sub>10</sub> emissions, it is difficult to directly compare CAQIEI with previous emission inventories since CAQIEI not only  
806 contains anthropogenic and biomass-burning emissions, but also coarse-dust emissions. As a result, the estimated emissions  
807 of PM<sub>10</sub> by CAQIEI are substantially higher than those by previous inventories, especially over the NW, Central and NE  
808 regions (Fig. 11), which are the typical natural windblown dust-source regions in China (Zeng et al., 2020). Besides the  
809 naturally windblown dust of arid desert regions (Prospero et al., 2002), large amounts of coarse-dust emissions also stem from  
810 anthropogenic sources, including anthropogenic fugitive, combustion and industrial dust from urban sources (AFCID) (Philip  
811 et al., 2017), and anthropogenic windblown dust from human-disturbed soils due to changes in land-use practices, deforestation  
812 and agriculture (Tegen et al., 1996). Therefore, although the other regions are not typical natural windblown dust-source  
813 regions in China, there are still high levels of coarse-dust emissions from anthropogenic sources there (also called “urban  
814 dust”), which may be the main reason for the large deviation in the estimated PM<sub>10</sub> emissions between CAQIEI and previous  
815 inventories. On the one hand, although AFCID is included in MEIC, ABaCAS, HTAPv3 and EDGARv6, it is difficult for  
816 current bottom-up emission inventories to completely represent fugitive sources (Philip et al., 2017). On the other hand, the  
817 anthropogenic windblown dust emissions haves not been included in current bottom-up emission inventories, which is an  
818 important source of coarse dust in urban areas according to the estimations of Li et al. (2016) and the another important  
819 contributor to the differences between CAQIEI and previous emission inventories.

#### 820 4.3.1.6 NMVOCs

821 For NMVOC emissions, since CAQIEI includes both anthropogenic and natural sources, its estimated NMVOC emissions  
822 are much higher than those estimated by previous emission inventories. After consideration of natural sources, the CAQIEI  
823 suggests close estimations of the NMVOC emissions with the MEIC, HTAPv3 and CEDS inventories on the national scale,  
824 with their differences being about 1.5–12.5%. The estimated NMVOC emission by ABaCAS and EDGARv6 are slightly lower  
825 than CAQIEI by 17.8% and 24.6%, respectively. On the regional scale, the CAQIEI suggests higher NMVOC emissions over  
826 the northern China (NCP, NE and NW), with the top-down estimated NMVOC emissions about 30.4–81.4%, 27.3–72.1%,  
827 79.3–116.8%, and 8.7–57.5% higher than those of the previous emission inventories. In contrast, the CAQIEI suggests lower

828 NMVOC emissions over the SE region, with the estimated NMVOC emissions of CAQIEI being about 21.2–27.6% lower  
829 than those of MEIC, ABaCAS, HTAPv3 and CEDS. These results are consistent with the previous inversion results based on  
830 the satellite observations, which suggest higher NMVOC emissions over the NCP region and lower NMVOC emissions over  
831 the south China (Souri et al., 2020). Over the SW region, CAQIEI shows good agreement with MEIC, ABaCAS and CEDS,  
832 with CAQIEI being slightly lower than these inventories by 1.0–8.9%, but is lower than HTAPv3 and EDGARv6 by about  
833 38.6% and 29.1%, respectively. Again, it should be noted that the comparisons of NMVOC emission are conducted on the  
834 basis of natural emissions estimated by CAMS and GFAS, and could be more sensitive to the used natural sources than other  
835 species considering the larger contributions of the natural source to the NMVOC emissions.

### 836 4.3.2 Seasonality

837 Figure 12 presents the monthly profiles of different air pollutants obtained from different emission inventories. Note that  
838 the natural sources have been added to the previous inventories to facilitate the comparisons. The results show that different  
839 emission inventories give similar monthly profiles of NO<sub>x</sub> and CO emissions, with higher emissions during wintertime and  
840 lower emissions during summertime, which suggests relatively lower uncertainty in the estimated monthly profiles for these  
841 two species. For SO<sub>2</sub> emissions, CAQIEI yields stronger monthly variation than the other inventories, with a higher proportion  
842 from January to March and lower proportion during summertime. Due to the influences of dust emissions, the top-down  
843 estimated PM<sub>2.5</sub> and PM<sub>10</sub> emissions show higher proportions than the other emission inventories during the spring season,  
844 especially for PM<sub>10</sub>. However, the proportion of emissions during autumn and winter are lower than in the other inventories.  
845 The monthly profiles of NMVOC emissions are generally consistent, with higher emissions during summer due to the enhanced  
846 biogenic emissions. However, the profile of CAQIEI is flatter than the previous inventories, and suggests a higher proportion  
847 during springtime. In addition, the timings of peak values of NMVOC emissions are also different between CAQIEI and the  
848 previous inventories, with CAQIEI showing peak values during May–July but the other inventories suggesting peaks during  
849 June–August.

### 850 4.3.3 Emission changes during 2015–2018

851 The top-down estimated emission changes of different air pollutants during 2015–2018 were also compared with previous  
852 emission inventories. Figure 13 shows the time series of the total emissions of different species from 2013 to 2020 obtained  
853 from the CAQIEI and other emission inventories. Comparisons of the emission changes over the regional scales are also  
854 presented in Fig. S10–S15. Before the comparison, we firstly analyze the trends of natural sources in China to investigate their  
855 influences on the emission changes of different species based on the CAMS emission inventory and GFAS. Note that we only  
856 consider the soil, biogenic and biomass-burning emissions for the natural sources; the trends of dust emissions in China are  
857 not analyzed, which may lead to uncertainty when comparing the emission changes of PM<sub>2.5</sub> and PM<sub>10</sub>. As shown in Fig. S16,  
858 the natural sources of NO<sub>x</sub> and NMVOC emissions changed little during 2013–2018. The other species had small decreasing  
859 trends from 2013 to 2018. However, considering the small contributions of natural sources to their emissions, these small  
860 trends would not significantly influence their emission trends. For the dust emissions, previous studies have indicated a  
861 declining trend in dust activity in China from 2001 to 2020 (Wu et al., 2022; Wang et al., 2021), due to weakened surface wind  
862 and increased vegetation cover and soil moisture. These results suggest that the emission trends in the CAQIEI would be  
863 mainly driven by the anthropogenic sources for the gaseous air pollutants based on the estimations of CAMS and GFAS, while  
864 its estimated emission trends of PM<sub>2.5</sub> and PM<sub>10</sub> would be influenced by the declining trends in dust emissions in China, which  
865 should be noted when comparing the emission changes of PM<sub>2.5</sub> and PM<sub>10</sub>.

866 As shown in Fig. 14, all the emission inventories agree that the NO<sub>x</sub>, SO<sub>2</sub>, CO, PM<sub>2.5</sub> and PM<sub>10</sub> emissions in China were  
867 reduced from 2015 to 2018, except for the increases of CO emissions estimated by TCR-2, which confirms the effectiveness  
868 of the emission control policies implemented during the clean air action plans. Meanwhile, most emission inventories agree

869 that SO<sub>2</sub> is the species with the largest emission reduction rate, followed by PM<sub>2.5</sub>, indicating better emission mitigation effects  
870 of these two species (Fig. 14). However, the CAQIEI suggested lower emission reduction rates than the other emission  
871 inventories for most species, especially for NO<sub>x</sub>, PM<sub>10</sub> and NMVOCs (Fig. 14). The estimated emission reduction rate of NO<sub>x</sub>  
872 obtained from CAQIEI is about -2.7%, which is lower than the values of MEIC (-9.7%), ABaCAS (-23.0%), HTAPv3  
873 (-13.0%) and CEDS (-9.0%). As we discussed in Sect. 4.2.2.2, the small reductions of NO<sub>x</sub> emission in CAQIEI would be  
874 related to the increased vehicle emissions and the undesirable mitigation effects of the industry control. In fact, these factors  
875 have been considered in some bottom-up emission inventories, such as MEIC. The differences between our inversion results  
876 and previous inventories thus reflect uncertainty in the quantifications of the effects of these factors on the NO<sub>x</sub> emissions due  
877 to the lack of sufficient statistics on mobile vehicle or other sectors. Our inversion results suggest larger adverse effects of  
878 these two factors on the reductions of NO<sub>x</sub> emissions in China. According to Fig. S17, the differences between CAQIEI and  
879 these inventories mainly occur in the SE, SW, NW and Central regions, with the emission reduction rate estimated by CAQIEI  
880 being substantially lower than those estimated by previous inventories. In particular, CAQIEI suggests increases of NO<sub>x</sub>  
881 emissions over the Central region, which is opposite to the previous emission inventories. Better agreement is achieved over  
882 the NCP and NE regions, with the emission reduction rate estimated by CAQIEI being closer to those of MEIC, HTAPv3 and  
883 CEDS. The NO<sub>x</sub> emission reduction rates estimated by EDGARv6 (-3.3%) and TCR-2 (-1.7%) are closer to our results on  
884 the national scale, but they estimated lower NO<sub>x</sub> emission reduction rate than our estimate over the NCP and NE regions.

885 Similarly, the emission reduction rate of PM<sub>10</sub> obtained from CAQIEI (-10.8%) is lower than those estimated by MEIC  
886 (-27.9%), ABaCAS (-33.0%) and HTAPv3 (-27.8%) on the national scale (Fig. 14). A lower PM<sub>10</sub> emission reduction rate  
887 of CAQIEI than these inventories also exist in the different regions of China, except SW (Fig. S17). In particular, different  
888 from previous emission inventories, CAQIEI suggests that PM<sub>10</sub> emissions may have actually increased over the Central region.  
889 Considering that dust emissions may have decreased from 2015 to 2018 owing to weakened dust events (Wang et al., 2021),  
890 the increase in PM<sub>10</sub> emissions over the Central region may reflect the increases in anthropogenic sources. Meanwhile, we also  
891 found that CAQIEI estimated the emission reduction rate of PM<sub>10</sub> to be smaller than that of PM<sub>2.5</sub>. This is different from  
892 previous emission inventories, which show similar emission reduction rates for PM<sub>2.5</sub> and PM<sub>10</sub>. Considering that PM<sub>10</sub>  
893 emissions include PM<sub>2.5</sub> and PM<sub>10-2.5</sub> emissions, the lower emission reduction rate of PM<sub>10</sub> than PM<sub>2.5</sub> in CAQIEI suggests that  
894 PM<sub>10-2.5</sub> emissions may have decreased slower than PM<sub>2.5</sub> emissions from 2015 to 2018.

895 In terms of NMVOCs, most previous inventories, including MEIC, EDGARv6 and CEDS, suggest a weak decrease in  
896 China, with the estimated rates of change in emissions ranging from -0.8% to -4.6%. The emission reduction rate estimated  
897 by ABaCAS is larger, reaching up to -14.2%. In contrast, the CAQIEI suggests an opposite emission change to these  
898 inventories, with estimated NMVOC emissions increasing by 26.6% from 2015 to 2018. HTAPv3 also suggests an increase in  
899 NMVOC emissions, but with a much lower rate of increase (2.7%). Similar results could also be found on the regional scale  
900 (Fig. S17), especially over the NCP, NE and Central regions, where NMVOC emissions could have increased by 38.0%, 38.3%  
901 and 60.0%, respectively, according to the estimates of CAQIEI. As we discussed in Sect. 4.2.2.2, the increases of NMVOC  
902 emission estimated in CAQIEI may be related to the increased anthropogenic NMVOC emissions from the chemical industry,  
903 solvent use, and vehicles. Therefore, similar to the NO<sub>x</sub> emissions, the differences between CAQIEI and previous inventories  
904 reflects the uncertainty in the quantifications of the impacts of these factors, and suggest larger adverse effects of these factors  
905 on the emission reductions of NMVOC emission than the previous inventories.

906 The differences in the estimated emission reduction rates between CAQIEI and previous inventories are relatively smaller  
907 for SO<sub>2</sub> and PM<sub>2.5</sub> emissions. The emission reduction rate of SO<sub>2</sub> estimated by CAQIEI is close to that estimated by MEIC and  
908 CEDS, ranging from -34.7% to -44.3%. ABaCAS and HTAPv3 estimate a larger emission reduction rate of about -58.5%  
909 and -53.7%, respectively. EDGARv6 and TCR-2 may underestimate the reduction rate of SO<sub>2</sub>, with estimates of only about  
910 -7.0% and -9.1%, respectively. This may be because EDGARv6 underestimates the FGD (flue-gas desulfurization devices)  
911 penetration or SO<sub>2</sub> removal efficiencies of FGD in China. On the regional scale (Fig. S17), the top-down estimated SO<sub>2</sub>

912 emission reduction rate agrees reasonably with that of MEIC over the NCP, NE and SE regions, but these inventories estimate  
913 different SO<sub>2</sub> emission reduction rates over the SW, NW, and Central regions. The reduction rates estimated by MEIC over  
914 the SW and Central regions is higher than those given by CAQIEI, but lower over the NW region. The other emission  
915 inventories also give different emission reduction rates, suggesting large uncertainty in the estimated SO<sub>2</sub> emission reduction  
916 rates over these three regions. In terms of PM<sub>2.5</sub>, CAQIEI's estimated emission reduction rate agrees well with those of MEIC  
917 and HTAPv3 on the national scale, which is about 24–27% from 2015 to 2018. The emission reduction rate of PM<sub>2.5</sub> estimated  
918 by EDGARv6 are lower than our estimates and other inventories, which were about 9%. On the regional scale, our results  
919 show good consistency with MEIC and HTAPv3 over the NCP, NE, SE and SW regions, but they have large differences over  
920 the NW and SW regions.

921 Different from the other species, the CO emission reduction rate estimated by CAQIEI (–21.3%) is higher than in most  
922 of the previous inventories, including MEIC (–13.0%), ABaCAS (–11.6%), EDGARv6 (–4.7%), and CEDS (–11.7%),  
923 suggesting larger mitigation effects on CO emissions than other inventories. HTAPv3 agrees with our results, with an estimated  
924 emission reduction rate of about –22.0%. On the regional scale (Fig. S17), our result is consistent with MEIC over the NCP  
925 and SE regions, with estimated emission reduction rates for CO of around 24% and 15%, respectively, while in other regions  
926 the emission reduction rate estimated by CAQIEI is higher than that estimated by MEIC. The TCR-2 shows opposite changes  
927 in CO emissions compared with the other inventories insofar as it suggests increases of CO emissions over different regions  
928 of China. Since the emissions in TCR-2 are constrained by satellite observations, the differences between our results and those  
929 of TCR-2 highlight that the observations used to constrain the emissions may have a large influence on the estimated emission  
930 changes. In this case, the estimated changes of CO emissions by CAQIEI are more consistent with those estimated by other  
931 bottom-up inventories. Considering this, the TCR-2 may have uncertainties in its estimated changes of CO emissions in China  
932 from 2015 to 2017, which could be related the suboptimal performance of the data assimilation caused by the underestimated  
933 background errors of CO or too short assimilation window for the CO emission estimates (Miyazaki et al., 2020)~~In this case,~~  
934 ~~the assimilation of surface observations (our study) is shown to be superior to the assimilation of satellite observations (TCR-~~  
935 ~~2), as our results are more consistent with other bottom up inventories.~~

#### 936 4.4 Uncertainty estimation of CAQIEI

937 Finally, the uncertainty of the inversed emission inventory product is estimated in this section to facilitate users'  
938 understanding of the data's accuracy. Within the framework of EnKF, the analysis perturbation  $\mathbf{X}^a$  estimated by using Eq. (3)  
939 could provide the information regarding the uncertainty of the inversed emission inventory. The Coefficient of variation  
940 (hereinafter, CV), defined as the standard deviation divided by the average, with a larger value denoting higher uncertainty, is  
941 calculated based on the analysis perturbation to measure the uncertainty of the inverse emission inventory. Based on this  
942 method, the uncertainty (CV) of the a posteriori emission was estimated as follows: 92.3% (PM<sub>2.5</sub>), 88.8% (PM<sub>10</sub>), 26.7%  
943 (SO<sub>2</sub>), 46.8% (CO), 31.8% (NO<sub>x</sub>) and 65.5% (NMVOC). However, it should be noted that such uncertainty was only calculated  
944 under the framework of the EnKF constructed in this study, which is dependent on the assigned value of the a priori emission  
945 uncertainty, observation errors and the number of assimilated observations. In addition, we only considered the a priori  
946 emission uncertainty and the observation errors during the inversion. The influences of the other error sources, such as  
947 uncertainty in the chemistry transport model, meteorology simulations and the inversion method were not considered.  
948 Therefore, the current estimated uncertainty should be considered as a lower bound for the real uncertainty. More systematic  
949 analysis that thoroughly consider the uncertainty sources regarding the emission inversion should be conducted in future to  
950 give a more accurate estimation of the uncertainty in our products.

952 A long-term, top-down emissions inventory of major air pollutants in China was developed and validated in this study by  
953 assimilating surface observations from CNEMC using the modified EnKF method and NAQPMS. It includes gridded emission  
954 maps of NO<sub>x</sub>, SO<sub>2</sub>, CO, primary PM<sub>2.5</sub>, primary PM<sub>10</sub>, and NMVOCs in China from 2013 to 2020, on a monthly basis, with a  
955 horizontal resolution of 15 km × 15 km. This new top-down emissions inventory, named CAQIEI, provides new insights into  
956 the air pollutant emissions and their changes in China during the country's two clean air action periods. The estimated total  
957 emissions for the year 2015 in China are 25.2 Tg of NO<sub>x</sub>, 17.8 Tg of SO<sub>2</sub>, 465.4 Tg of CO, 15.0 Tg of PM<sub>2.5</sub>, 40.1 Tg of PM<sub>10</sub>  
958 and 46.0 Tg of NMVOCs. Comparisons of CAQIEI with previous inventories, including MEIC, ABaCAS, HTAPv3,  
959 EDGARv6, CEDS and TCR-2, on the basis of the natural emissions obtained from CAMS and GFAS showed reasonable  
960 agreement for the estimation of NO<sub>x</sub>, SO<sub>2</sub> and NMVOC emissions in China. The PM<sub>2.5</sub> emissions obtained from CAQIEI (13.2  
961 Tg) are slightly higher than in the previous emission inventories (8.3–11.1 Tg), while the CO emissions estimated by CAQIEI  
962 (426.8 Tg) are substantially higher than in previous inventories (120.7–237.7 Tg). However, the reasons for such a large gap  
963 are still not clear, but might be attributable to both the underestimation of CO sources (e.g., anthropogenic, biomass-burning  
964 and chemical-production sources) ~~according to previous model simulation and inversion studies~~ (Bergamaschi et al., 2000;  
965 Miyazaki et al., 2012; Petron et al., 2002; Petron et al., 2004; Tang et al., 2013; Gaubert et al., 2020), and/or the overestimation  
966 of CO sinks in the model (Müller et al., 2018). In addition, comparisons with previous inversion studies suggest there are larger  
967 differences in the top-down estimated CO emissions based on surface and satellite observations. Our inversion results are  
968 consistent with previous inversions based on surface observations, but are much higher than those based on satellite  
969 observations, suggesting large uncertainty in inversion-estimated CO emissions in China. Therefore, more research is needed  
970 to better understand the reasons behind the negative biases in CO simulation, and to explain the differences between our results  
971 and those of previous inventories. Similar to situation with CO emissions, the PM<sub>10</sub> emissions estimated by CAQIEI (37.7 Tg)  
972 are also substantially higher than in previous inventories (11.1–15.9 Tg). However, this will be mainly associated with the  
973 emissions of coarse dust, which were not included in previous inventories. The estimation of dust emissions in China is subject  
974 to high levels of uncertainty, with the estimated dust fluxes based on different dust emission schemes differing by several  
975 orders of magnitude (Zeng et al., 2020). Therefore, our inversion results could provide a reference for the magnitude of coarse-  
976 dust emissions in China, which could then help to reduce the large uncertainty in estimations of dust emissions in China.

977 Several potential important deficiencies in current emission estimations were also indicated by CAQIEI on the regional  
978 scale. For example, the CAQIEI suggests substantially higher air pollutant emissions than the previous emission inventories  
979 over the NW and Central regions. Thus, the air pollutant issues may be more severe than we expected over these two regions.  
980 Meanwhile, our inversion results suggest higher NMVOC emissions over the northern China but suggest lower NMVOC  
981 emissions in southern China, which is consistent with the previous inversion studies based on the satellite. China is now facing  
982 increasingly severe O<sub>3</sub> pollution and has an urgent need for a coordinated control of O<sub>3</sub> and PM<sub>2.5</sub>. Our results may provide  
983 valuable information on the NMVOC emissions in China, which is important for a proper understanding of O<sub>3</sub> pollution and  
984 the development of effective control strategies nationally. Higher emissions were also found in the NE region based on our  
985 inversion results. The NE region is a typical area for open-area biomass burning, with significant emissions from straw  
986 combustion (Wu et al., 2020b). The higher emissions estimated by our inversion result may indicate higher biomass-burning  
987 emissions over there. This is consistent with recent estimations of biomass-burning emissions by Xu et al. (2023) and Wu et  
988 al. (2020b), who showed higher biomass-burning emissions in China than previous estimations, including those of GFEDv4.1s  
989 (<https://www.globalfiredata.org/data.html>), FINNv1.5 (<https://www.acom.ucar.edu/Data/fire/>), and GFASv1.2  
990 (<https://www.ecmwf.int/en/forecasts/dataset/global-fire-assimilation-system>).

991 Based on CAQIEI, we further quantified the emission changes of different air pollutants in China during the two clean  
992 air action plans. The results confirmed the effectiveness of these campaigns on the mitigation of air pollutant emissions in  
993 China, with estimated emission reductions of 15.1% for NO<sub>x</sub>, 54.5% for SO<sub>2</sub>, 35.7% for CO, 44.4% for PM<sub>2.5</sub>, and 33.6% for



994 PM<sub>10</sub> from 2015 to 2020. In contrast, NMVOC emissions increased by 21.0% from 2015 to 2020. Comparisons of the estimated  
995 emission reduction rates during the two clean air action plans suggested that emission reductions were larger during the 2018–  
996 2020 than during 2015–2017. The estimated rates of change in emissions were 5.9% for NO<sub>x</sub>, –23.8% for SO<sub>2</sub>, –9.8% for CO,  
997 –14.5% for PM<sub>2.5</sub>, –7.2% for PM<sub>10</sub>, and 27.6% for NMVOCs during 2015–2017, which were smaller than the –12.1% for NO<sub>x</sub>,  
998 –23.5% for SO<sub>2</sub>, –18.3% for CO, –26.6% for PM<sub>2.5</sub>, –25.5% for PM<sub>10</sub>, and –4.5% for NMVOCs during 2018–2020. On the  
999 one hand, this is due to the fact that more sectors were controlled during the 2018–2020 action plan. Besides the industrial and  
1000 power sectors, which were the main points of control in the 2013–2017 action plan, the residential sector, transportation sector,  
1001 and non-point sources like blowing-dust emissions, were also strengthened in the 2018–2020 action plan. Consequently, the  
1002 emission reduction rates of CO, PM<sub>2.5</sub> and PM<sub>10</sub> during 2018–2020 were higher than those during the 2015–2017 when the  
1003 2013–2017 action plan was implemented. However, the reduction of SO<sub>2</sub> emissions was similar during the two action plan  
1004 periods. This is because most SO<sub>2</sub> emissions stem from the industrial sector and power plants, which together contribute about  
1005 77% of all emissions (Zheng et al., 2018). Thus, the additional control of other sectors in the 2018–2020 action plan may not  
1006 have significantly impacted the mitigation of SO<sub>2</sub> emissions. On the other hand, strict emission controls were implemented or  
1007 strengthened in more areas of China during the 2018–2020 action plans. For example, the inversion results indicated that there  
1008 were obvious increases of SO<sub>2</sub>, NO<sub>x</sub>, PM<sub>2.5</sub>, PM<sub>10</sub> and NMVOC emissions during 2015–2017 over the Central region,  
1009 especially in the Fenwei Plain area, where the emission controls were relatively weak during the 2013–2017 action plan.  
1010 However, all species showed obvious emission reductions almost the whole China during the 2018–2020 action plan.

1011 The estimated rates of change in emissions during 2015–2018 were also compared with those estimated by previous  
1012 emission inventories. Although both CAQIEI and previous inventories showed declines of air pollutant emissions in China,  
1013 the emission reduction rates estimated by CAQIEI were generally smaller than those estimated by previous inventories,  
1014 especially for NO<sub>x</sub>, PM<sub>10</sub> and NMVOCs, suggesting a smaller mitigation effects of the air pollution control measures than the  
1015 previous emission inventories suggested. In particular, China's NMVOC emissions were shown to have increased by 26.6%  
1016 from 2015 to 2018, especially over NCP (38.0%), NE (38.3%) and Central (60.0%). CO was found to be an exception insofar  
1017 as the emission reduction rate estimated by CAQIEI was larger than that of most previous emission inventories, except in the  
1018 NCP region. The estimated emission reduction rates of SO<sub>2</sub> and PM<sub>2.5</sub> were relatively closer to those of previous inventories,  
1019 suggesting better consistency in the estimated emission reduction for these two species.

1020 Overall, the inversion inventory developed in this study could provide us with value information on the complex variations  
1021 of air pollutant emissions in China during its two recent clean air action periods, which could help improve our understanding  
1022 of air pollutant emissions and related changes in air quality in China. For example, the increases of O<sub>3</sub> and nitrate  
1023 concentrations may be associated with the undesirable emission reduction effects of the 2013–2017 action plans. The estimated  
1024 lower NO<sub>x</sub> emission reduction rate by CAQIEI may also help explain the weak responses of nitrogen deposition fluxes to the  
1025 clean air action plans. Meanwhile, this top-down emissions inventory can be used to supply the input data for CTMs or server  
1026 as a comparable reference for future inversion studies based on other methods or observation data, which is expected to  
1027 improve the performance of model simulations and air quality forecasts, and facilitate the development of inversion method.

## 1028 **6 Limitations**

1029 However, due to the complexity of the emission estimation, it is inevitable that there are some limitations in our inversion  
1030 results. Here We summarise some issues that might affect the quality of the CAQIEI which were known at the time of  
1031 publication to assist the potential users in properly using this data products.

1032 (1) The changes in the number of observation sites would induce spurious emission trends during 2013–2014, especially  
1033 over western China, although the influence of the number of observation sites is smaller over the NCP and SE regions because  
1034 of their higher density of observation sites. Therefore, it is recommended that not to use the emissions in 2013 and 2014 when

1035 analyzing the emission trends in China. This limitation makes it difficult to estimate the overall emission control effects of  
1036 2013 – 2017 action plan. Consequently, the emission change rate during the 2015–2017 were sampled in this study to represent  
1037 the emission control effects of the 2013–2017 action plan, but it may not necessarily reflect the overall reduction rate of the  
1038 action plan for the entire period. In addition, although the number of observation sites has become stable since 2015, the limited  
1039 number of observation sites makes it difficult to fully constrain China’s air pollutant emissions, especially for the natural  
1040 sources considering that the majority of the observation sites are located in the urban areas. Therefore, the uncertainty in the  
1041 estimated emissions over the remote areas are expected to be higher than those over the urban areas, especially for the species  
1042 with large amount of natural emission, such as PM and NMVOC. For example, the coarse-dust emissions over western China  
1043 are expected to be underestimated by CAQIEI because of the limited availability of observation sites. Therefore, adding  
1044 observations there will help improve the accuracy of the inversion estimates. For example, -simultaneous assimilation of the  
1045 surface and satellite observation may help alleviate this problem and provide more constrains on the emissions without surface  
1046 observations.

1047 (2) The natural and anthropogenic emissions are not differentiated in our inversion method, leading to higher emissions  
1048 of PM<sub>10</sub> and NMVOCs than in other emission inventories. This also hinders the comparisons of our inversion results with the  
1049 previous inventories. Therefore, potential readers should be aware of that the current comparisons of our inversion results and  
1050 previous inventories are on the basis of the natural emissions estimated by CAMS and GFAS, which does not necessarily  
1051 indicate large uncertainties in anthropogenic sources within the bottom-up inventories. The impacts are expected to be smaller  
1052 for the NO<sub>x</sub>, SO<sub>2</sub> and CO due to the small contributions of natural sources to their emission, but would be larger for NMVOC  
1053 and PM which has large amount of natural emission. Assimilation of isotope data, speciated PM<sub>2.5</sub> and NMVOC observations  
1054 may help differentiate the natural and anthropogenic emissions, and address this problem in future.

1055 (3) The NMVOC emissions may have larger uncertainty than the other species. On the one hand, a significant amount of  
1056 NMVOC emission would originate from suburban or rural regions. Therefore, although the O<sub>3</sub> observations at the urban sites  
1057 could provide information on the NMVOC emissions over the suburban or rural areas according to covariance estimated by  
1058 the ensemble simulation, the NMVOC emissions may not be fully constrained due to the lack of observation sites over the  
1059 suburban or rural areas. On the other hand, due to the lack of long-term NMVOC observations, the NMVOC emissions were  
1060 constrained by the O<sub>3</sub> concentrations in this study. Although the feasibility of this approach has been demonstrated by previous  
1061 inversion studies, the nonlinear NO<sub>x</sub>-VOC-O<sub>3</sub> interactions could inevitably introduces greater uncertainty into the inversion of  
1062 NMVOC than other species. Therefore, more attention should be paid while using the inversion results of NMVOC, and more  
1063 robust analysis of the effects of nonlinear NO<sub>x</sub>-VOC-O<sub>3</sub> interactions and the number of observation sites should be performed  
1064 in future to better illustrate the feasibility of assimilating O<sub>3</sub> to constrain the NMVOC emissions.

1065 (4) The errors in the meteorological simulation and the CTMs were not considered in the emission inversions, which  
1066 would lead to uncertainty in our estimated emissions. For example, the errors in the simulated wind would influence the  
1067 transportation of the air pollutant and lead to uncertainty in the emissions distributions. According to the evaluation results of  
1068 meteorological simulations (Table S1), the simulated relative humidity is generally lower than the observations, which may  
1069 weaken the formation of secondary aerosol. On the contrary, the simulated precipitation was higher than the observation for  
1070 most regions which would lead to overestimations of the wet removal of air pollutants. As a result, there may be a positive  
1071 tendency in the inversed emission inventory due to the errors in the simulated relative humidity and precipitation. Besides  
1072 these parameters, the accuracy of the simulated boundary layer is also important for the performance of the emission inversions  
1073 (Du et al., 2020), although it was not evaluated currently due to the lack of observation. If the WRF systematically  
1074 underestimates the boundary layer, the vertical diffusions of the air pollutants would be suppressed, which would lead to  
1075 overestimated surface air pollutant concentrations and a negative tendency in the inverse emission inventory. However, it is  
1076 difficult to consider the meteorological and model errors in the assimilation process. it is difficult to quantify the influences of  
1077 the meteorological errors on the emission inversions, as the errors in the meteorological simulation and chemical transport

1078 model interact with each other. More comprehensive analysis should be conducted in the future to better understand the impacts  
1079 of the meteorological and model errors on the inverse emission inventory. A multi-model inversion framework, for example  
1080 that of Miyazaki et al. (2020a), may help alleviate the influences of model errors on emission inversions in future. Using other  
1081 models (e.g., WRF-Chem, CMAQ) to validate our inversion inventory could also help us assess the impacts of model  
1082 uncertainty on the emission inversions. Meanwhile, because of the many uses that require a rapid update of emissions, it may  
1083 be time to organize an intercomparison study focused on the emission inversions.

1084 (4)(5) Current inversion emission inventory is mainly assessed by the surface observations and previous emission  
1085 inventories. more independent observations, such as the satellite observation data, should be used in future to further validate  
1086 the inversion results of this study and its derived findings. For example, the independent measurements from field campaign  
1087 or satellite retrievals (e.g., TropOMI CO data) can help validate the reliability of the much higher a posterior CO emissions in  
1088 CAQIEI than the previous inventories in the future.

## 1089 7 data availability

1090 The CAQIEI inventory can be freely download at <https://doi.org/10.57760/sciencedb.13151> (Kong et al., 2023), which  
1091 includes monthly grid maps of the air pollutant emissions from 2013 to 2020. The contained species include NO<sub>x</sub>, SO<sub>2</sub>, CO,  
1092 primary PM<sub>2.5</sub>, primary PM<sub>10</sub> and NMVOC. The horizontal resolution is 15km. There are totally 8 Network Common Data  
1093 Form files (NetCDF), which were named by the date and contains the monthly emissions of different air pollutants in China  
1094 in each year. The description of the content of each NetCDF file and some important notes when using this dataset are also  
1095 available in README.txt on the website.

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## 1097 Tables

1098 **Table 1. Corresponding relationships between the chemical observations and adjusted emissions**

Species	Description	Observations used for inversions of this species
BC	Black carbon	PM <sub>2.5</sub>
OC	Organic carbon	PM <sub>2.5</sub>
PMF	Fine-mode unspciated aerosol	PM <sub>2.5</sub>
PMC	Coarse-mode unspciated aerosol	PM <sub>10</sub> – PM <sub>2.5</sub>
NO <sub>x</sub>	Nitrogen oxide	NO <sub>2</sub>
SO <sub>2</sub>	Sulfur dioxide	SO <sub>2</sub>
CO	Carbon monoxide	CO
NMVOCs	Non-methane volatile organic compounds	MDA8h O <sub>3</sub>

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1106 **Table 2. Evaluation statistics of the *a posteriori* (*a priori*) model simulation for different species<sup>a</sup>**

	PM <sub>2.5</sub> (µg/m <sup>3</sup> )				PM <sub>10</sub> (µg/m <sup>3</sup> )			
	R	MBE	NMB (%)	RMSE	R	MBE	NMB (%)	RMSE
Hourly	0.77 (0.53)	2.1 (13.3)	4.5 (28.6)	32.4 (55.6)	0.72 (0.44)	-3.7 (-11.5)	-4.6 (-14.3)	53.1 (74.4)
Daily	0.89 (0.61)	2.1 (13.3)	4.4 (28.4)	20.0 (46.3)	0.88 (0.51)	-3.7 (-11.2)	-4.6 (-14.1)	31.6 (62.2)
Monthly	0.94 (0.68)	2.1 (13.3)	4.5 (28.3)	11.7 (32.5)	0.90 (0.56)	-3.6 (-11.3)	-4.5 (-14.1)	21.2 (44.1)
Yearly	0.94 (0.62)	2.2 (11.9)	4.4 (24.3)	9.1 (27.7)	0.89 (0.52)	-3.8 (-13.4)	-4.6 (-16.1)	18.5 (38.7)
	SO <sub>2</sub> (µg/m <sup>3</sup> )				NO <sub>2</sub> (µg/m <sup>3</sup> )			
	R	MBE	NMB (%)	RMSE	R	MBE	NMB (%)	RMSE
Hourly	0.64 (0.16)	-1.8 (19.0)	-9.1 (93.8)	24.9 (58.7)	0.67 (0.45)	-1.2 (-0.9)	-3.9 (-2.7)	19.9 (25.5)
Daily	0.80 (0.20)	-1.8 (19.0)	-9.2 (94.5)	16.0 (51.4)	0.80 (0.51)	-1.2 (-0.8)	-3.7 (-2.6)	12.8 (20.1)
Monthly	0.85 (0.20)	-1.9 (18.9)	-9.3 (93.1)	12.4 (45.8)	0.84 (0.57)	-1.2 (-0.8)	-3.8 (-2.6)	9.4 (15.6)
Yearly	0.83 (0.18)	-2.4 (17.0)	-10.8 (75.9)	11.6 (42.4)	0.82 (0.63)	-1.3 (-1.6)	-3.9 (-5.0)	8.1 (13.0)
	CO (mg/m <sup>3</sup> )				O <sub>3</sub> (µg/m <sup>3</sup> )			
	R	MBE	NMB (%)	RMSE	R	MBE	NMB (%)	RMSE
Hourly	0.69 (0.38)	-0.1 (-0.4)	-8.8 (-45.6)	0.6 (0.8)	0.71 (0.51)	5.6 (-8.4)	9.5 (-14.0)	34.9 (41.6)
Daily	0.81 (0.42)	-0.1 (-0.4)	-8.6 (-45.5)	0.4 (0.7)	0.71 (0.40)	5.7 (-8.4)	9.5 (-14.1)	26.1 (33.8)
Monthly	0.83 (0.42)	-0.1 (-0.4)	-8.7 (-45.7)	0.3 (0.7)	0.76 (0.47)	5.6 (-8.4)	9.4 (-14.1)	19.6 (26.0)
Yearly	0.82 (0.27)	-0.1 (-0.5)	-9.0 (-47.6)	0.3 (0.7)	0.53 (0.11)	5.1 (-7.8)	8.7 (-13.4)	14.2 (20.5)

<sup>a</sup>The time series of the air pollutant concentrations at each station were firstly catenated into a single vector. Then the values of each evaluation metric were calculated based on the catenated time series of the observed and simulated concentrations.

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1127 **Table 3. Inversion-estimated emissions (Tg/yr) of different species in China as well as the six regions for year 2015**

	China	NCP	SE	NE	SW	NW	Central
NO <sub>x</sub>	25.2	5.1	7.1	4.5	4.2	1.2	3.2
SO <sub>2</sub>	17.8	3.5	3.3	4.0	2.6	0.8	3.6
CO	465.4	82.2	106.7	78.7	82.8	32.6	82.3
PM <sub>2.5</sub>	14.9	2.7	3.3	3.1	2.9	1.2	1.9
PM <sub>10</sub>	40.1	8.7	7.5	8.2	5.5	4.1	6.2
NMVOC	46.0	9.0	13.7	8.5	7.8	2.7	4.2

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1134 **Table 4. The calculated annual trends of PM<sub>2.5</sub> and PM<sub>10</sub> emissions in China based on CAQIEI**

	PM <sub>2.5</sub> (Tg/year)			PM <sub>10</sub> (Tg/year)		
	2015–2020	2015–2017	2018–2020	2015–2020	2015–2017	2018–2020
China	-1.4*	-1.1	-1.5	-2.6*	-1.4	-4.6
NCP	-0.32*	-0.30	-0.32	-0.64*	-0.88	-0.99
SE	-0.32*	-0.21	-0.44	-0.52*	-0.48	-0.84
NE	-0.24*	-0.25	-0.11	-0.52*	-0.22	-0.73
SW	-0.21*	-0.26	-0.20	-0.40*	-0.26	-0.56
NW	-0.09	-0.08	-0.12	-0.20*	-0.32	-0.32
Central	-0.15	0.01	-0.32	-0.27	-0.32	-1.14

1135 \* Trend is significant at the 0.05 significance level

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1154 **Table 5. The calculated annual trends of the four gaseous emissions in China based on CAQIEI**

	SO <sub>2</sub> (Tg/year)			CO (Tg/year)		
	2015–2020	2015–2017	2018–2020	2015–2020	2015–2017	2018–2020
China	-2.1*	-2.1	-1.3	-36.0*	-22.8	-33.5
NCP	-0.57*	-0.69	-0.21	-8.4*	-4.30	-7.23
SE	-0.34*	-0.39	-0.20	-6.1*	-3.54	-8.37
NE	-0.44*	-0.44	-0.21	-6.2*	-1.74	-3.91
SW	-0.22*	-0.27	-0.17	-3.8*	-2.36	-4.54
NW	-0.08*	-0.08	-0.08	-3.0*	-0.73	-2.95
Central	-0.46*	-0.25	-0.40	-8.7*	-10.14	-6.55
	NO <sub>x</sub> (Tg/year)			NMVOC (Tg/year)		
	2015–2020	2015–2017	2018–2020	2015–2020	2015–2017	2018–2020
China	-0.67	0.74	-1.6	1.9	6.3	-1.3
NCP	-0.32	0.05	-0.40	0.66	1.37	-0.42
SE	-0.22	0.18	-0.49	0.50	1.73	-0.24
NE	-0.17	0.03	-0.19	0.03	0.79	-0.49
SW	-0.06	0.10	-0.26	0.23*	0.43	0.03
NW	-0.03	0.11	-0.06	0.10	0.69	-0.27
Central	0.04	0.28	-0.16	0.55*	1.33	0.09

\* Trend is significant at the 0.05 significance level

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1179 **Table 6 The top-down estimated CO emissions in China from previous inventories**

Reference	Region	Period	Method	Assimilated observation	<i>A priori</i> CO emission (kt/day)	<i>A posteriori</i> CO emission (kt/day)
Feng et al. (2020)	China	December 2013			586.4	1678.0
	Mainland	December 2017	EnKF with	Surface	499.3	1388.1
	NCP	December 2013	CMAQ model	observation	143.9	394.3
		December 2017			120.5	340.7
Muller et al. (2018)	China	2013	4DVar with IMAGES model	IASI CO observation with different constraints on OH levels	454.8	367.1–553.4
Gaubert et al. (2020)	Central China	May 2016	DART/CAM-CHEM	MOPITT CO observation	193.6	220.3
	North China				93.5	163.6
Jiang et al. (2017)	East China	2013	4DVar with GEOS-Chem	MOPITT CO observation	564.5	439.5–484.4
		2014				430.4–481.1
		2015				397.5–439.7
Zheng et al. (2019)	China	2010–2017 average	Bayesian inversion	MOPITT CO, OMI HCHO, and GOSAT CH <sub>4</sub> observation	-	444.4

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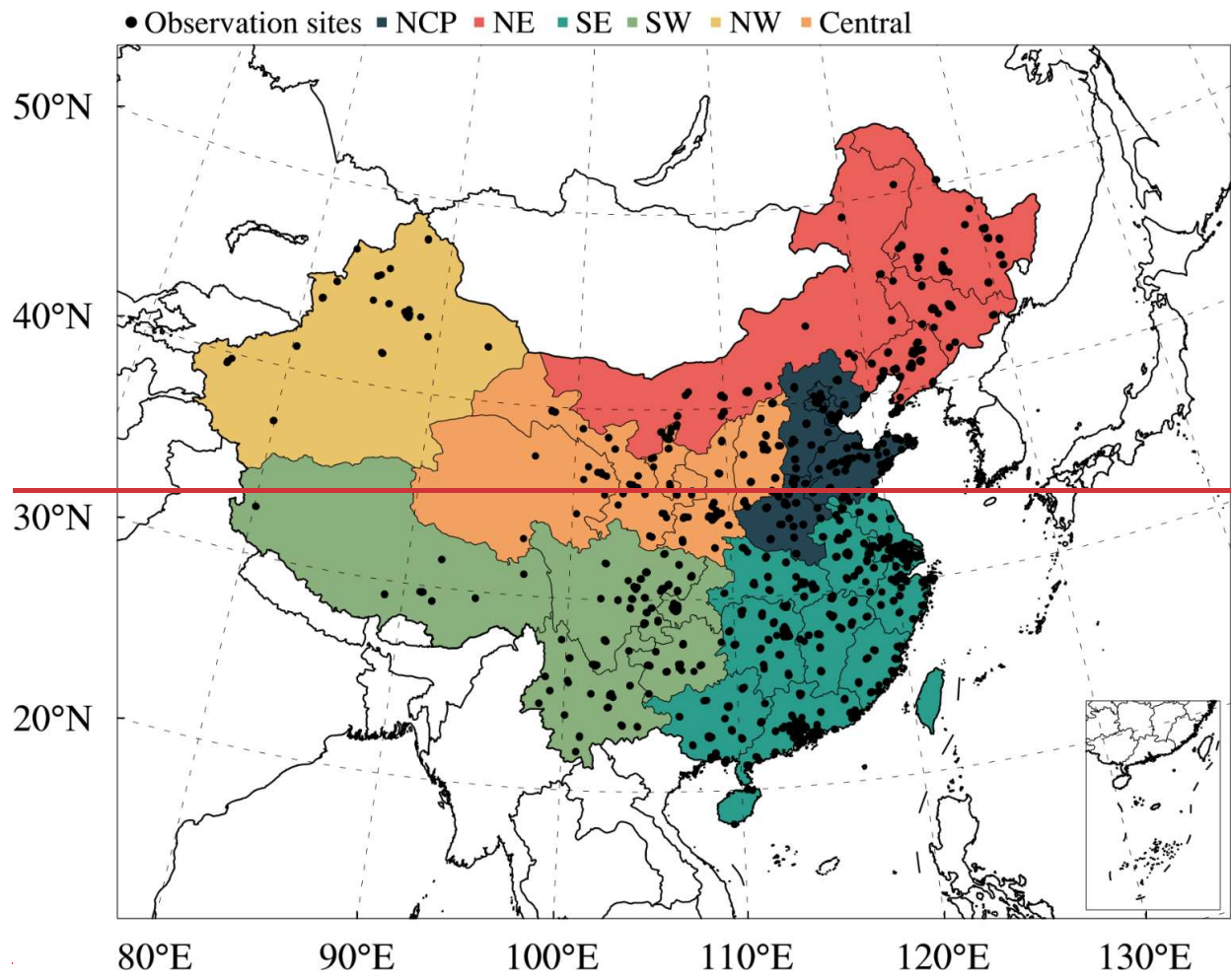
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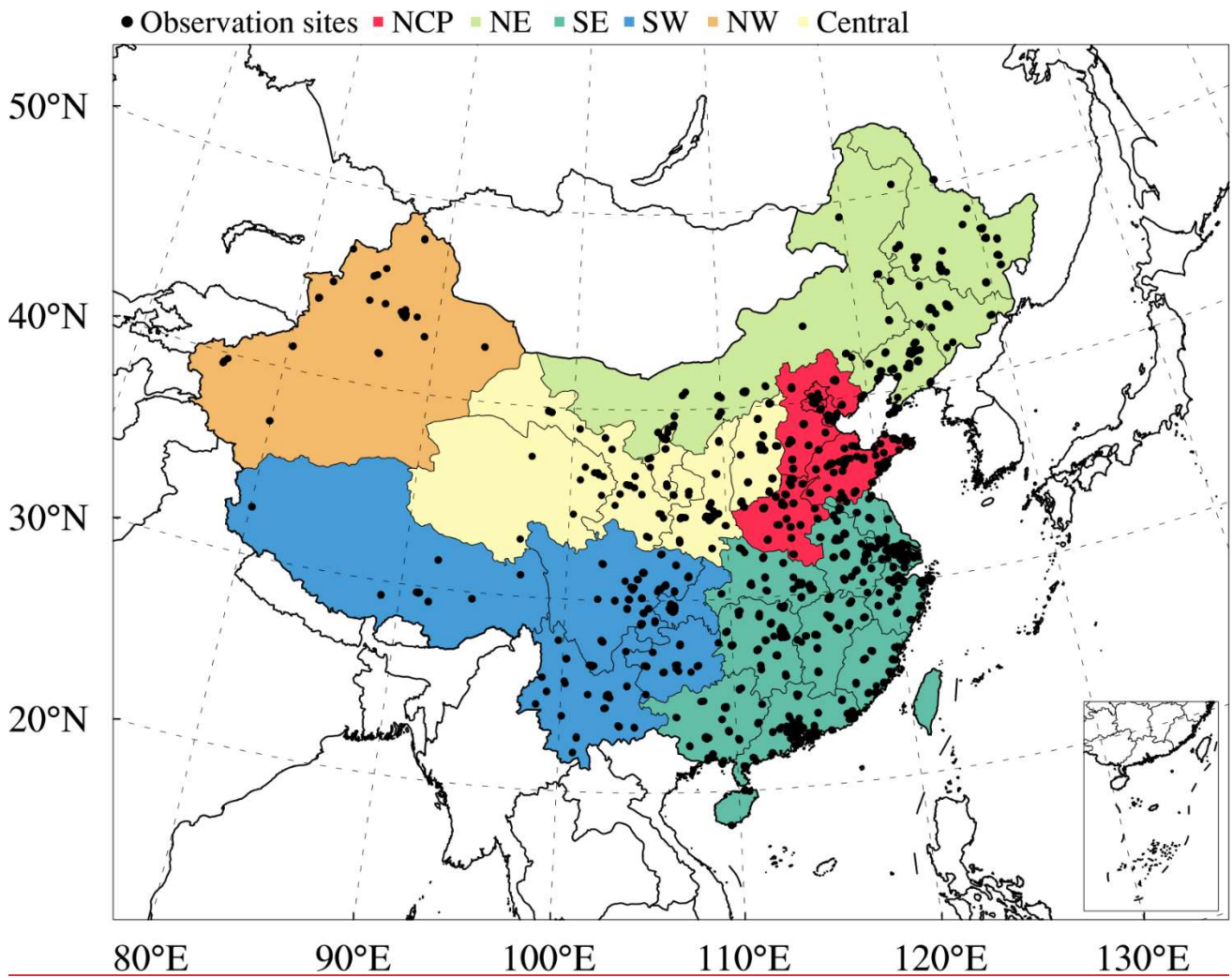
1194 **Figures**

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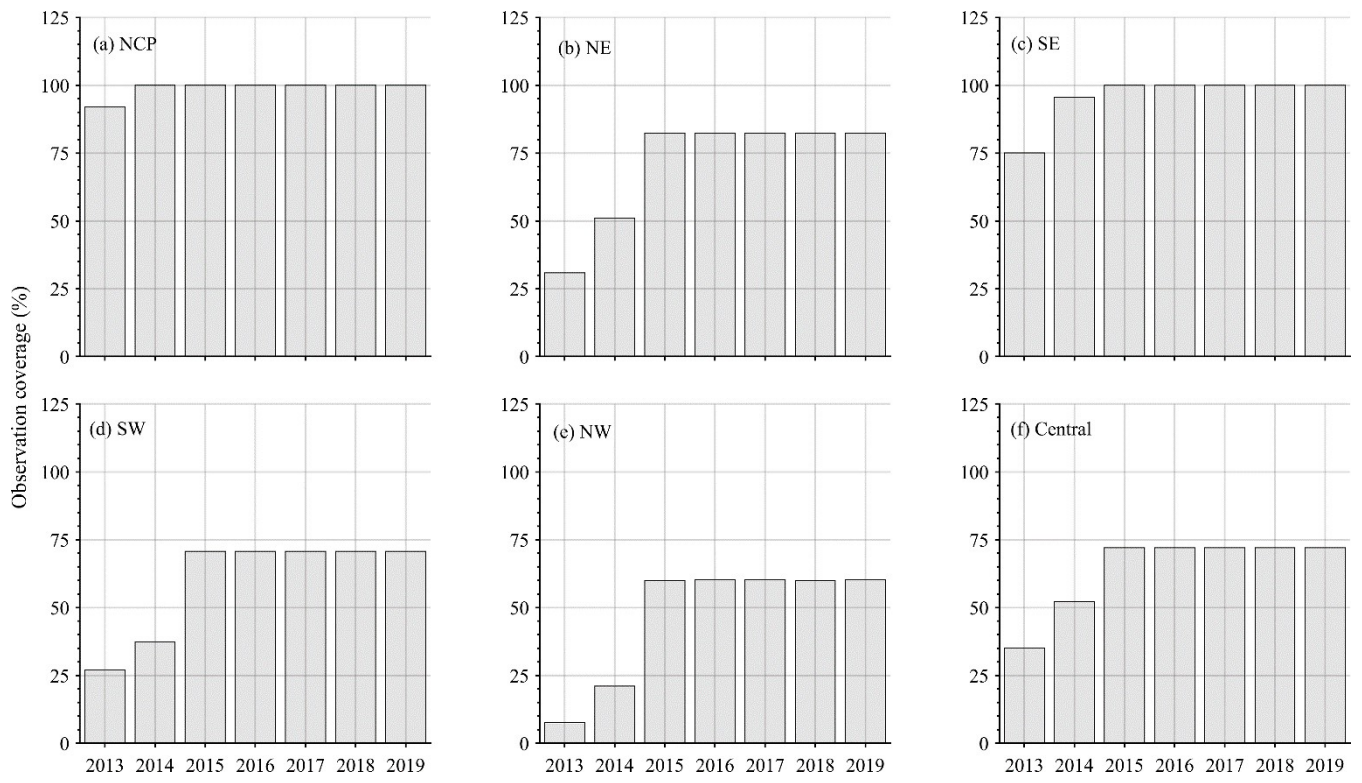


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1198 **Figure 1: Modeling domain of the ensemble simulation overlaid with the distributions of observation sites from CNEMC. Different**  
 1199 **colors denote the different regions in mainland China—namely, the North China Plain (NCP), Northeast China (NE), Southwest**  
 1200 **China (SW), Southeast China (SE), Northwest China (NW) and Central China (Central).**

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1204 **Figure 2: Time series of the observational coverage from 2013 to 2020 over different regions of China.**

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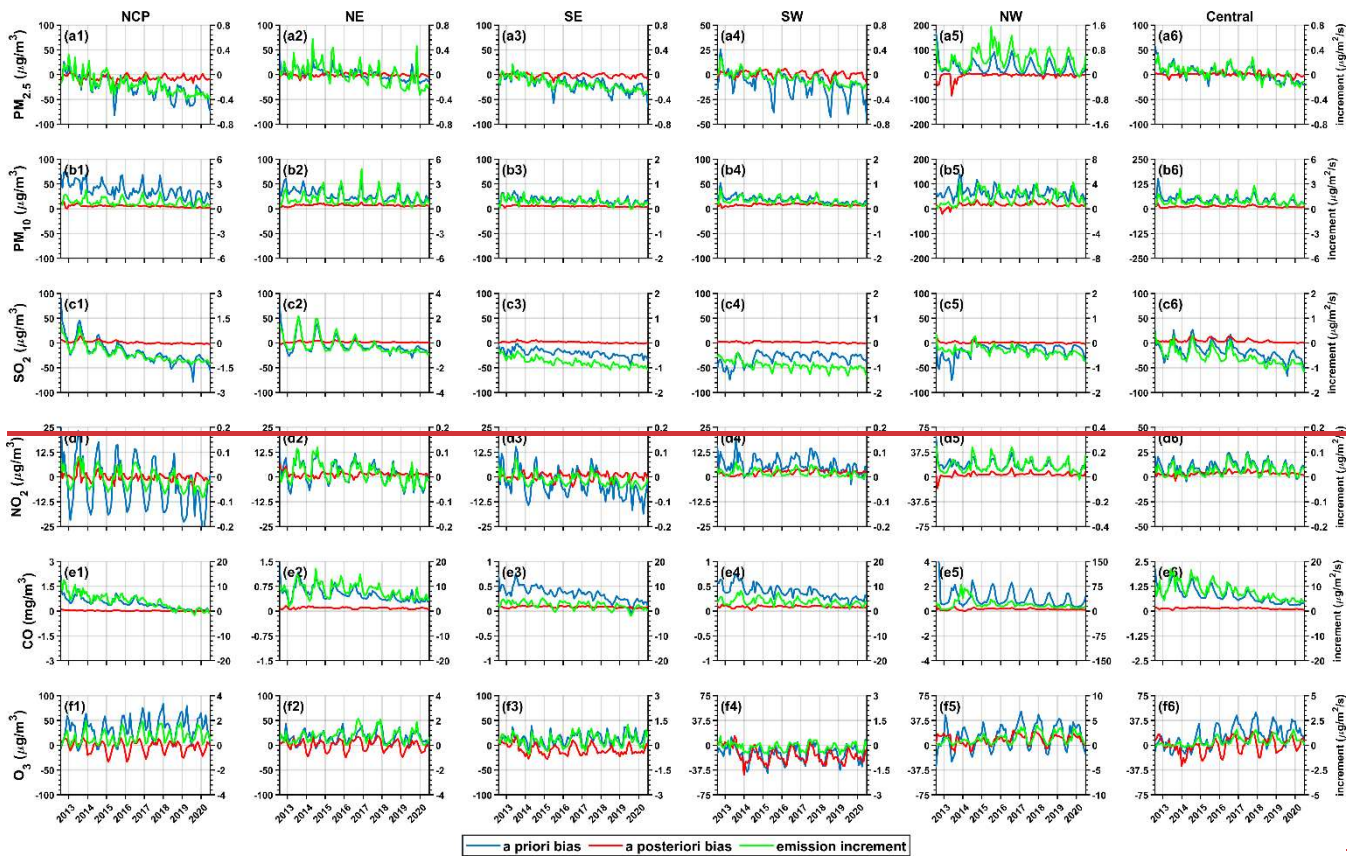
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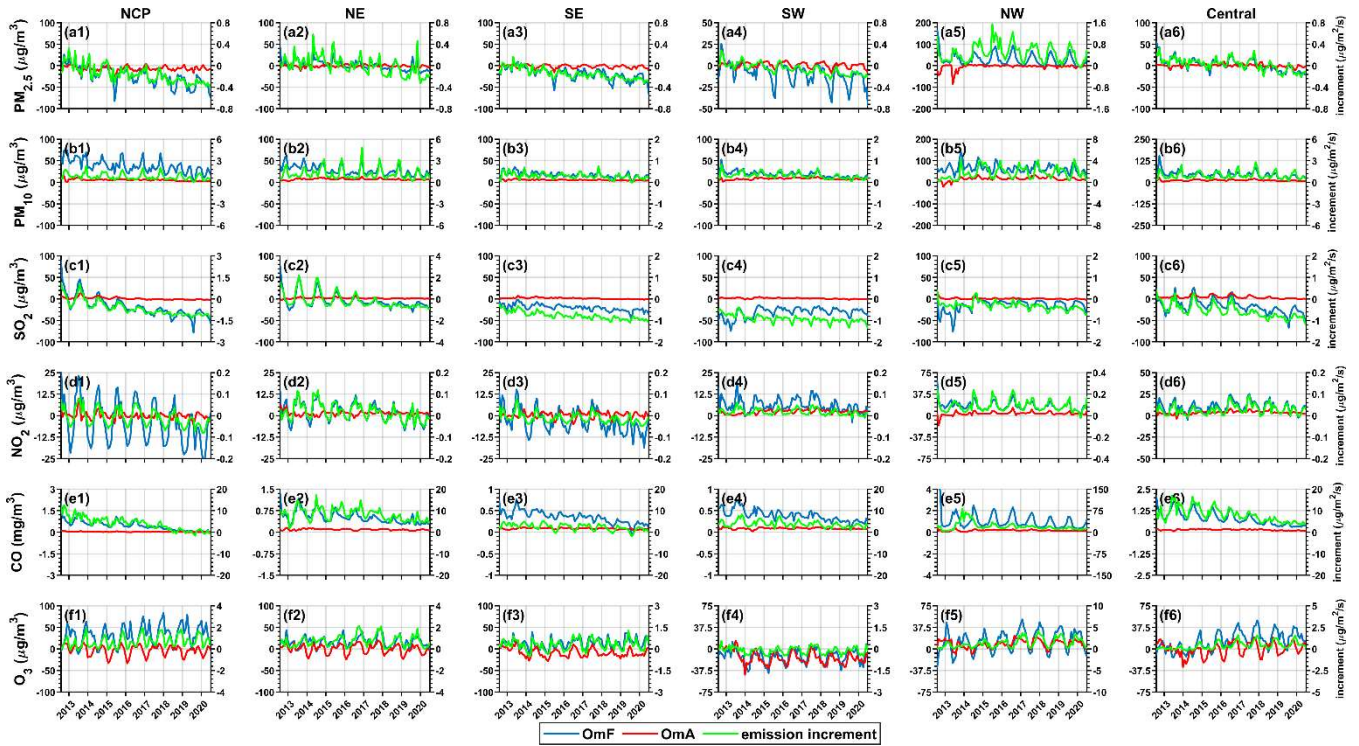
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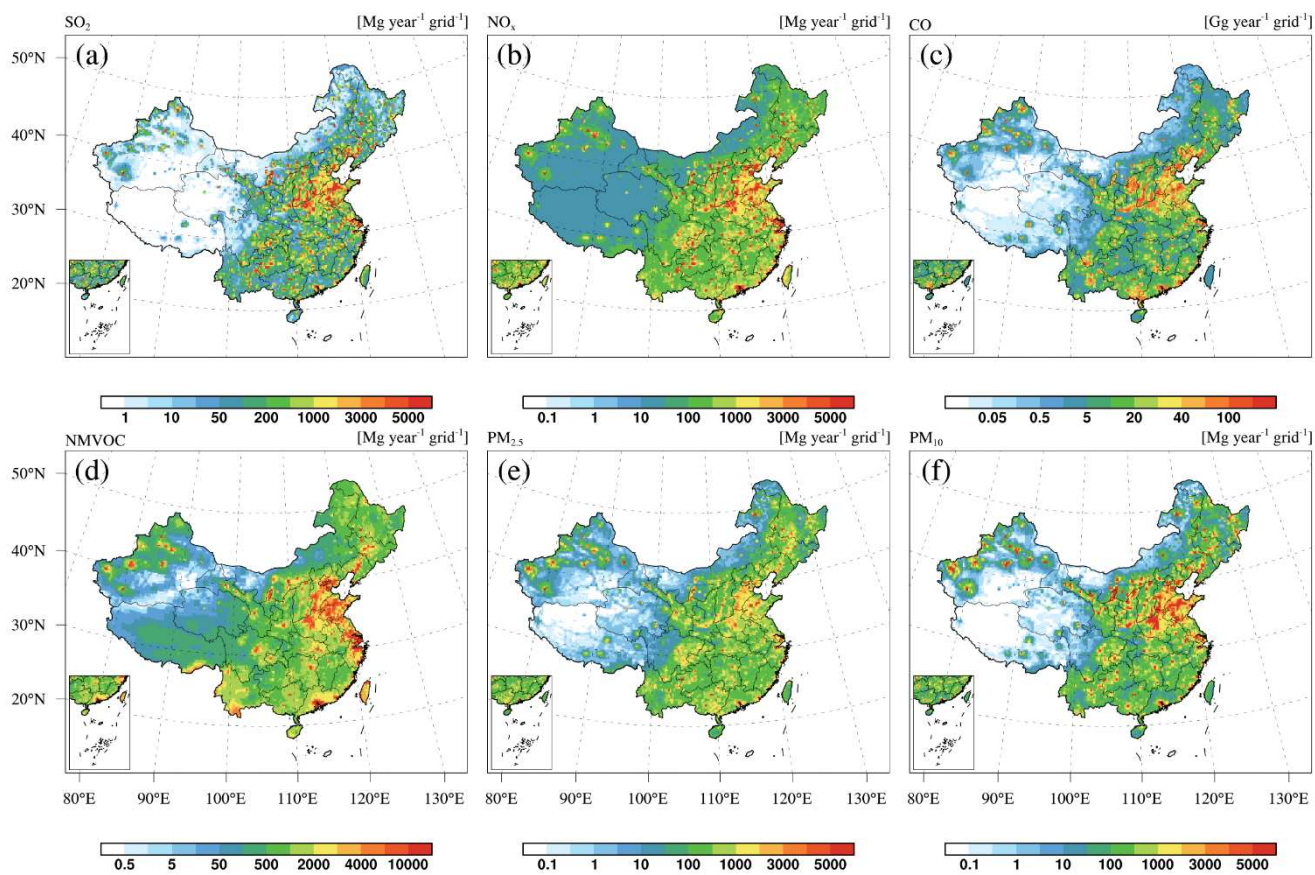


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214 **Figure 3: Time series of the *a priori* bias (blue lines), the *a posteriori* bias (red lines), and the emission increment (green lines) from**  
 215 **2013 to 2020 for different species over the six regions of China.**

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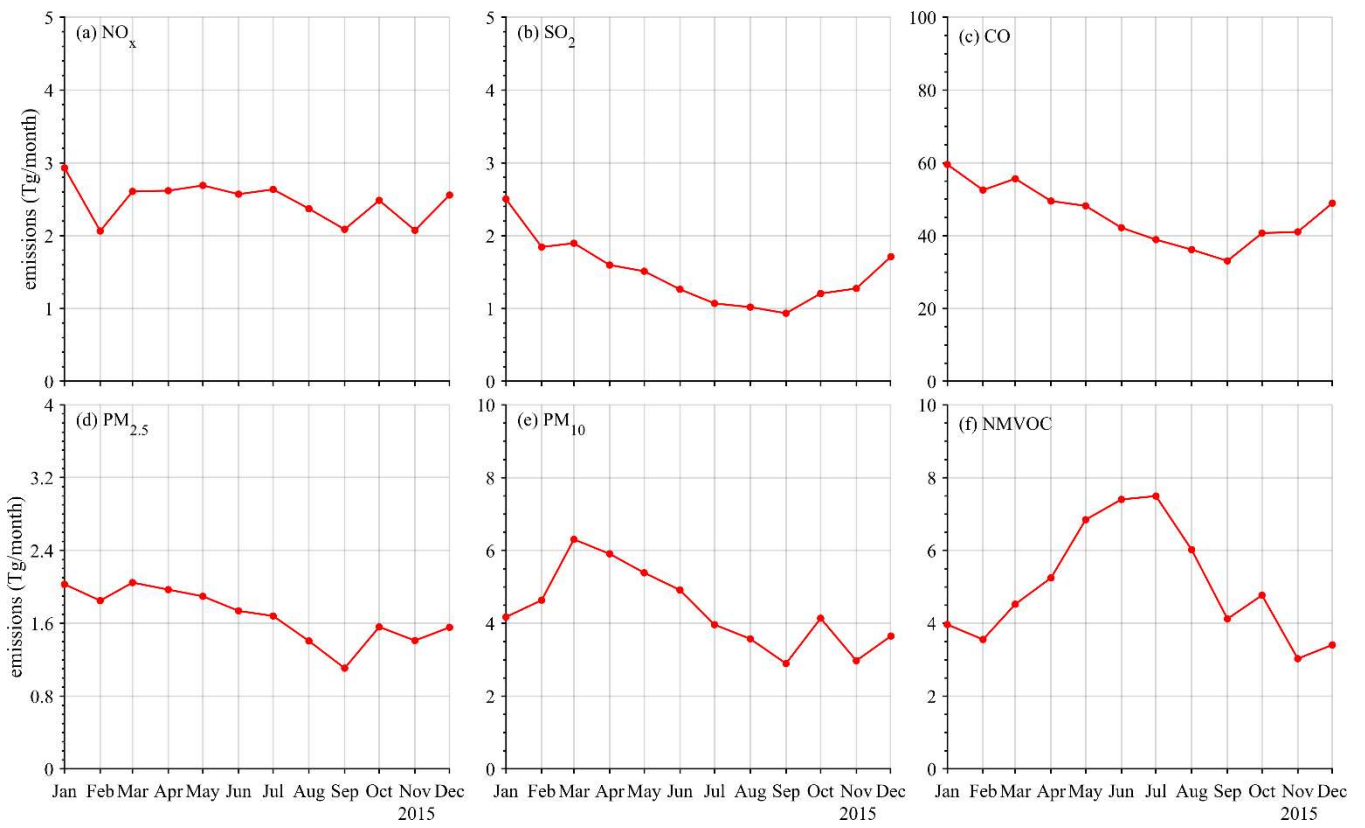
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1219 **Figure 4: Spatial distributions of the emissions of (a)  $\text{SO}_2$ , (b)  $\text{NO}_x$ , (c) CO, (d) NMVOCs, (e)  $\text{PM}_{2.5}$ , and (f)  $\text{PM}_{10}$  in 2015 obtained**  
 1220 **from CAQIEI.**

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1225 **Figure 5: Monthly series of total emissions of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub>, and (f) NMVOCs in China for year 2015**  
 1226 **obtained from CAQIEI.**

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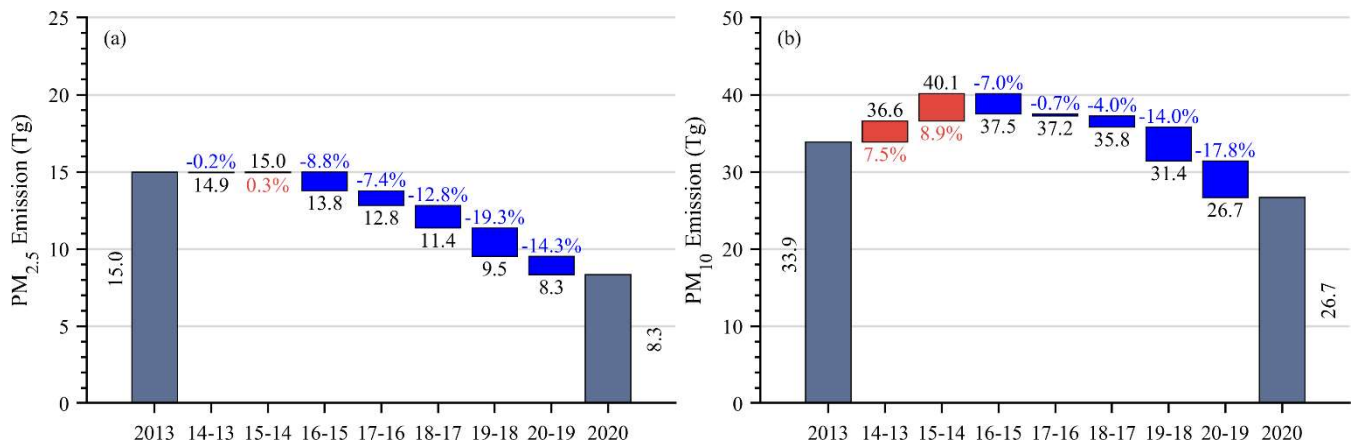
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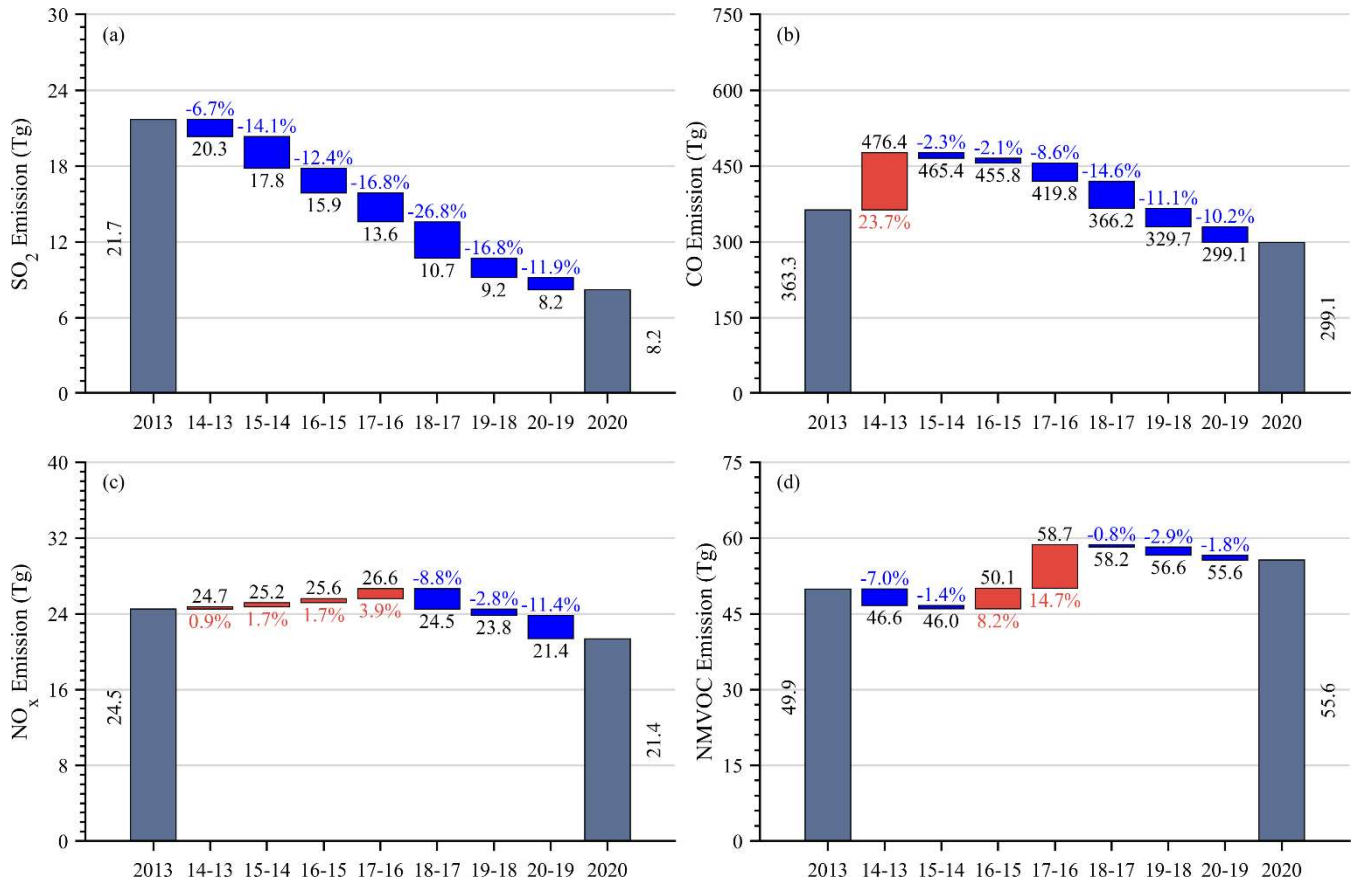
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**Figure 6: Emission changes in (a) PM<sub>2.5</sub> and (b) PM<sub>10</sub> obtained from CAQIEI from 2013 to 2020.**

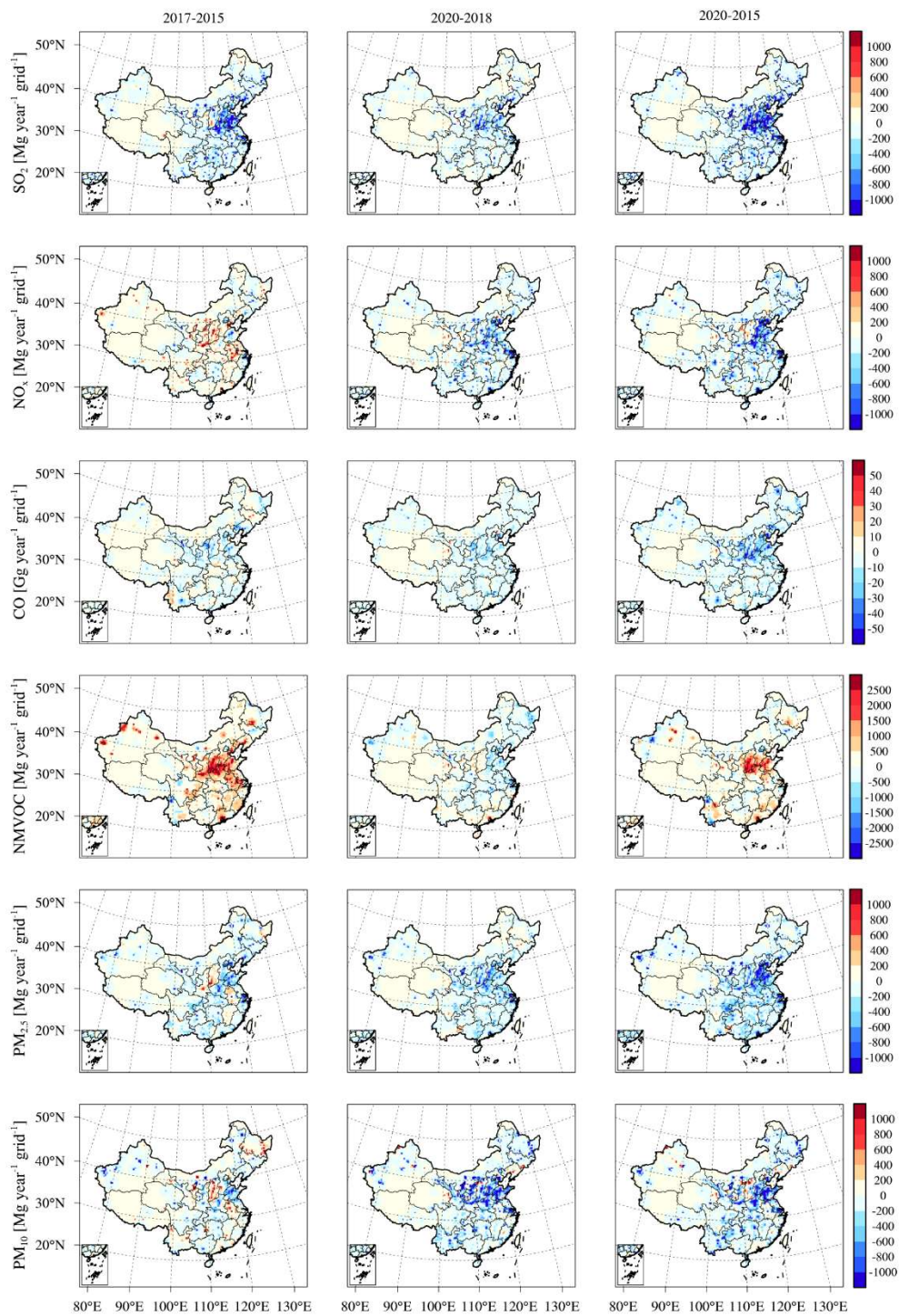


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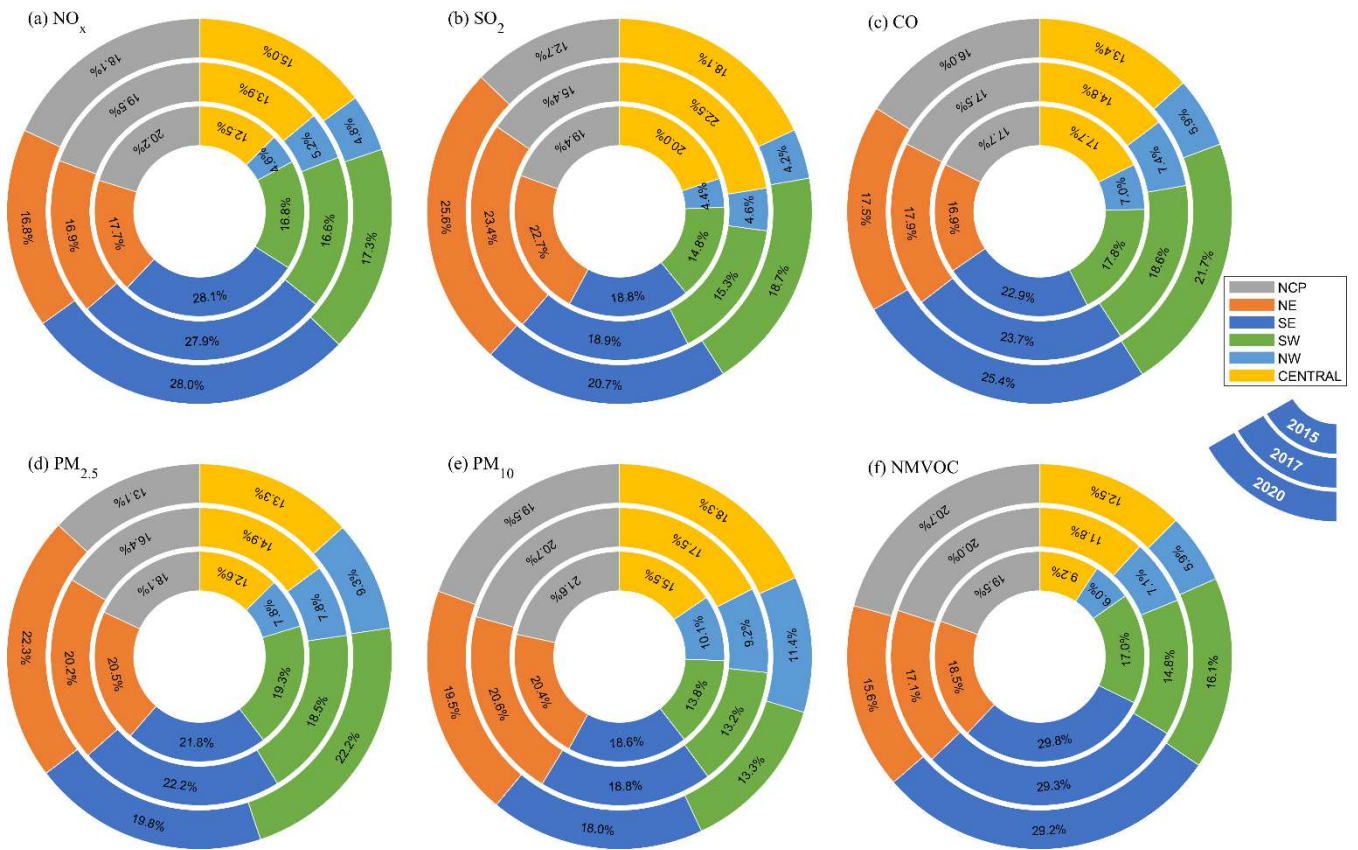
**Figure 7: Emission changes in (a) SO<sub>2</sub>, (b) CO, (c) NO<sub>x</sub>, and (d) NMVOCs obtained from CAQIEI from 2013 to 2020.**

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1248 **Figure 8: Spatial distributions of the emission changes of different species during 2015–2017 (left panels), 2018–2020 (middle panels),**  
 1249 **and 2015–2020 (right panels) obtained from CAQIEI from 2013 to 2020.**



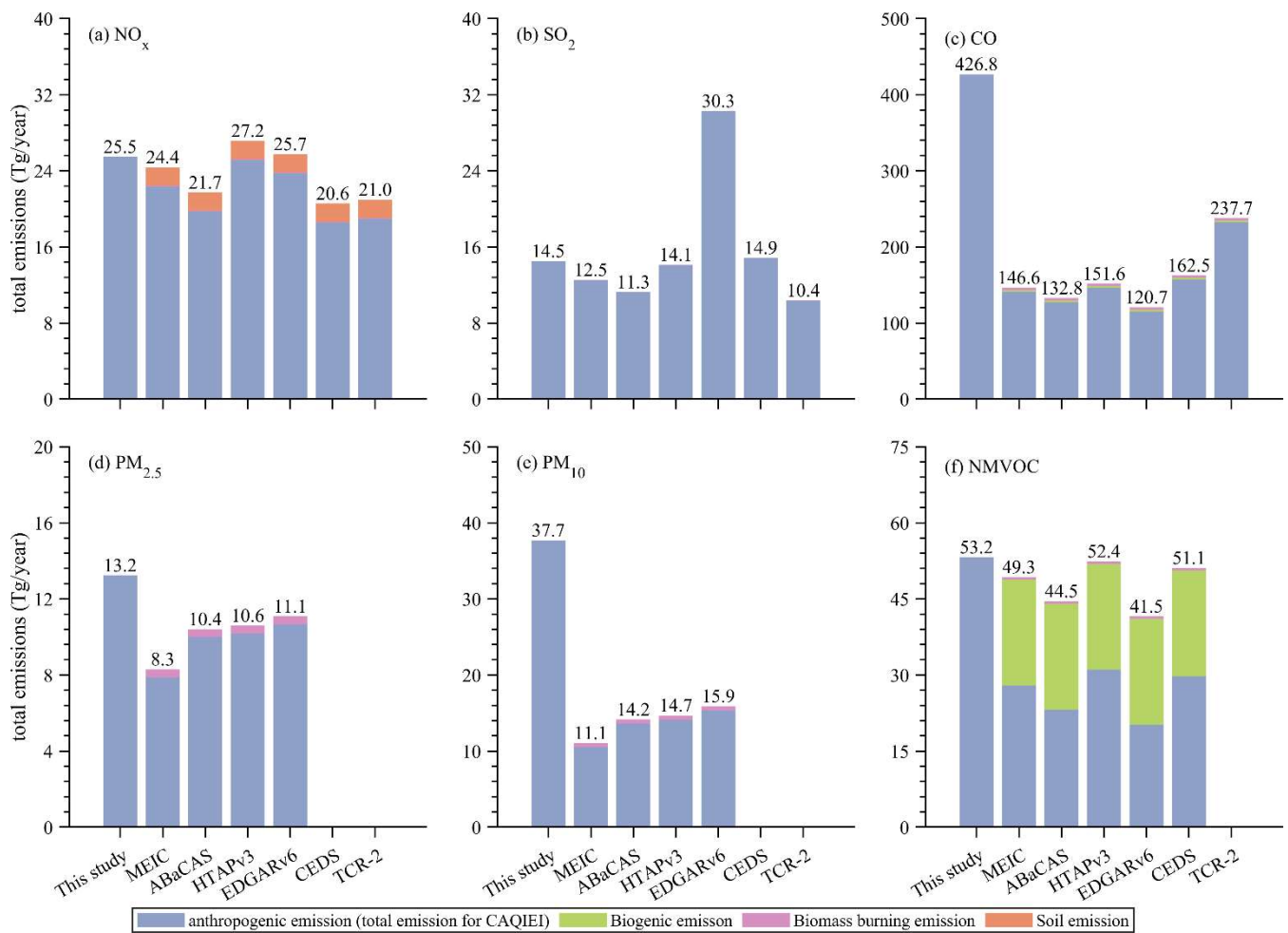
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1251 **Figure 9: Emission distributions of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub>, and (f) NMVOCs among different regions in China**  
 1252 **obtained from CAQIEI in 2015, 2017 and 2020.**

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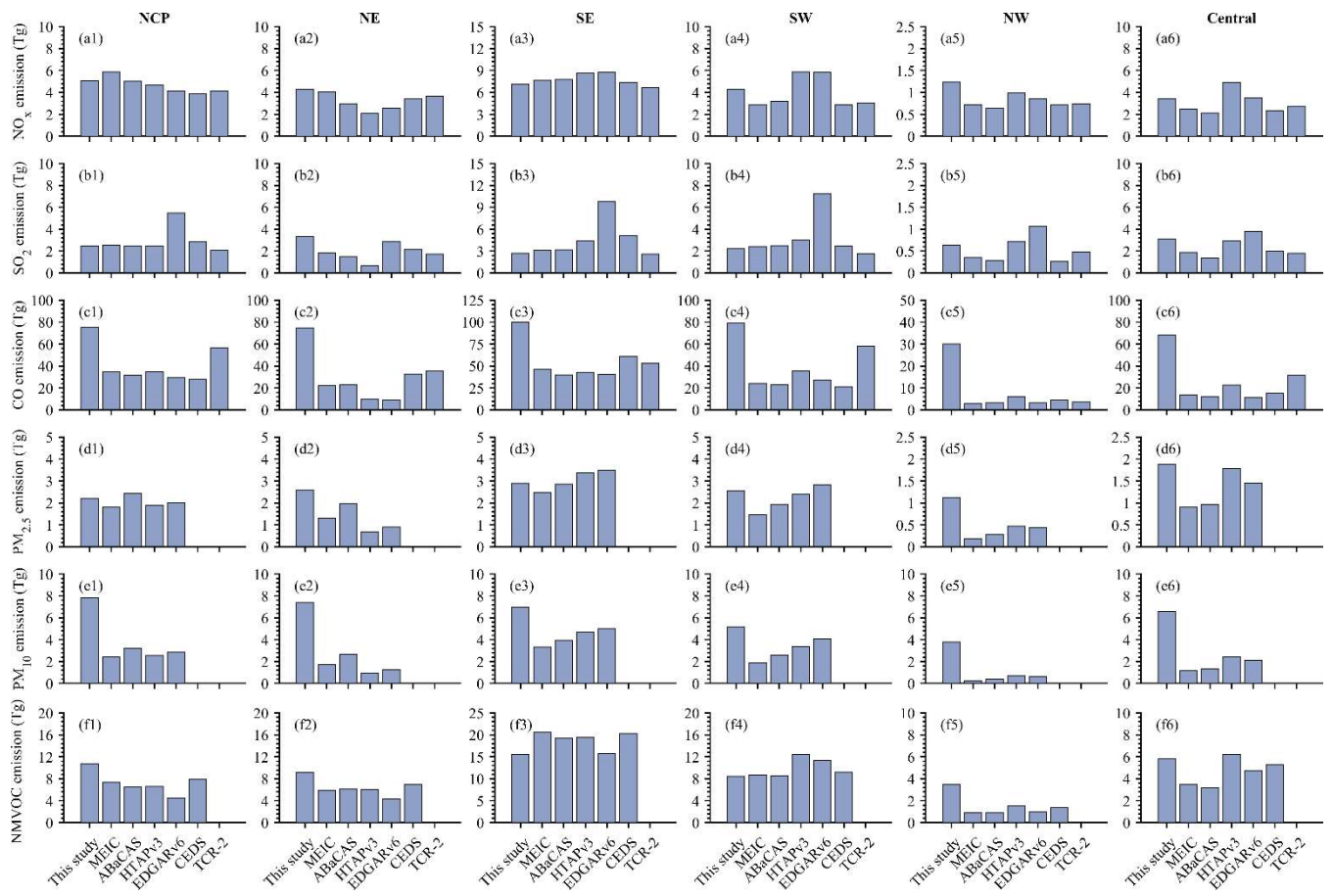




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1256 **Figure 10: Comparisons of the averaged emissions of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub>, and (f) NMVOCs over China**  
 1257 **from 2015 to 2018 between CAQIEI and previous inventories added with natural sources.**

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**Figure 11: Comparisons of the averaged emissions of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub>, and (f) NMVOCs over different regions in China from 2015 to 2018 between CAQIEI and previous inventories added with natural sources.**

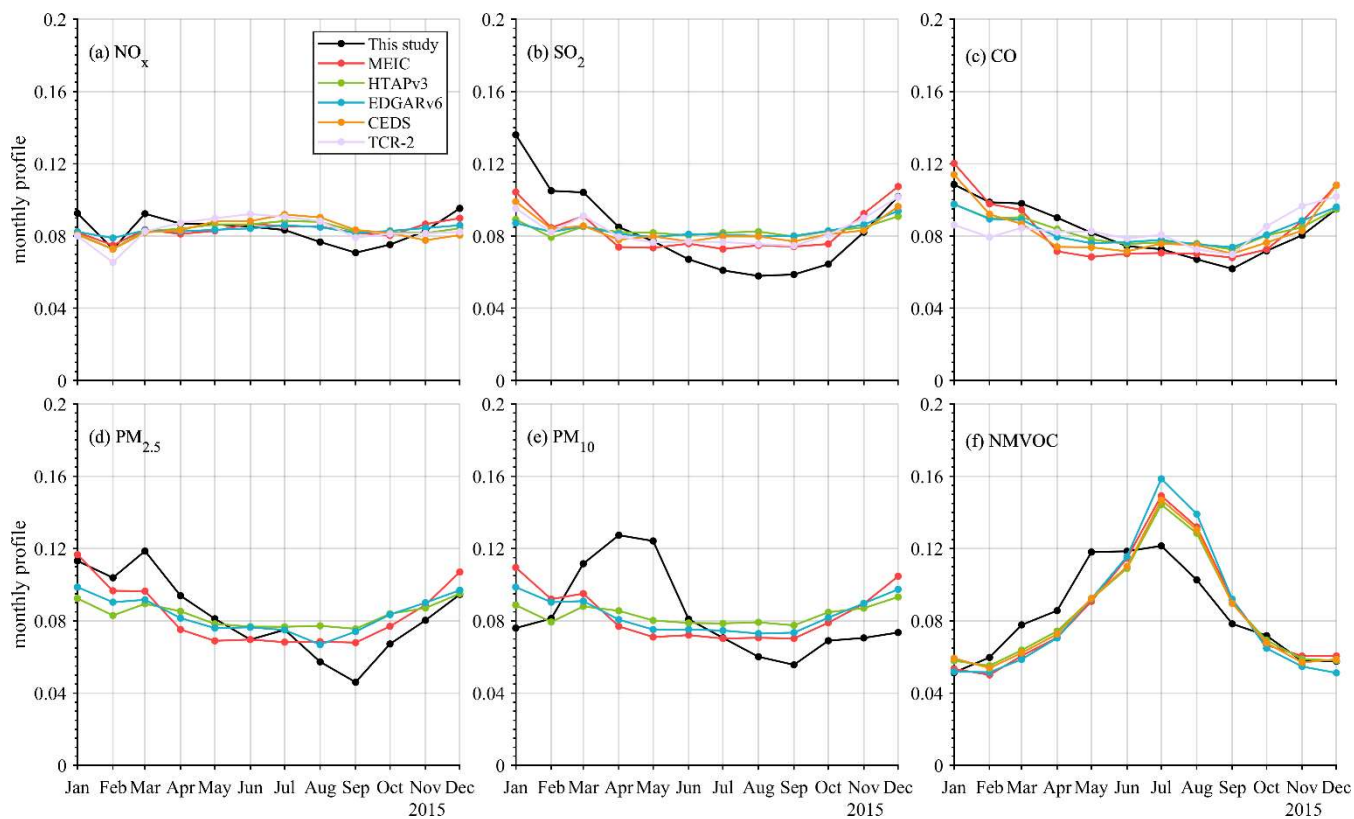
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1268 **Figure 12: Comparisons of the monthly profiles of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub>, and (f) NMVOCs over China**  
 1269 **averaged from 2015 to 2018 between CAQIEI and previous inventories added with natural sources.**

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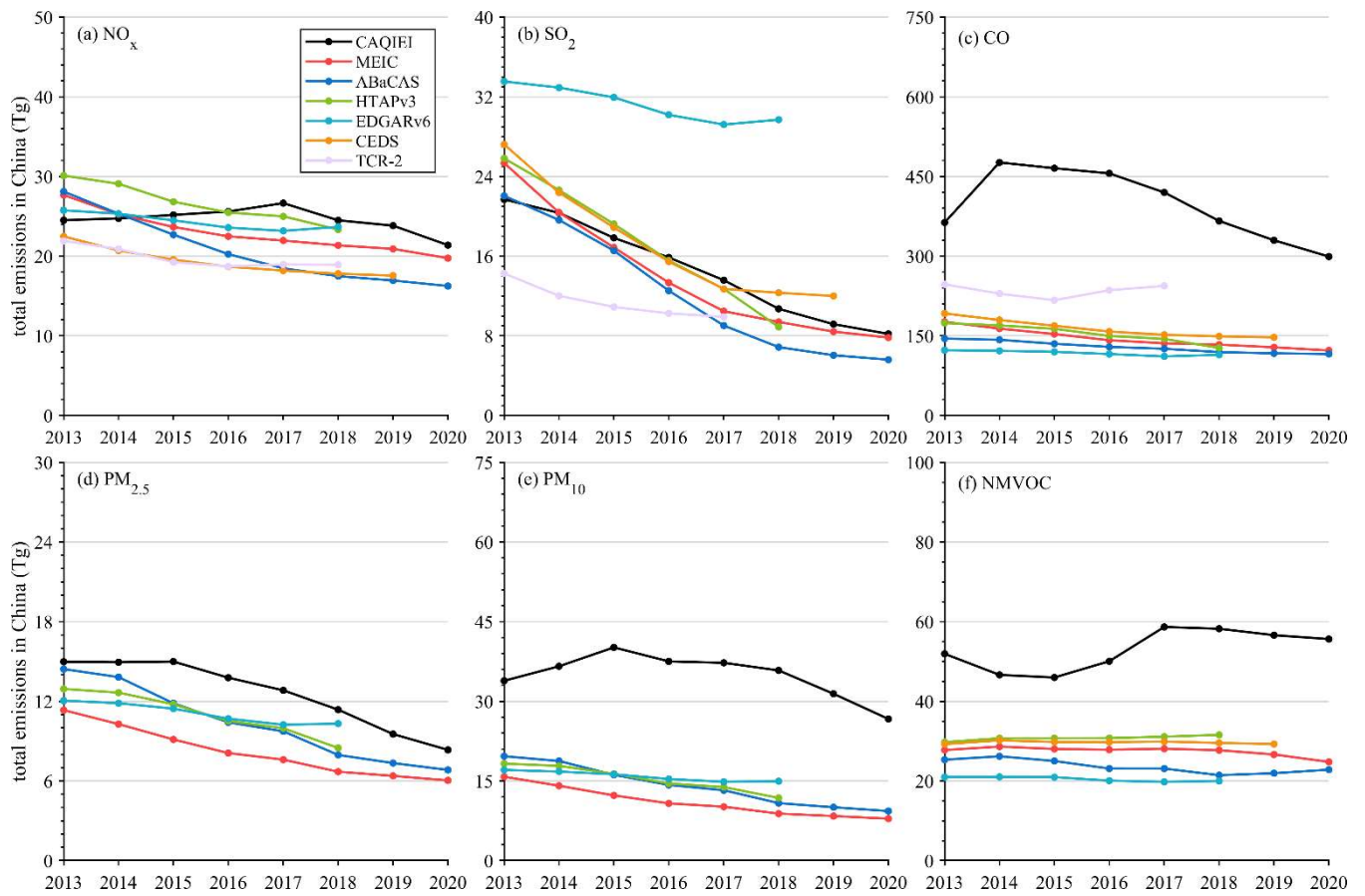
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**Figure 13: Time series of annual emissions of (a)  $\text{NO}_x$ , (b)  $\text{SO}_2$ , (c) CO, (d)  $\text{PM}_{2.5}$ , (e)  $\text{PM}_{10}$  and (f) NMVOC over China from 2013 to 2020 obtained from CAQIEI and previous inventories. Note that the natural sources were not included in the previous inventories in this figure.**

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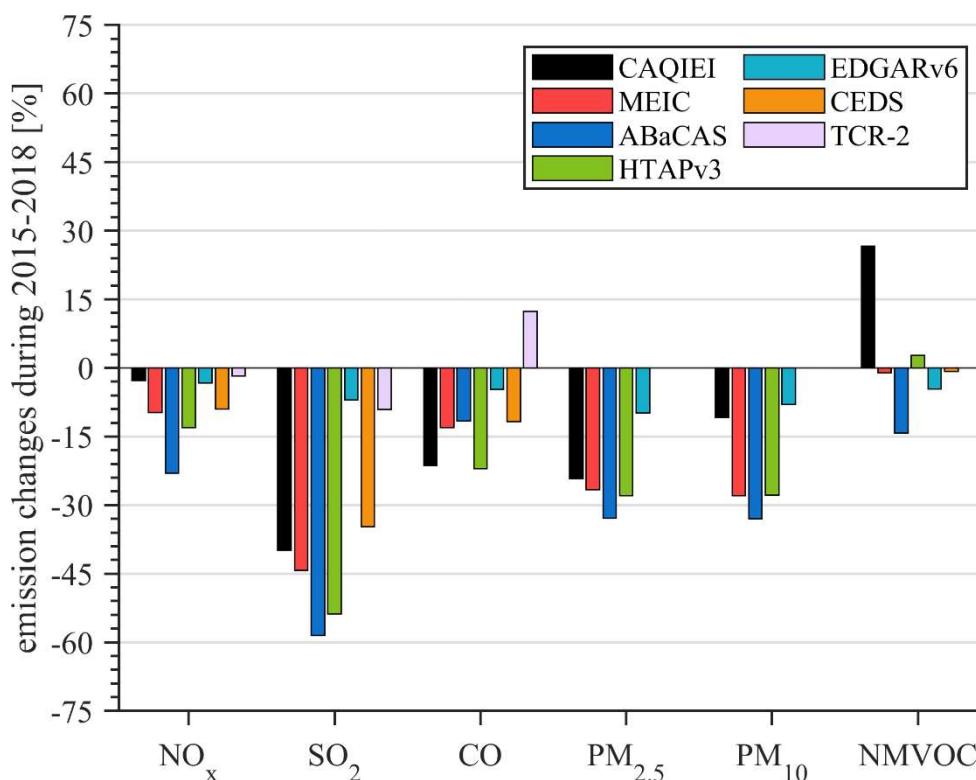
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1291 **Figure 14: Comparisons of the calculated emission changes of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub>, and (f) NMVOCs over**  
 1292 **China from 2015 to 2018 between CAQIEI and previous inventories. Note that the natural sources were not included in the**  
 1293 **calculation of the emission changes in this figure.**

1294 **Author contributions**

1295 X.T., Z.W., and J.Z. conceived and designed the project; L.K., H.W., X.T., and L.W. established the data assimilation system;  
 1296 Q.W. and L.K. performed the meteorology simulations; L.K., H.C., and J.L. conducted the ensemble simulation with the  
 1297 NAQPMS model; J.L., L.Z., W.W., B.L., Q.W., D.C. and Y.P. provided the air quality monitoring data; H.W. performed the  
 1298 quality control of the observation data; and L.K. performed the inversion estimation, generated the figures, and wrote the paper,  
 1299 with comments provided by G.R.C.

1300 **Competing interests**

1301 The authors declare no competing financial interest.

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 1305 provide us with ample computational resources to fill the requirement of the inversion of multiple years using the ensemble  
 1306 method at a high grid resolution of 15km.

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# Supplementary Material

## **Text S1: Evaluation of the meteorological simulation**

The performance of meteorological simulation is important for the inversion estimation since the meteorological parameters influence the transport, chemical and removal process of air pollutants and affect the estimation of flow-dependent background error covariance. Figure S18–S23 presents the comparisons of the simulated meteorological parameters, including zonal wind (U), meridional wind (V), temperature (T), relative humidity (RH) and precipitation, against the observations obtained from China Meteorological Administration (Figure S24). Evaluation statistics of meteorological simulation are also presented in Table S1. It shows that the WRF simulation can generally captured the main features of the different meteorological parameters over the different regions of China. The calculated correlation coefficient is 0.49–1.00 for different parameters, and the values of MB (RMSE) are -0.36–0.01 (0.3–0.52) m/s for U, -0.37–0.32 (0.32–0.80) m/s for V, -1.11–1.11 (0.6–2.17) °C for T, -11.2 to -2.59 (6.94–12.06) % for RH, and -2.05–37.35 (5.45–61.62) mm for precipitation. This suggests WRF simulation generally well reproduce the meteorological conditions for all regions of China, which is acceptable for the inversion estimates.

## **Text S2: Assessment of the influences of site differences on the emission inversions**

The emission increments at the observation sites (a posteriori minus a priori) for different species in China from 2013 to 2015 under the scenarios of fixed observation sites (blue lines) and varying observation sites (orange) were calculated to assess the influences of the site differences on the emission inversions (Fig. S2). In the fixed-site scenario, it is assumed that the number of observation sites remains constant at the 2013 level while in the varying-site scenario, the number of observation sites increases over time. The differences in emission increments between these two scenarios are used to analyze the impact of changes in the observation coverage on the emission inversions. Please note that, to simplify calculations, we only computed the emission increments at the locations of observation sites. Therefore, they may not be equal to the emission increments calculated for the entire grid as reported in the paper. However, they are still useful indicators for the effects of emission inversion. In addition, since we did not consider the temporal variation in the a priori emissions, the changes of the emission increments can be used to approximate the temporal variations of the a posterior emissions. It can be clearly seen that that there are obvious differences in the emission increments between the two scenarios. The emission increment is larger in the varying-site scenario than that in the fixed-site scenario for all species due to the increases of observation sites. Moreover, as indicated in Fig. S2, the changes of observation sites were shown to significantly affect the estimation of the emission trend in 2013 and 2014. Most of species showed decreasing trends in their inversed emission under the fixed-site scenario. However, under the varying-site scenario, the decreasing trends were smaller

for PM<sub>2.5</sub>, NO<sub>x</sub> and NMVOC, and the emissions of PM<sub>10</sub> and CO even showed increasing trends. This is due to that the emission increments were positive over most of observation sites for these species as demonstrated in Fig.3. Thus, the increases of observation site would lead to increases of positive emission increments and higher a posteriori emissions, which may counteract the decreasing trends or even lead to an opposite trend. These results provide the evidences that the increasing trends in the total emissions of PM<sub>10</sub> and CO from 2013 to 2015 seen in Fig. 6 and Fig. 7 are highly likely to be a spurious trend caused by the changes of observation coverage. The weak emission changes in PM<sub>2.5</sub> and NO<sub>x</sub> (Fig. 6 and Fig. 7) may also be related to the changes in the number of observation sites. The SO<sub>2</sub> emission is an except that its calculated trend is larger under the varying-site scenario than that under the fixed-site scenario. This is because that the emission increment for the SO<sub>2</sub> is generally negative over the most sites, thus the increased observation sites would lead to larger decreasing trend in the inversed emissions of SO<sub>2</sub>. To date, these results highlighted the significant influences of the site differences on the estimated emissions and their trends. Therefore, we recommend not to use the emission in 2013 and 2014 when analyze the trends of the emissions.

**Table S1** **Text S3: Comparisons of model performance driven by CAQIEI with that driven by more recent bottom-up emission inventories**

To obtain a better understanding of the accuracy of our inverse emission inventory, we conducted a one-year simulation of air pollution in China for year 2020 with more recent bottom-up emission inventories and compared its performance with that driven by the CAQIEI. The used bottom-up inventories in this simulation case includes the HTAPv3 (Crippa et al., 2023) inventory for the anthropogenic emissions outside China with a base year of 2018; the MEIC inventory for the anthropogenic emissions over China with a base year of 2020; the CAMS emission inventory (<https://ads.atmosphere.copernicus.eu/cdsapp#!/dataset/cams-global-emission-inventories?tab=overview>, last access: 19 June 2024) for the biogenic, soil and oceanic emissions; and the Global Fire Assimilation System (GFAS) (Kaiser et al., 2012) for the biomass burning emissions. Note that since the MEIC emission inventory does not include the ship, air and waste emissions. Emissions from these sectors over China were provided by the HTAPv3 emission inventory. For clarity, in following content, we name this simulation case as the MEIC-HTAPv3 based on the anthropogenic emission inventory used.

Figure S25 shows the time series of hourly concentrations of different air pollutants in China obtained from observation and simulation driven by the CAQIEI and more recent bottom-up inventories. Comparisons of the evaluation statistics of these two simulation scenarios are also presented in Table S3. It shows that updating the bottom-up emission inventories to a more recent year

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59 does improve the model performance compared to the outdated a priori emission inventory (Table 2), suggesting that the bottom-  
60 up emission inventory has to some extent captured the changes of air pollutant emissions in China. It is also encouraging to find  
61 that the model performance driven by CAQIEI and MEIC-HTAPv3 is similar for the PM<sub>2.5</sub>, PM<sub>10</sub>, and SO<sub>2</sub> over the NCP, NE, SE  
62 and SW regions, both significantly improved from the a priori emission inventory. This suggest that both the top-down and recent  
63 bottom-up emission inventories have good performance in capturing the emission changes of these species over these regions and  
64 they yield consistent estimations. However, the model simulation driven by MEIC-HTAPv3 still have negative biases in the CO  
65 concentrations possibly due to the underestimations of CO emissions as we illustrated in Sect.4.3.1.3. Similarly, due to the errors in  
66 the dust emission, there are negative biases in the simulated PM<sub>2.5</sub> and PM<sub>10</sub> concentrations over the western China driven by MEIC-  
67 HTAPv3. On the contrary, the simulated NO<sub>2</sub> concentrations in MEIC-HTAPv3 are higher than the observations over the NCP, NE  
68 and SE regions, which also partly contributes to the underestimated O<sub>3</sub> concentrations over these regions. The CAQIEI generally  
69 achieves better performance in simulating the air pollutant concentrations in China as indicated by higher values of correlation  
70 coefficient and lower values of bias and root mean square of error in the model simulation driven by CAQIEI than that driven by  
71 MEIC-HTAPv3 (Table S3).



**Table S1: Evaluation statistics for the meteorology simulation**

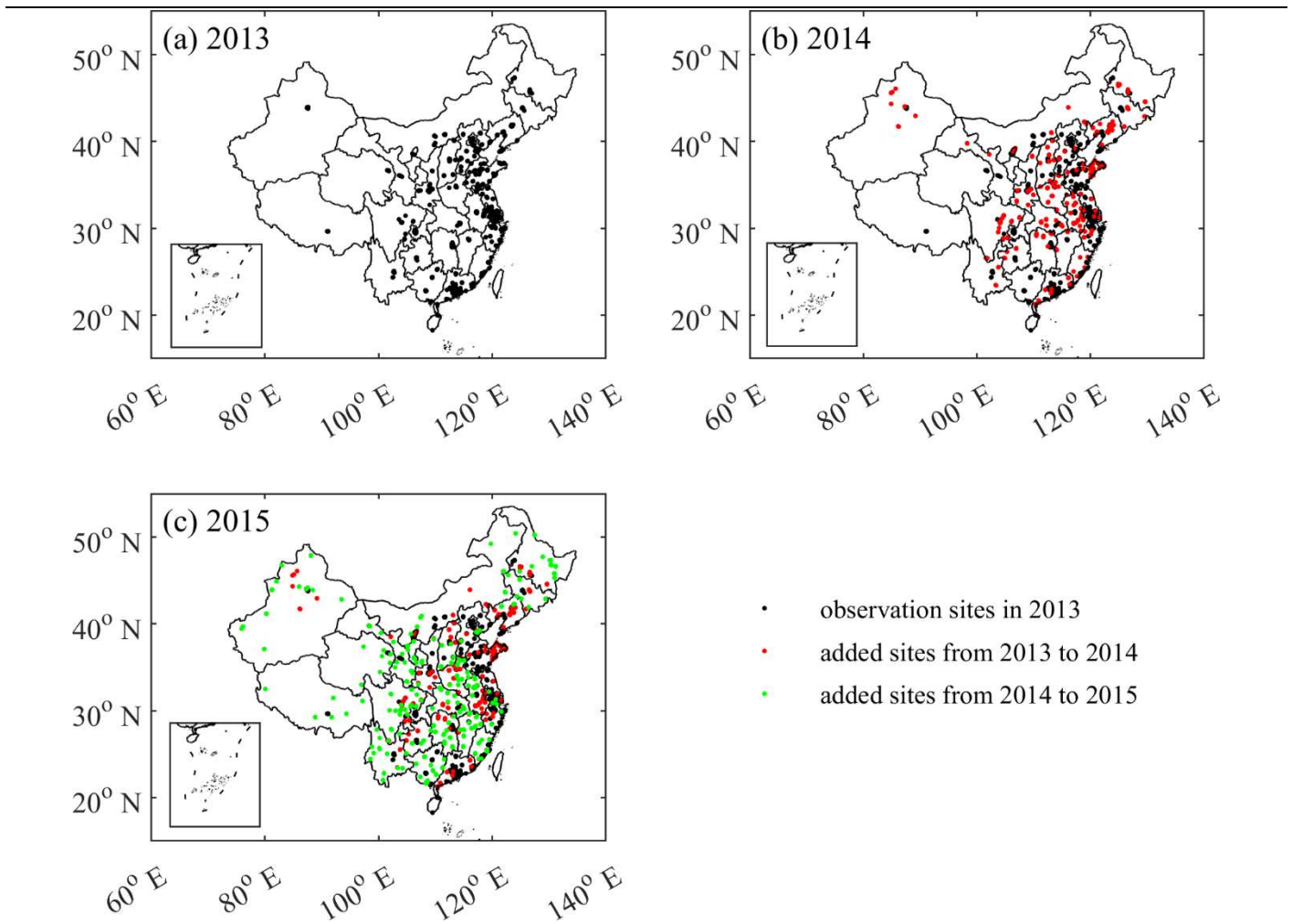
Region	U (m/s)			V (m/s)			T (°C)			RH (%)			Precipitation (mm/month)		
	R	MB	RMSE	R	MB	RMSE	R	MB	RMSE	R	MB	RMSE	R	MB	RMSE
<u>NCP</u>	<u>0.95</u>	<u>0.01</u>	<u>0.30</u>	<u>0.95</u>	<u>-0.02</u>	<u>0.49</u>	<u>1.00</u>	<u>-0.42</u>	<u>0.84</u>	<u>0.95</u>	<u>-11.24</u>	<u>11.66</u>	<u>0.95</u>	<u>3.74</u>	<u>18.56</u>
<u>NE</u>	<u>0.94</u>	<u>0.37</u>	<u>0.51</u>	<u>0.89</u>	<u>-0.08</u>	<u>0.49</u>	<u>0.99</u>	<u>-1.11</u>	<u>2.17</u>	<u>0.77</u>	<u>-2.59</u>	<u>7.18</u>	<u>0.97</u>	<u>12.09</u>	<u>19.76</u>
<u>SE</u>	<u>0.84</u>	<u>-0.27</u>	<u>0.37</u>	<u>0.98</u>	<u>-0.37</u>	<u>0.80</u>	<u>1.00</u>	<u>-0.40</u>	<u>0.60</u>	<u>0.88</u>	<u>-7.00</u>	<u>7.58</u>	<u>0.94</u>	<u>37.35</u>	<u>61.62</u>
<u>SW</u>	<u>0.63</u>	<u>-0.44</u>	<u>0.52</u>	<u>0.69</u>	<u>0.04</u>	<u>0.37</u>	<u>0.99</u>	<u>1.11</u>	<u>1.27</u>	<u>0.87</u>	<u>-5.84</u>	<u>6.94</u>	<u>0.92</u>	<u>16.85</u>	<u>40.18</u>
<u>NW</u>	<u>0.49</u>	<u>-0.36</u>	<u>0.51</u>	<u>0.58</u>	<u>0.32</u>	<u>0.43</u>	<u>0.99</u>	<u>0.83</u>	<u>1.91</u>	<u>0.79</u>	<u>-9.49</u>	<u>12.06</u>	<u>0.51</u>	<u>-2.05</u>	<u>5.45</u>
<u>CENTRAL</u>	<u>0.95</u>	<u>0.10</u>	<u>0.41</u>	<u>0.70</u>	<u>-0.08</u>	<u>0.32</u>	<u>1.00</u>	<u>-0.27</u>	<u>0.93</u>	<u>0.85</u>	<u>-8.59</u>	<u>10.30</u>	<u>0.97</u>	<u>4.64</u>	<u>10.87</u>

**Table S2 The average mean (standard deviation) of the calculated factor for the inflation of the ensemble member over different regions of China for different species**

	NCP	NE	SE	SW	NW	Central
PM <sub>2.5</sub>	1.0 (0.2)	1.7 (1.6)	1.0 (0.0)	6.8 (8.5)	3.1 (3.8)	3.9 (3.9)
PM <sub>10</sub>	1.4 (0.7)	7.2 (8.0)	2.4 (0.8)	78.1 (108.2)	26.3 (36.5)	36.0 (49.0)
SO <sub>2</sub>	1.4 (0.7)	4.1 (3.2)	2.3 (0.8)	176.1 (254.6)	7.8 (6.5)	58.6 (72.5)
NO <sub>x</sub>	1.0 (0.1)	1.7 (0.7)	1.2 (0.3)	8.1 (5.3)	2.8 (1.3)	5.4 (4.1)
CO	1.0 (0.1)	2.8 (2.3)	1.4 (0.4)	18.8 (16.8)	6.8 (6.9)	8.6 (10.0)
NMVOG	1.4 (0.6)	4.5 (4.4)	1.6 (0.5)	8.1 (8.6)	6.5 (5.8)	8.1 (10.1)

**Table S3: Evaluation statistics of the model simulation driven by CAQIEI (outside brackets) and more recent bottom-up inventories (inside brackets) in 2020**

	<u>PM<sub>2.5</sub></u> ( $\mu\text{g}/\text{m}^3$ )	<u>PM<sub>10</sub></u> ( $\mu\text{g}/\text{m}^3$ )	<u>SO<sub>2</sub></u> ( $\mu\text{g}/\text{m}^3$ )	<u>NO<sub>2</sub></u> ( $\mu\text{g}/\text{m}^3$ )	<u>CO</u> ( $\text{mg}/\text{m}^3$ )	<u>O<sub>3</sub></u> ( $\mu\text{g}/\text{m}^3$ )
<u>R</u>	<u>0.77 (0.53)</u>	<u>0.73 (0.44)</u>	<u>0.37 (0.19)</u>	<u>0.69 (0.45)</u>	<u>0.67 (0.40)</u>	<u>0.75 (0.48)</u>
<u>MB</u>	<u>3.6 (5.3)</u>	<u>-0.3 (-14.9)</u>	<u>0.3 (0.7)</u>	<u>-0.9 (6.7)</u>	<u>-0.06 (-0.4)</u>	<u>6.3 (-13.7)</u>
<u>NMB (%)</u>	<u>10.5 (15.8)</u>	<u>-0.5 (-25.9)</u>	<u>2.6 (7.6)</u>	<u>-3.4 (26.2)</u>	<u>-8.9 (-52.7)</u>	<u>10.2 (-22.1)</u>
<u>RMSE</u>	<u>24.6 (34.2)</u>	<u>37.4 (49.1)</u>	<u>10.9 (13.6)</u>	<u>15.9 (25.1)</u>	<u>0.4 (0.6)</u>	<u>30.3 (42.3)</u>



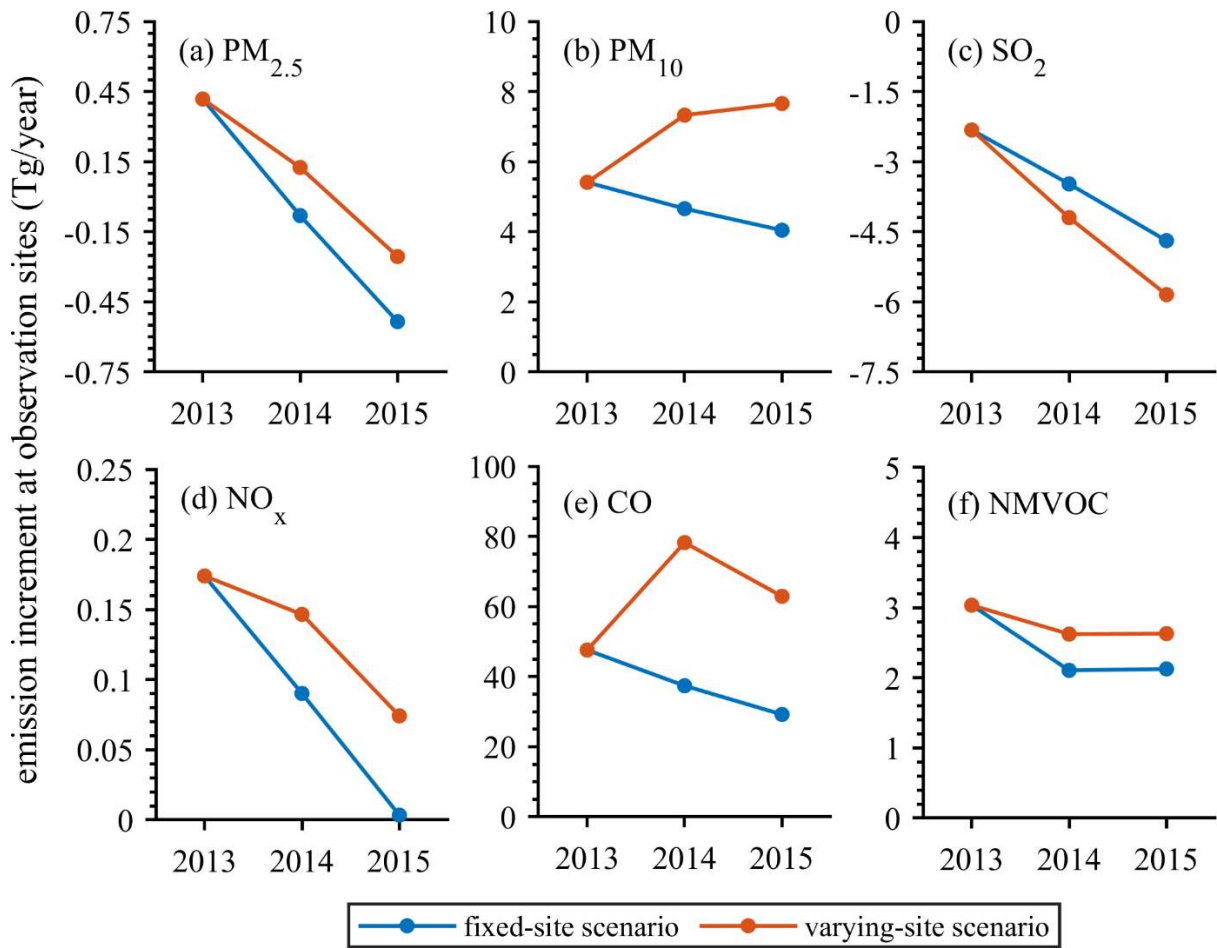
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Figure S1: Spatial distributions of observation sites in (a) 2013, (b) 2014 and (c) 2015. The observation sites in 2013 were marked as black dots, while the added observation sites from 2013 to 2014 and those from 2014 to 2015 were marked as red and green dots, respectively.



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 115 **Figure S2: The calculated total emission increments at the observation sites for different species under the fixed-site scenario and varying-**  
 116 **site scenario.**

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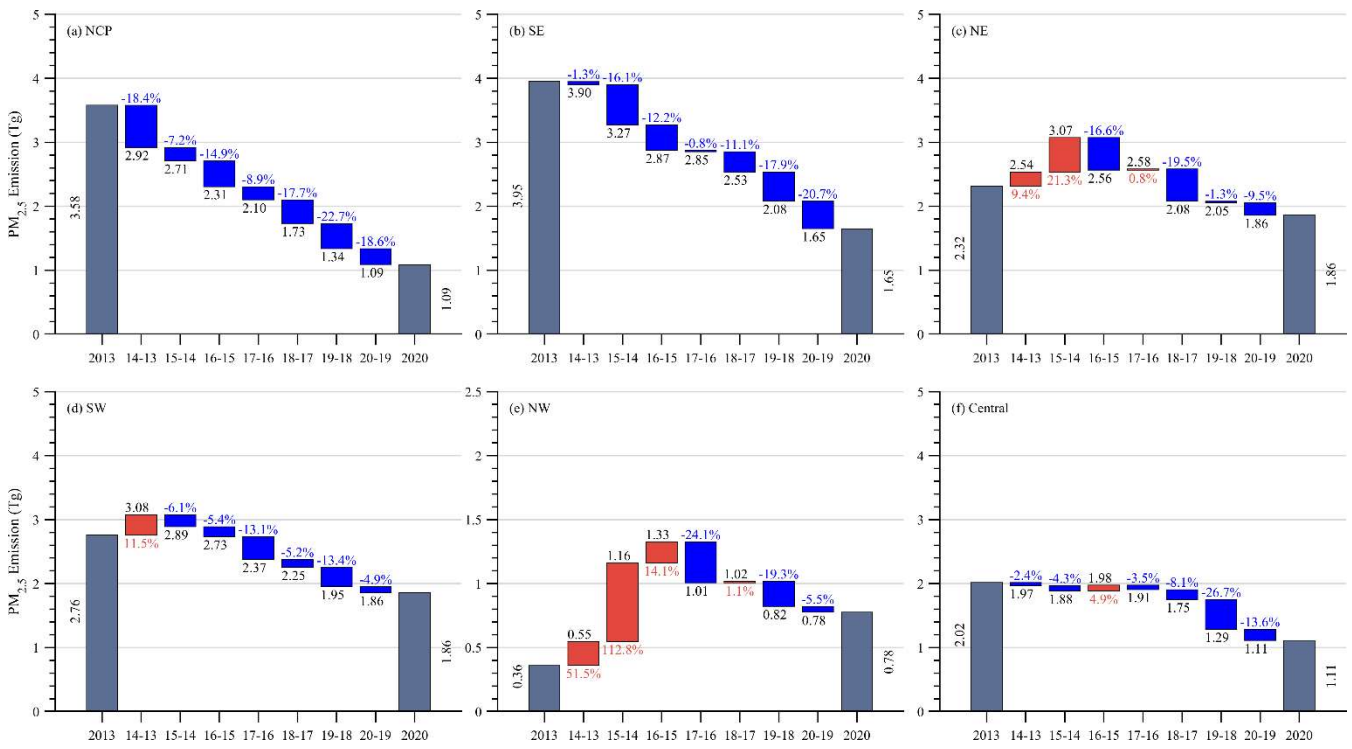


Figure S3: Emission changes of PM<sub>2.5</sub> from 2013 to 2020 over different regions of China obtained from CAQIET.

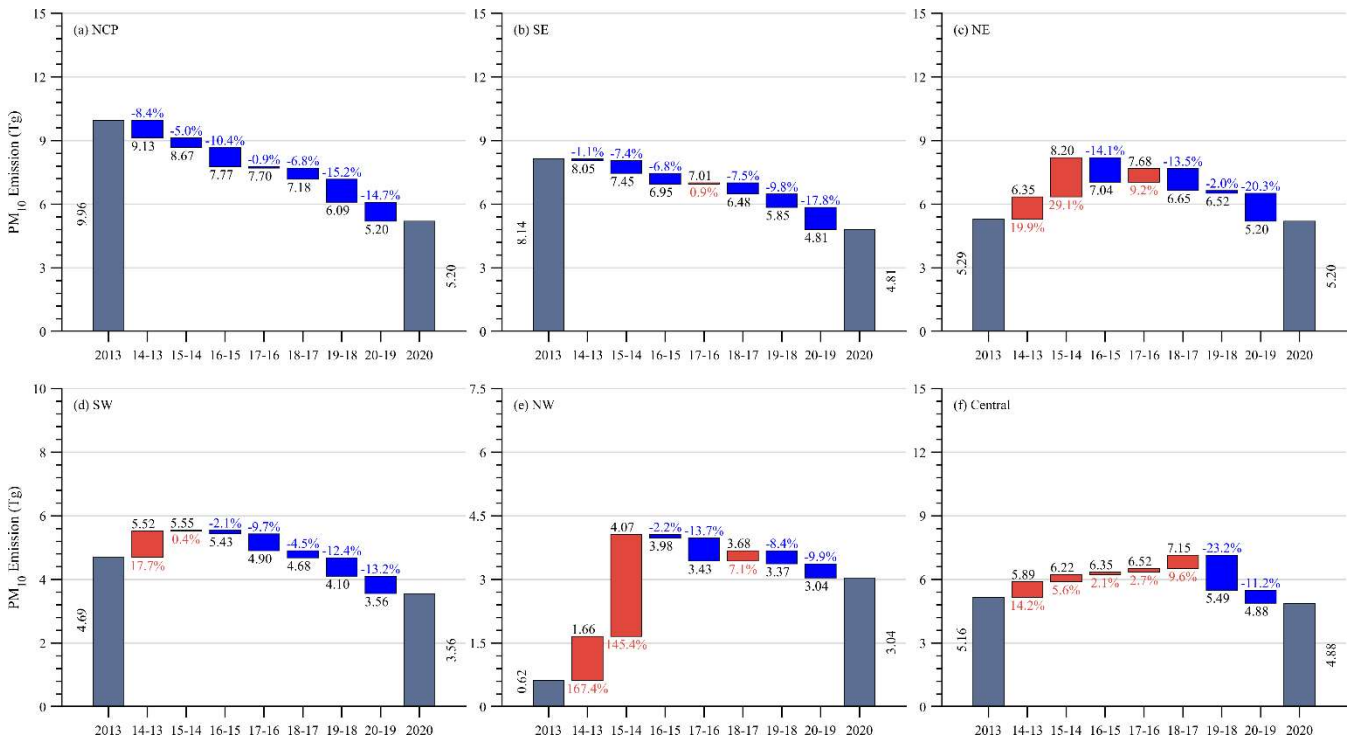


Figure S4: Same as Fig. S3 but for PM<sub>10</sub>.

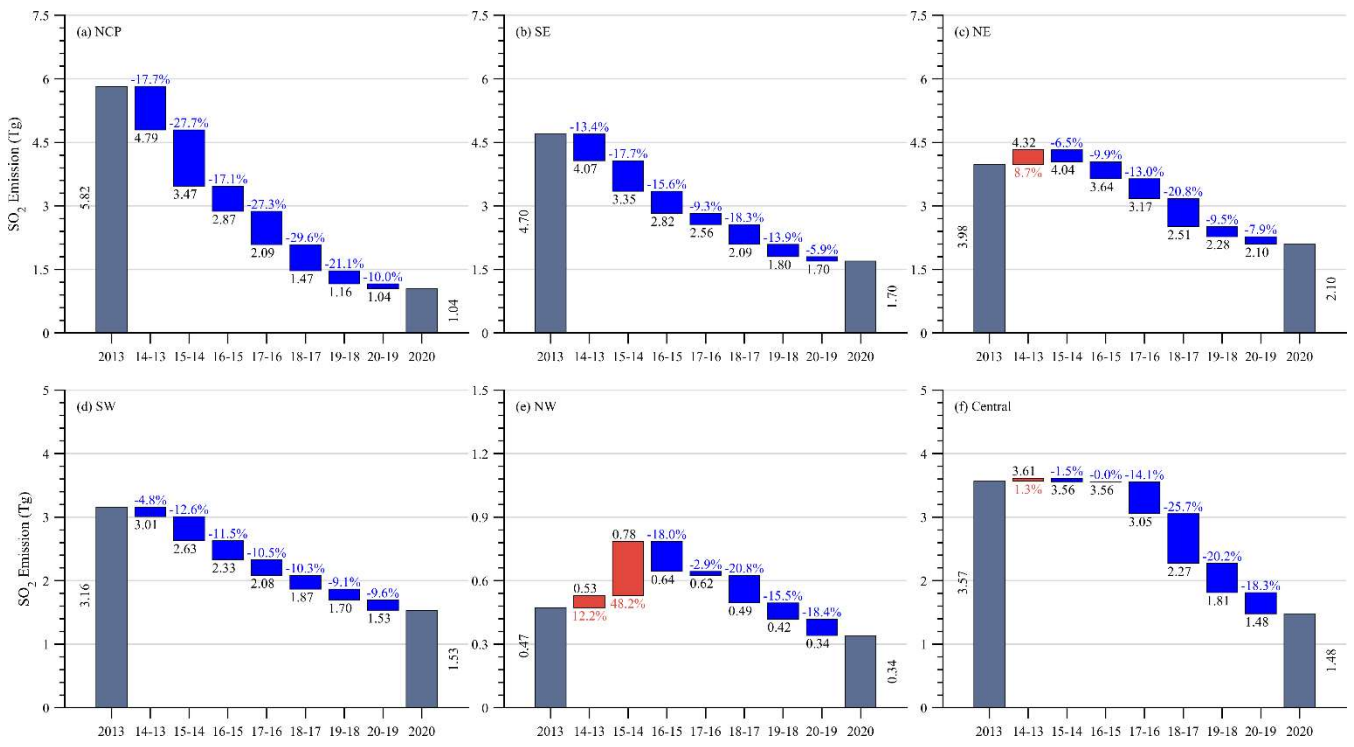


Figure S5: Same as Fig. S3 but for SO<sub>2</sub>.

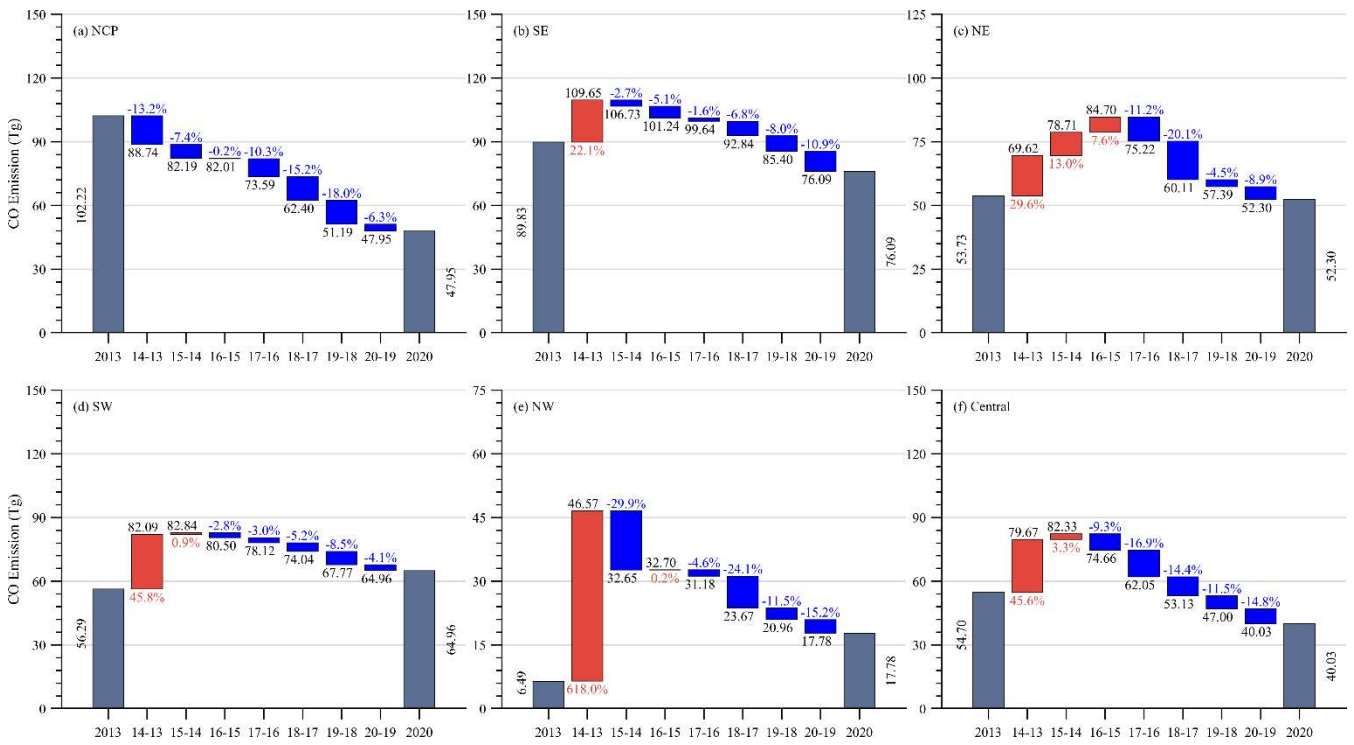


Figure S6: Same as Fig. S3 but for CO.

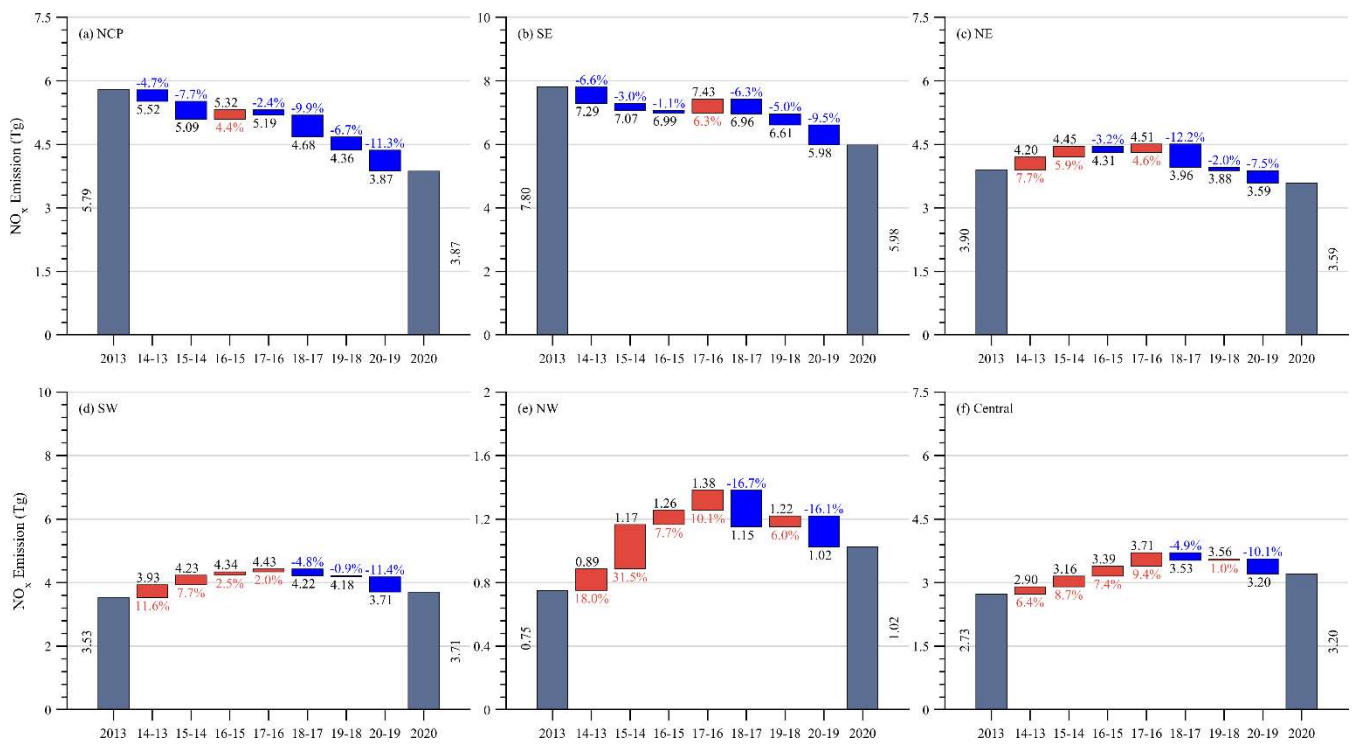


Figure S7: Same as Fig. S3 but for NO<sub>x</sub>.

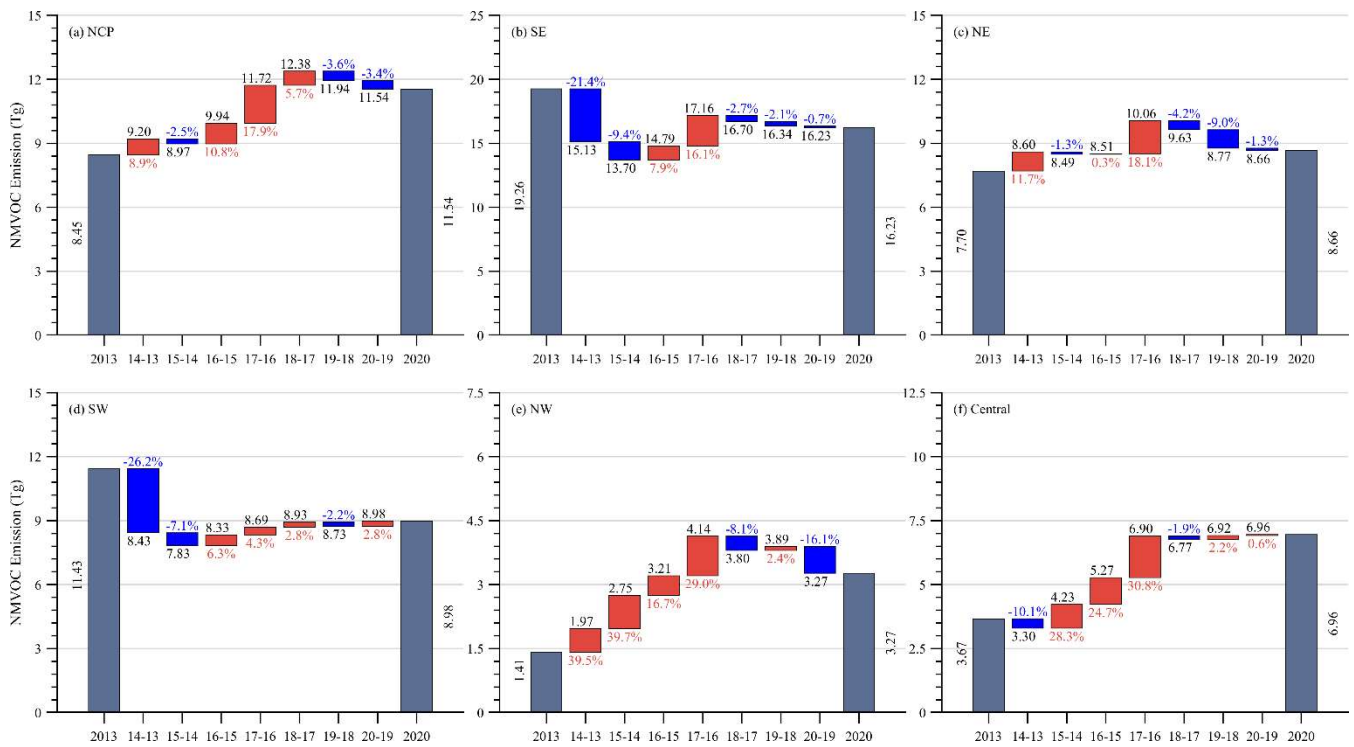
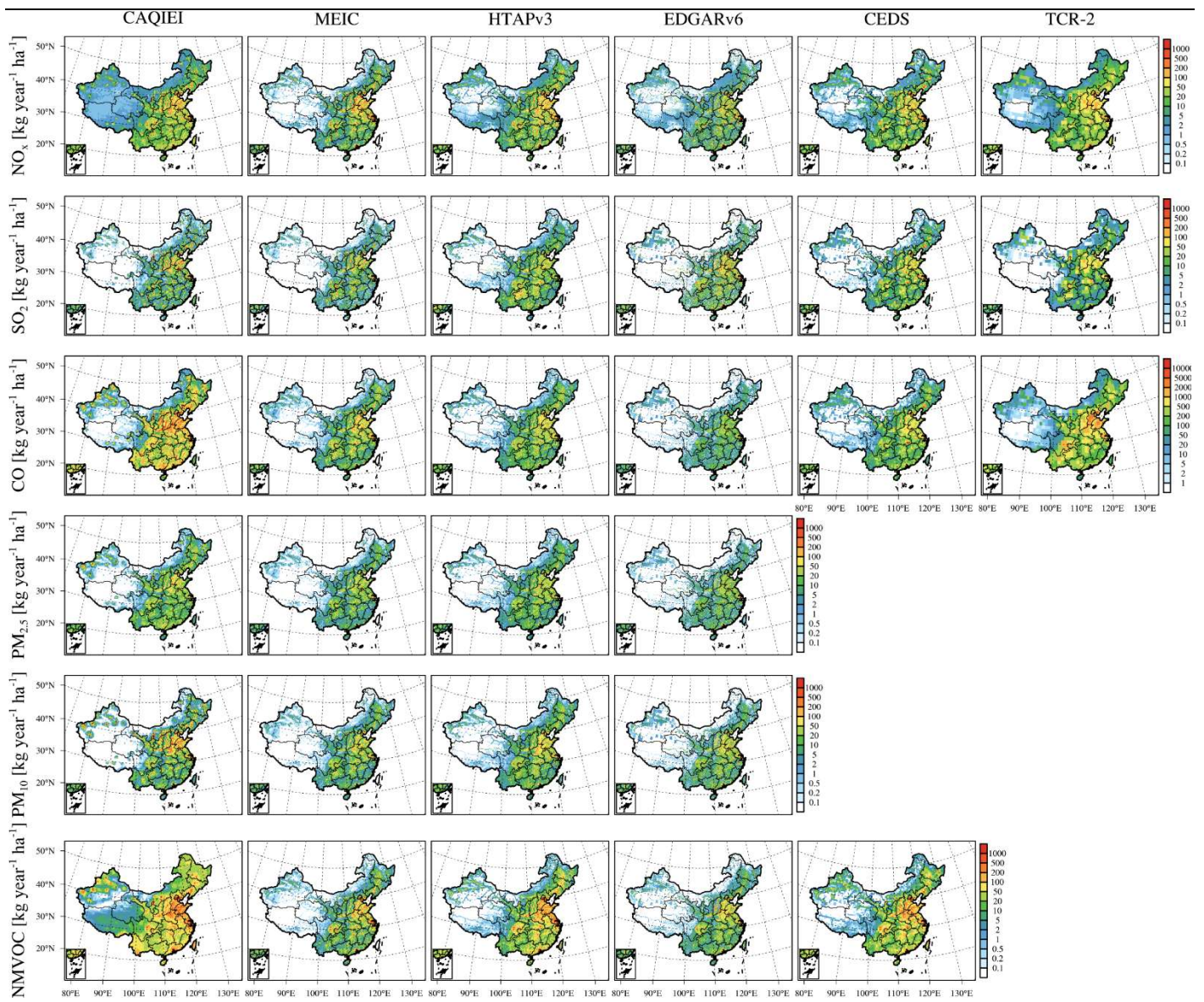


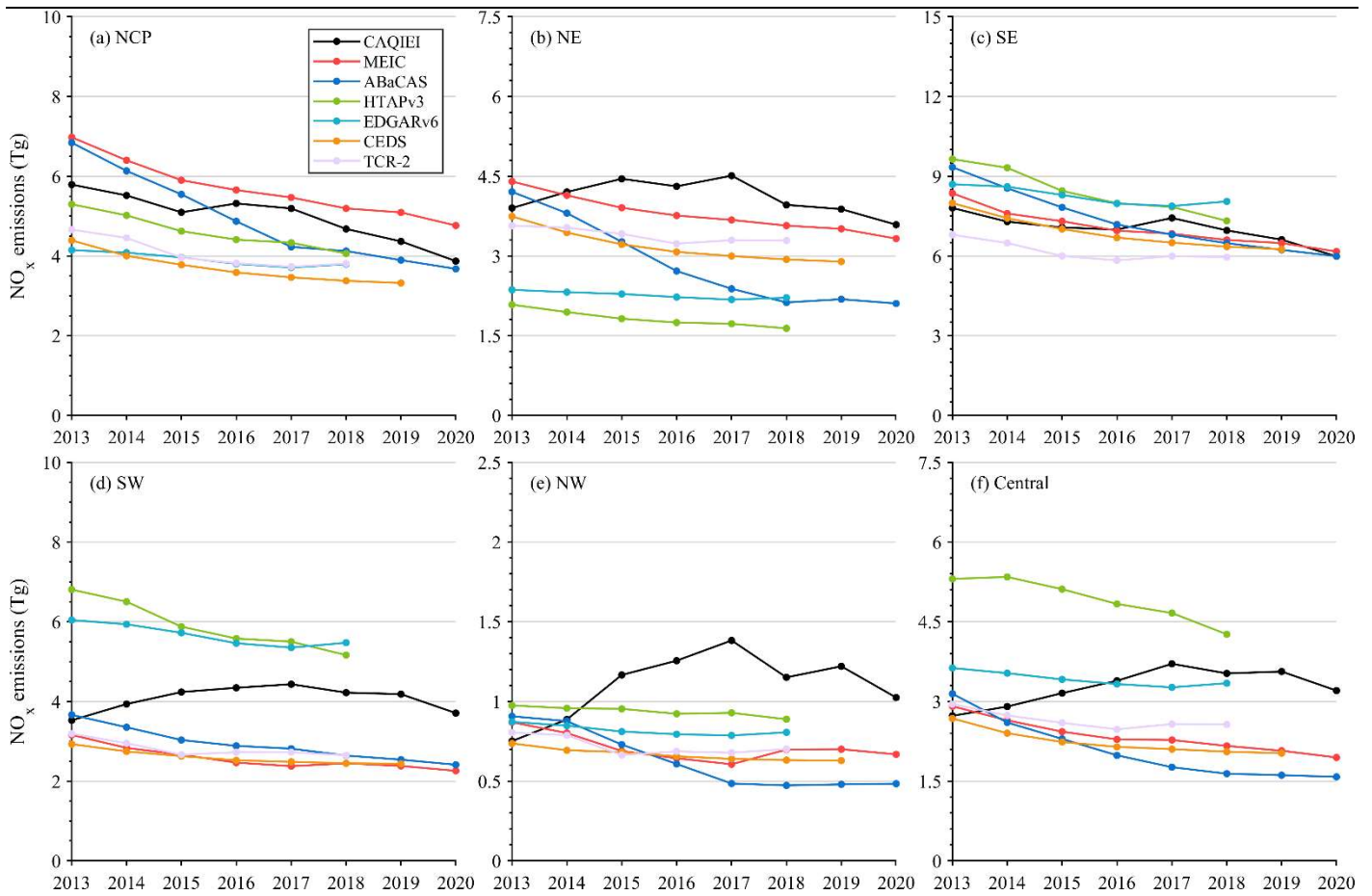
Figure S8: Same as Fig. S3 but for NMVOC.



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**Figure S9: Spatial distributions of the averaged emissions of different air pollutants in China during 2015–2018 obtained from CAQIEI, MEIC, HTAPv3, EDGARv6, CEDS and TCR-2. Note the due to absence of gridded products of the ABaCAS inventory, we did not provide its spatial distributions. Also, the natural sources were not added to the previous emission inventories in this figure because of the different spatial resolutions among these inventories.**





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 150 **Figure S10: Time series of annual NO<sub>x</sub> emissions over of different regions of China: (a) NCP, (b) NE, (c) SE, (d) SW, (e) NW and (f)**  
 151 **Central from 2013 to 2020 obtained from CAQIEI and previous inventories. Note that the natural sources were not included in the**  
 152 **previous inventories in this figure.**

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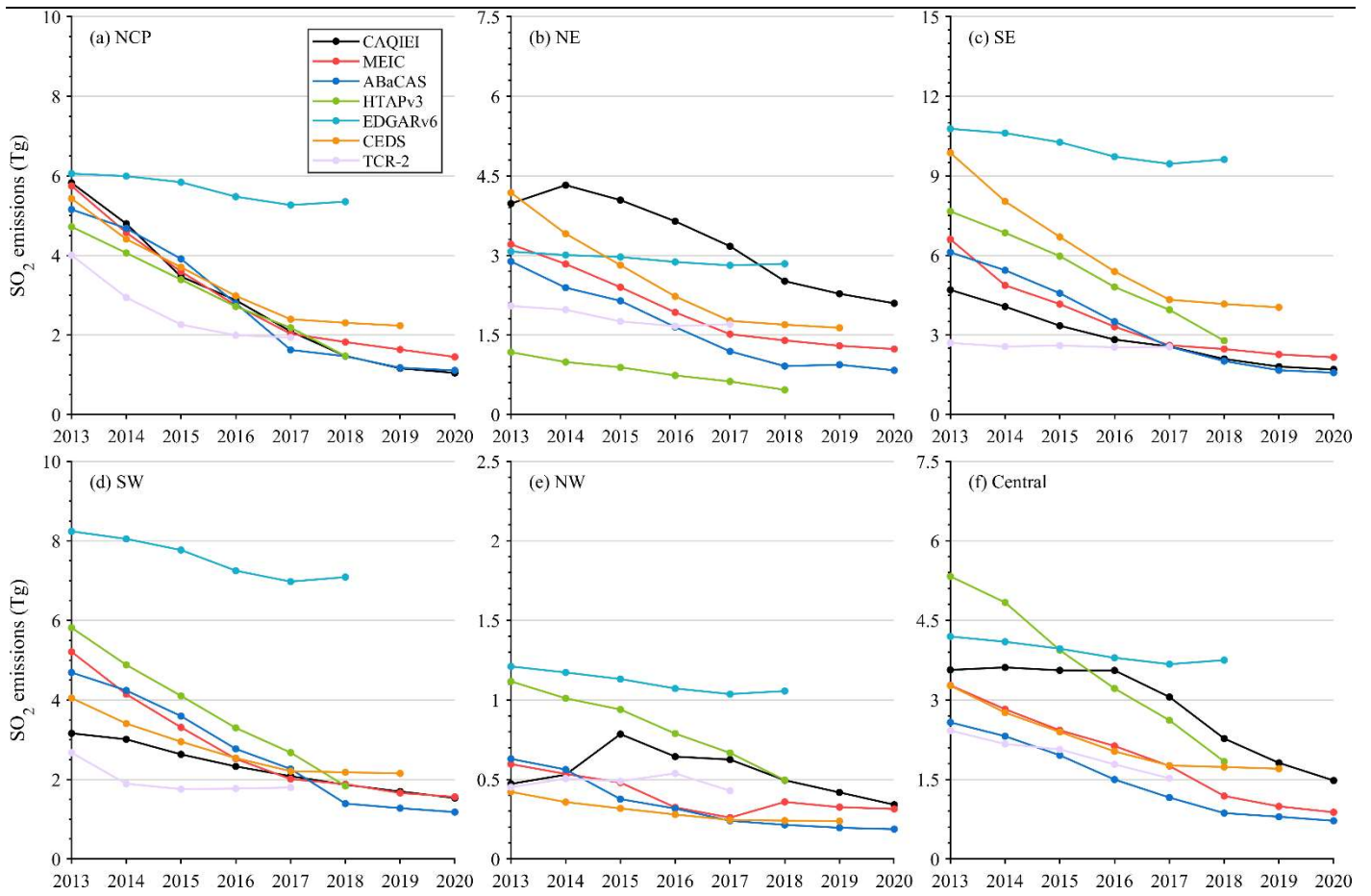


Figure S11: Same as Fig. S10 but for SO<sub>2</sub>.

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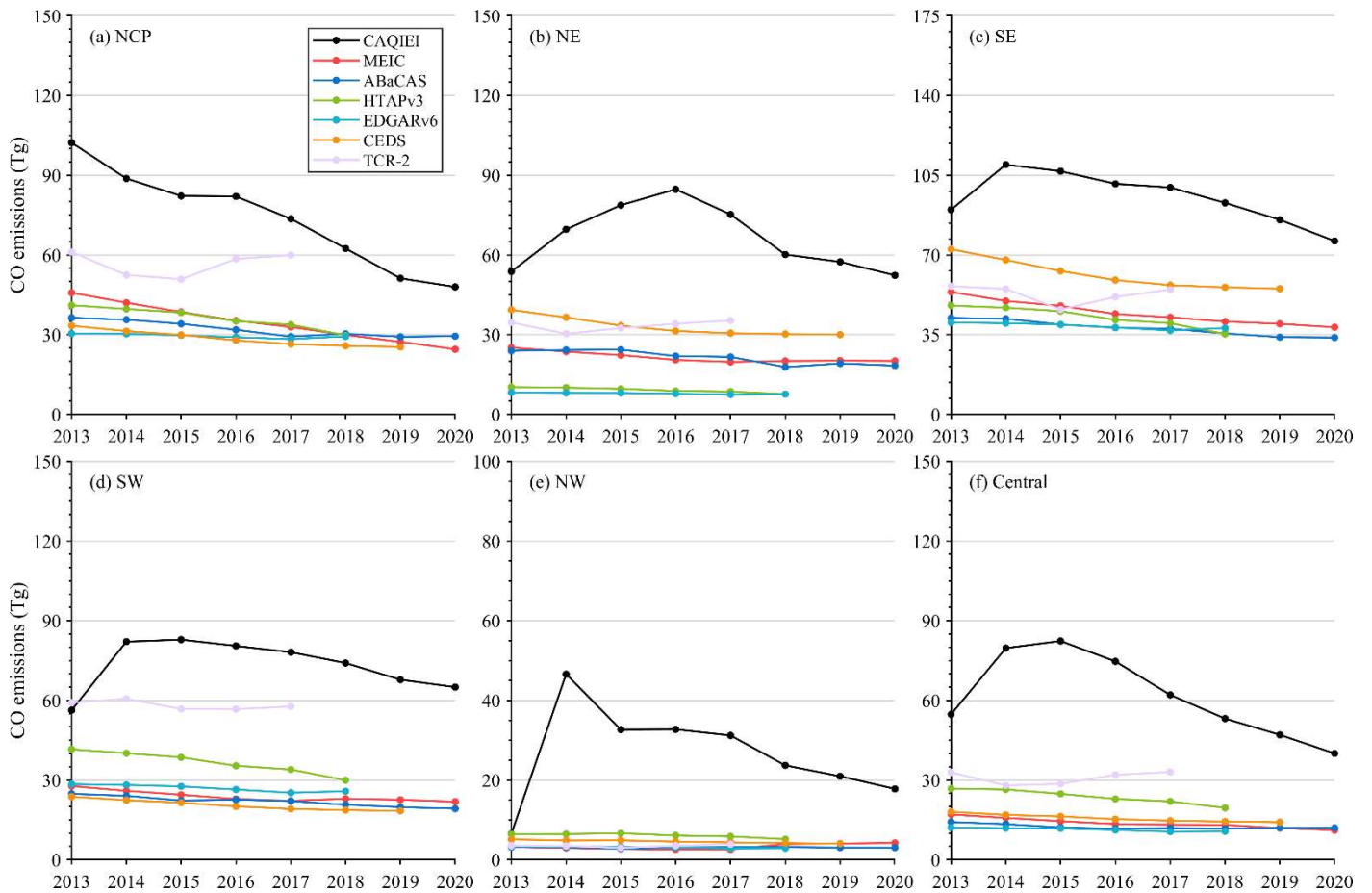


Figure S12: Same as Fig. S10 but for CO.

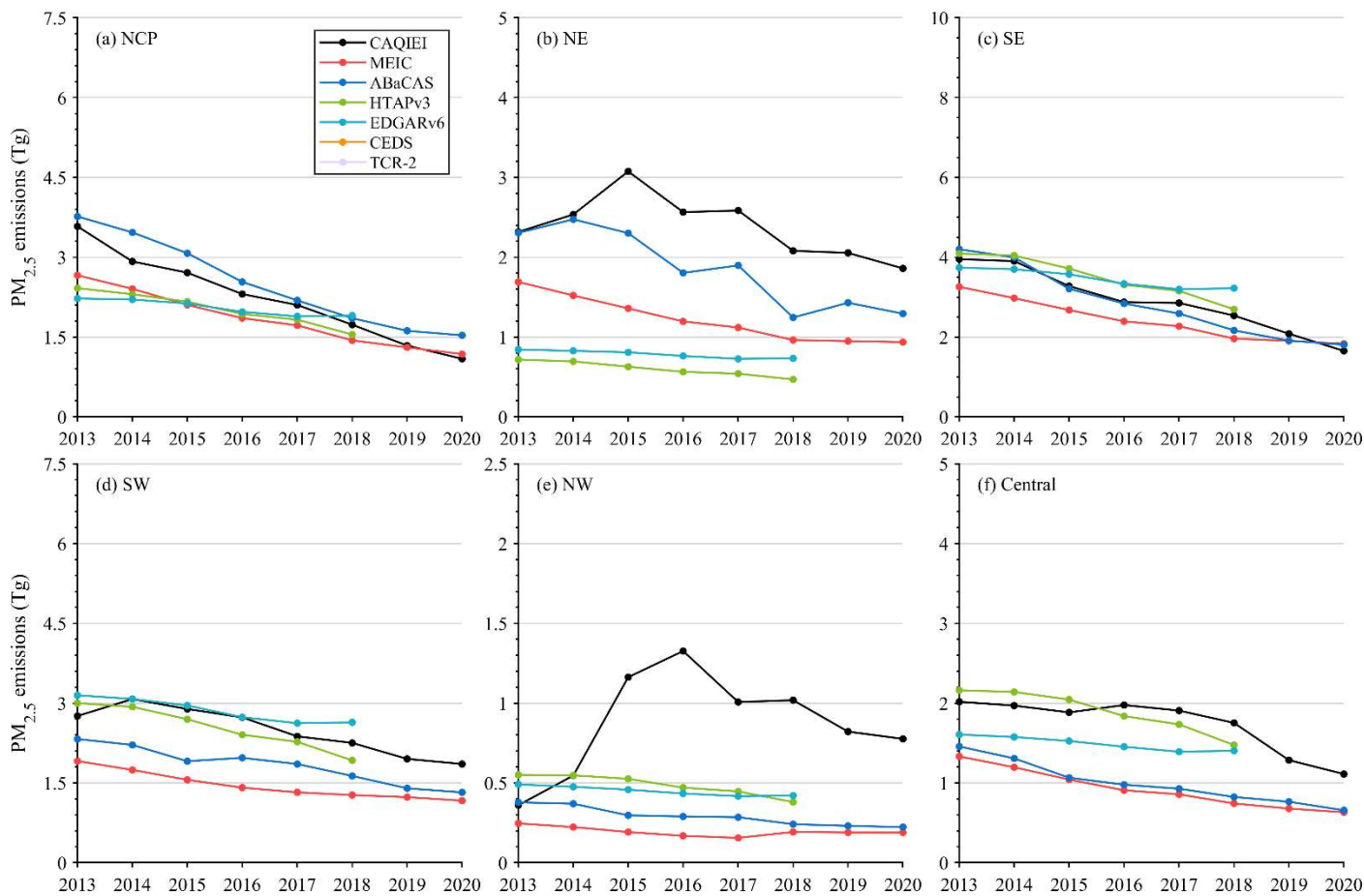


Figure S13: Same as Fig. S10 but for PM<sub>2.5</sub>.

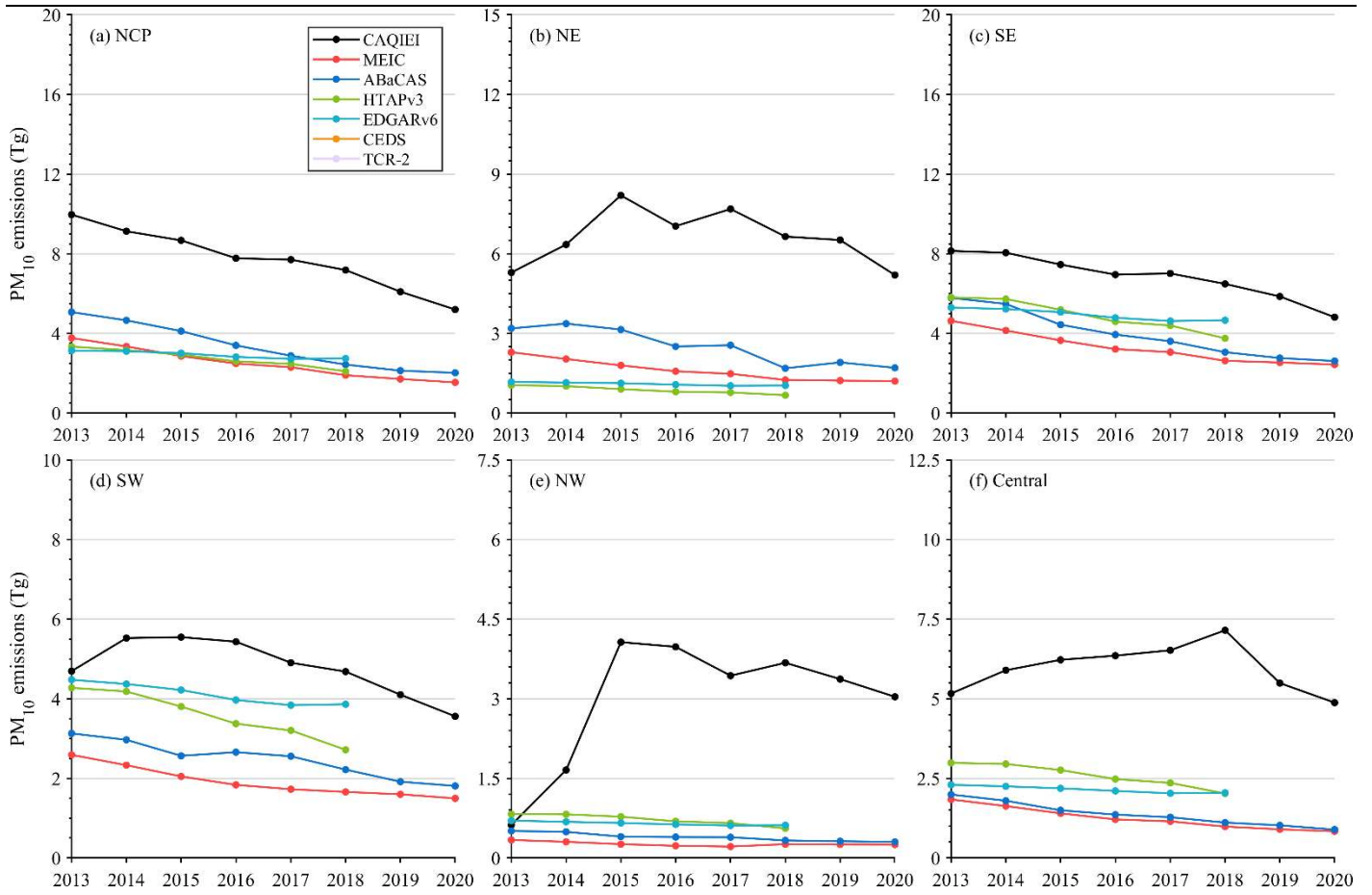


Figure S14: Same as Fig. S10 but for PM<sub>10</sub>

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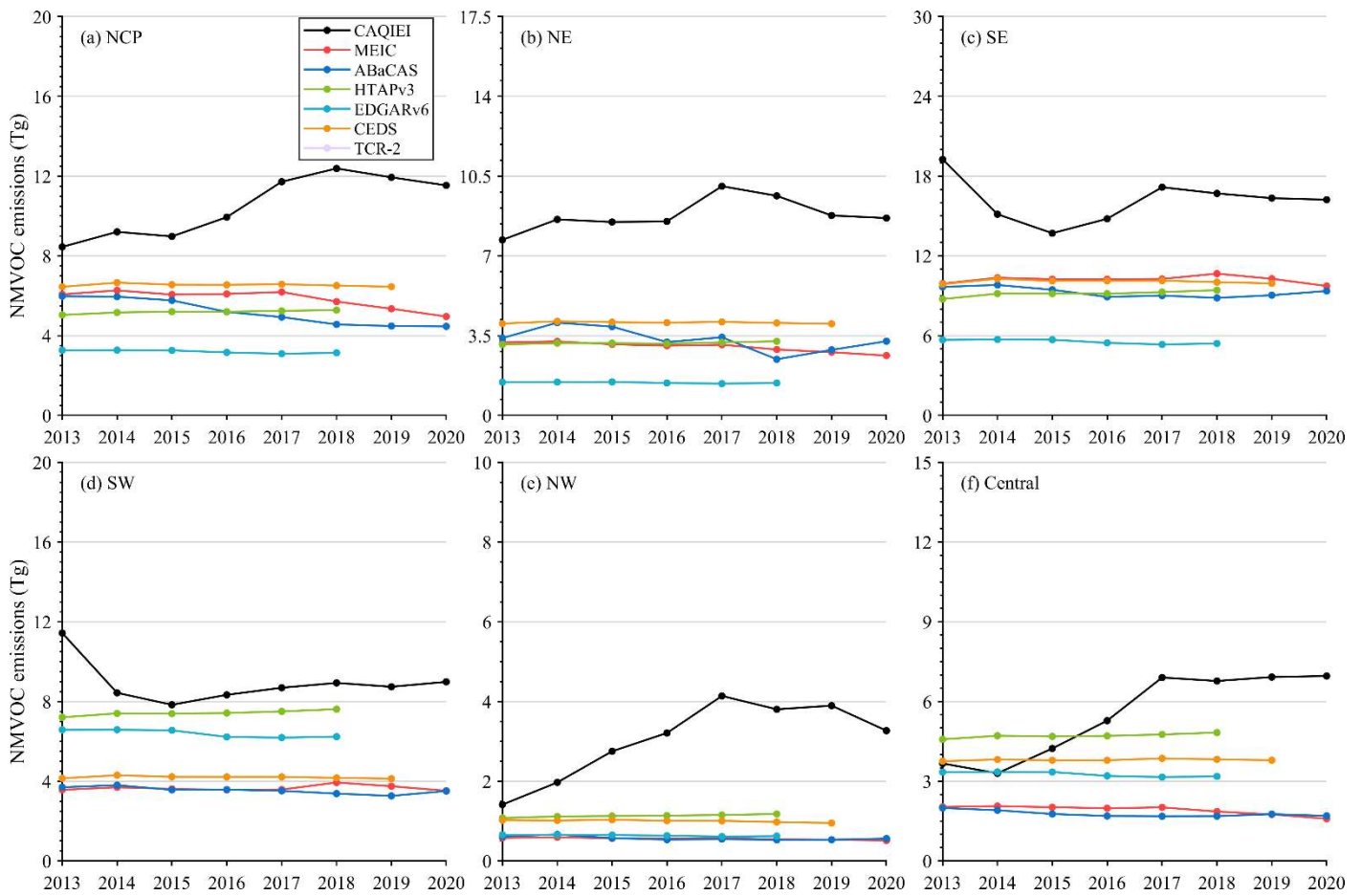
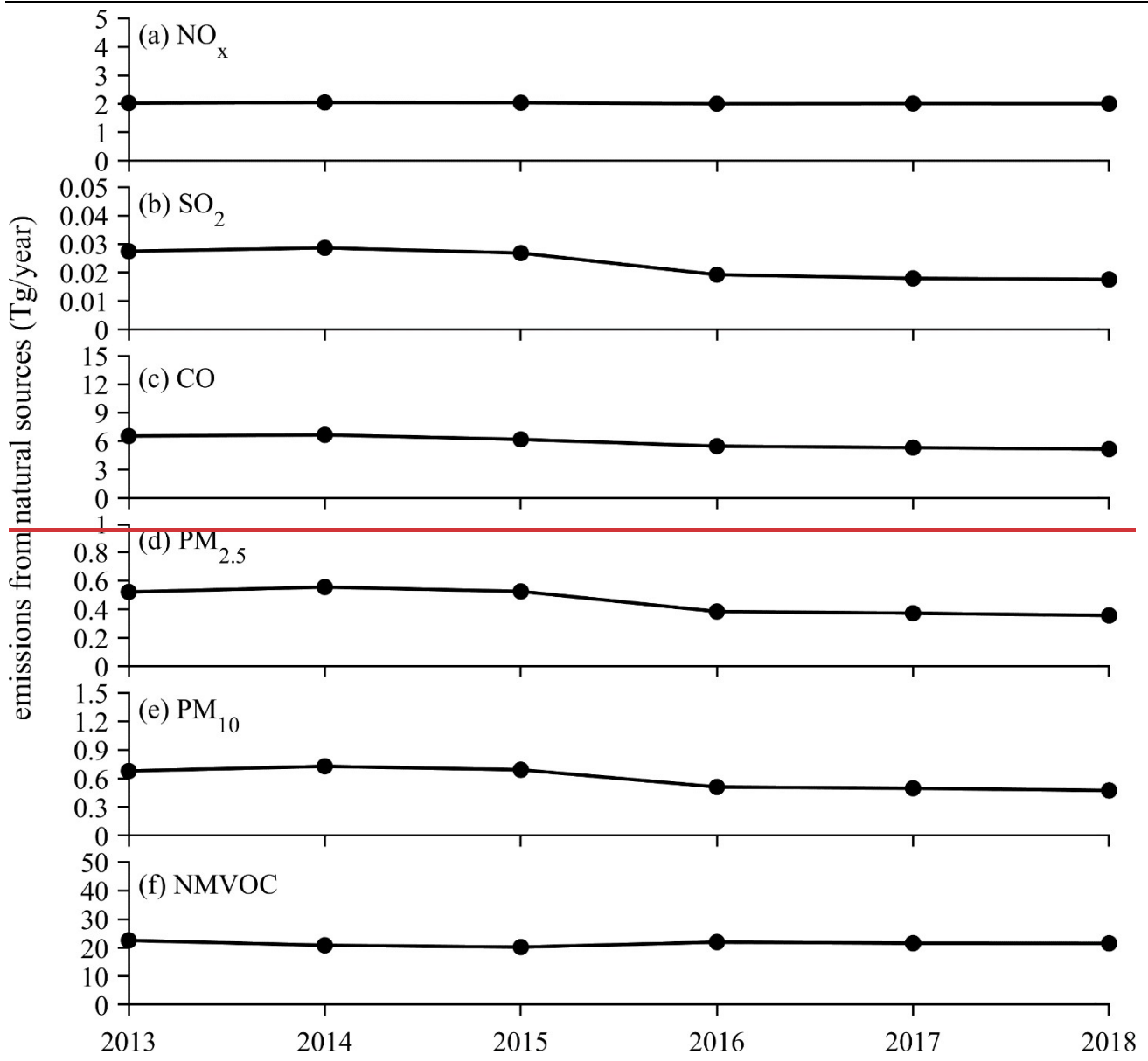


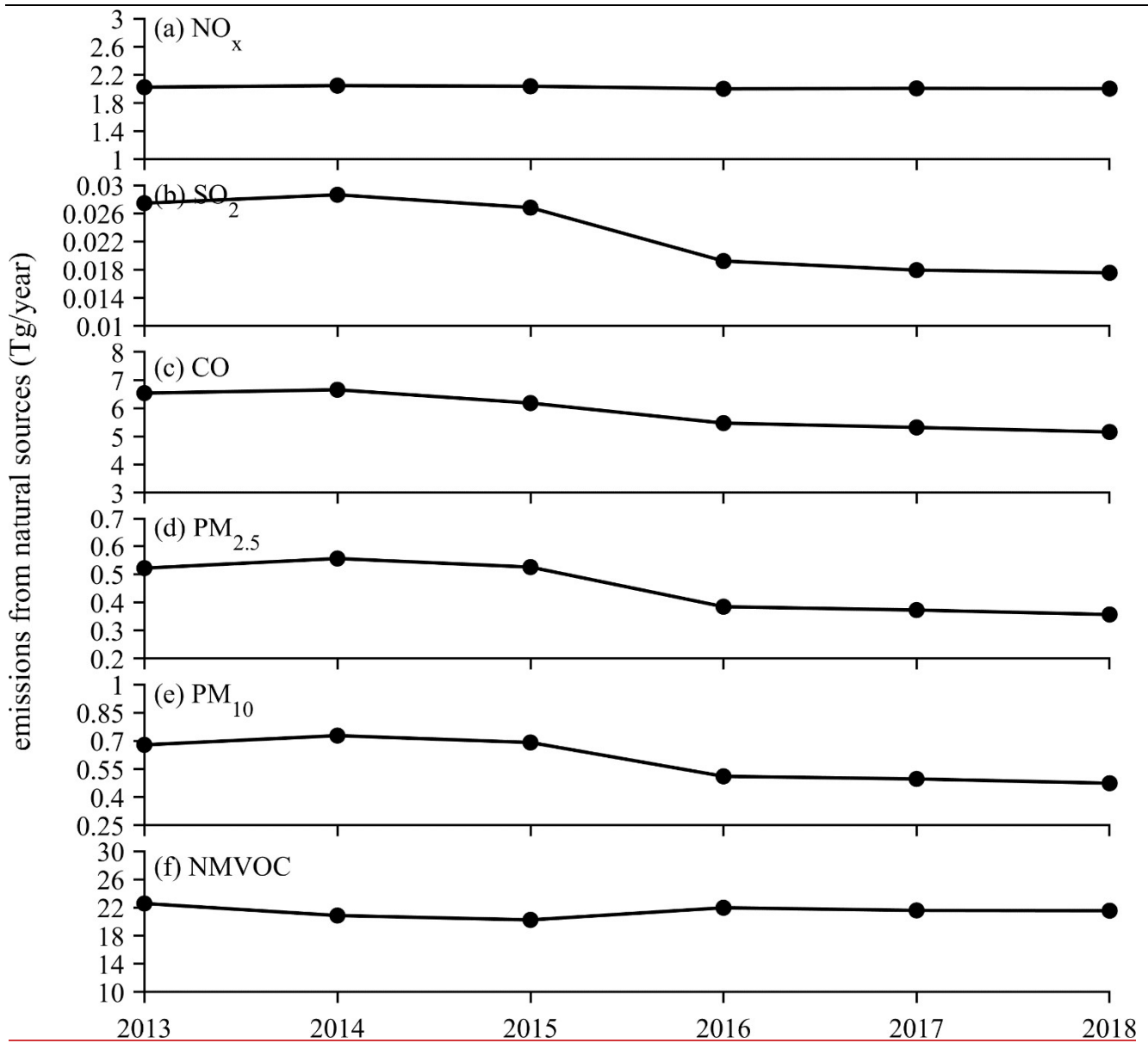
Figure S15: Same as Fig. S10 but for NMVOC.

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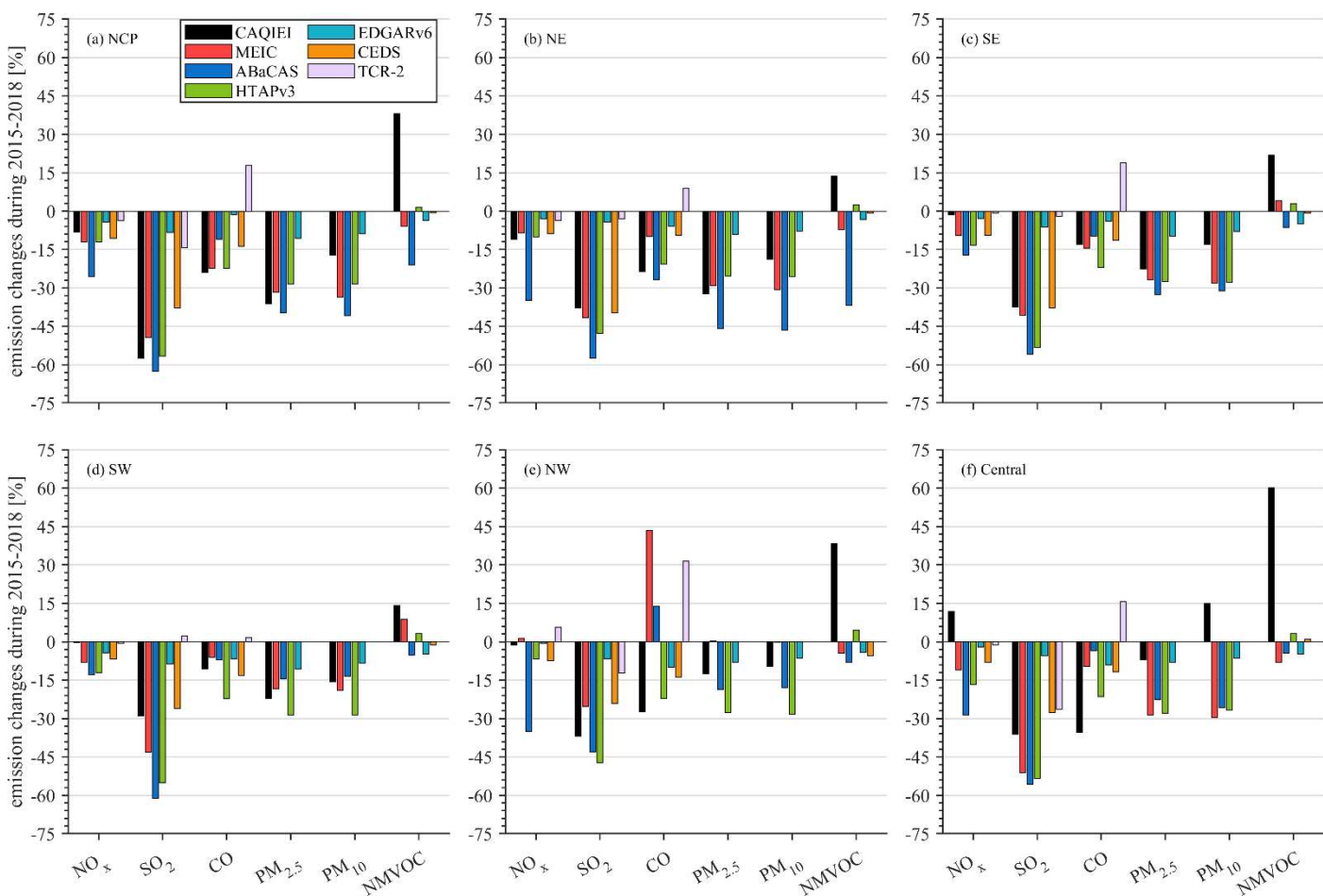
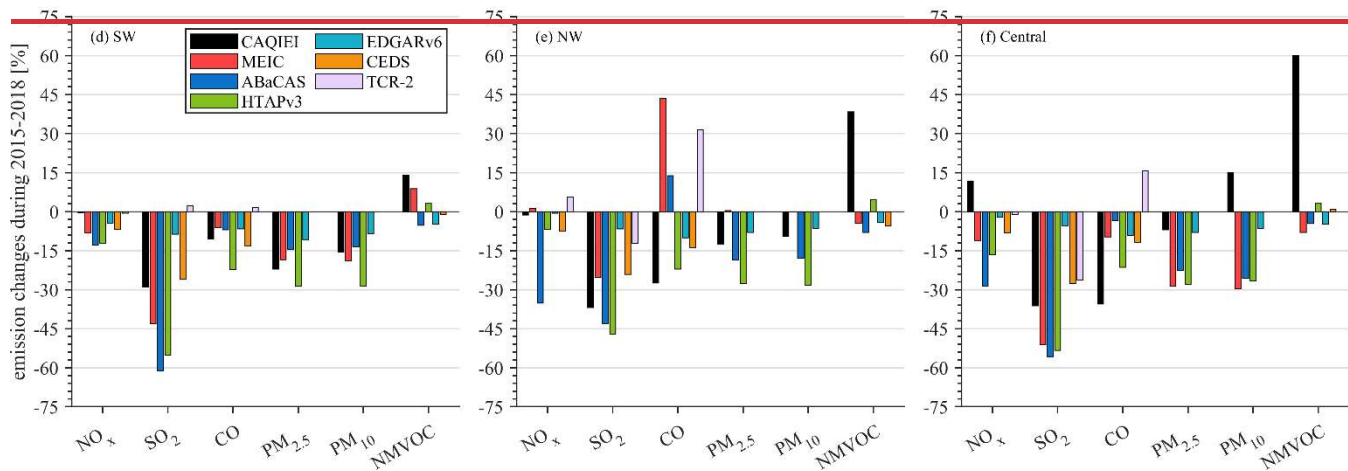
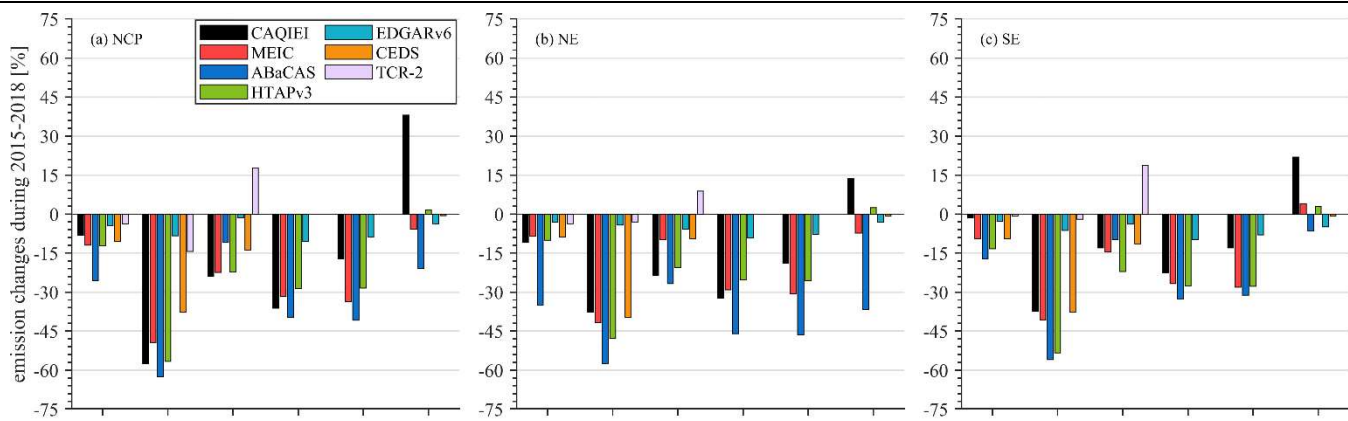
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 179 **Figure S16: Time series of annual natural emissions of (a) NO<sub>x</sub>, (b) SO<sub>2</sub>, (c) CO, (d) PM<sub>2.5</sub>, (e) PM<sub>10</sub> and (f) NMVOC in China from 2013**  
 180 **to 2018. The considered natural sources includes the biogenic, biomass burning and soil emissions.**

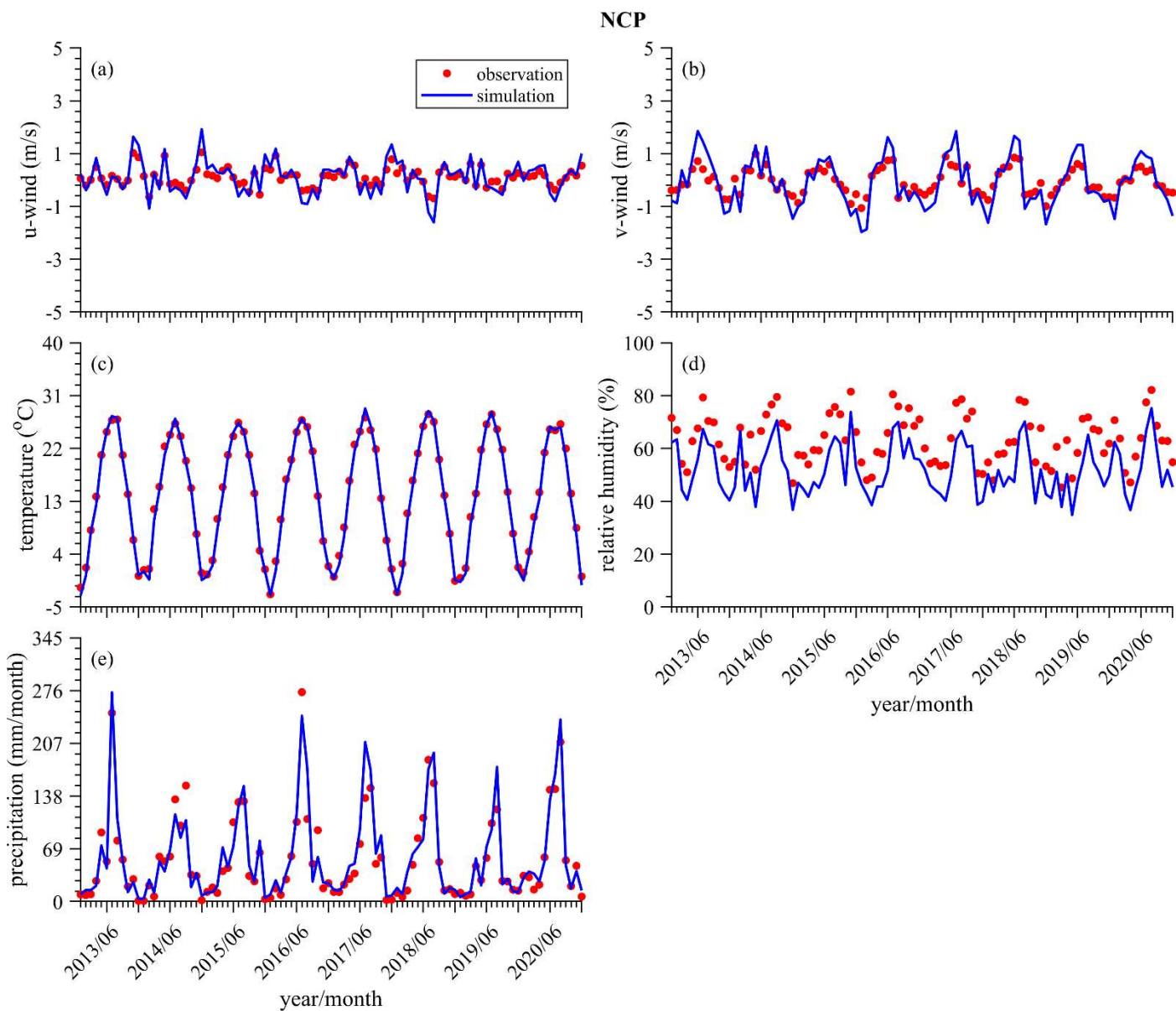




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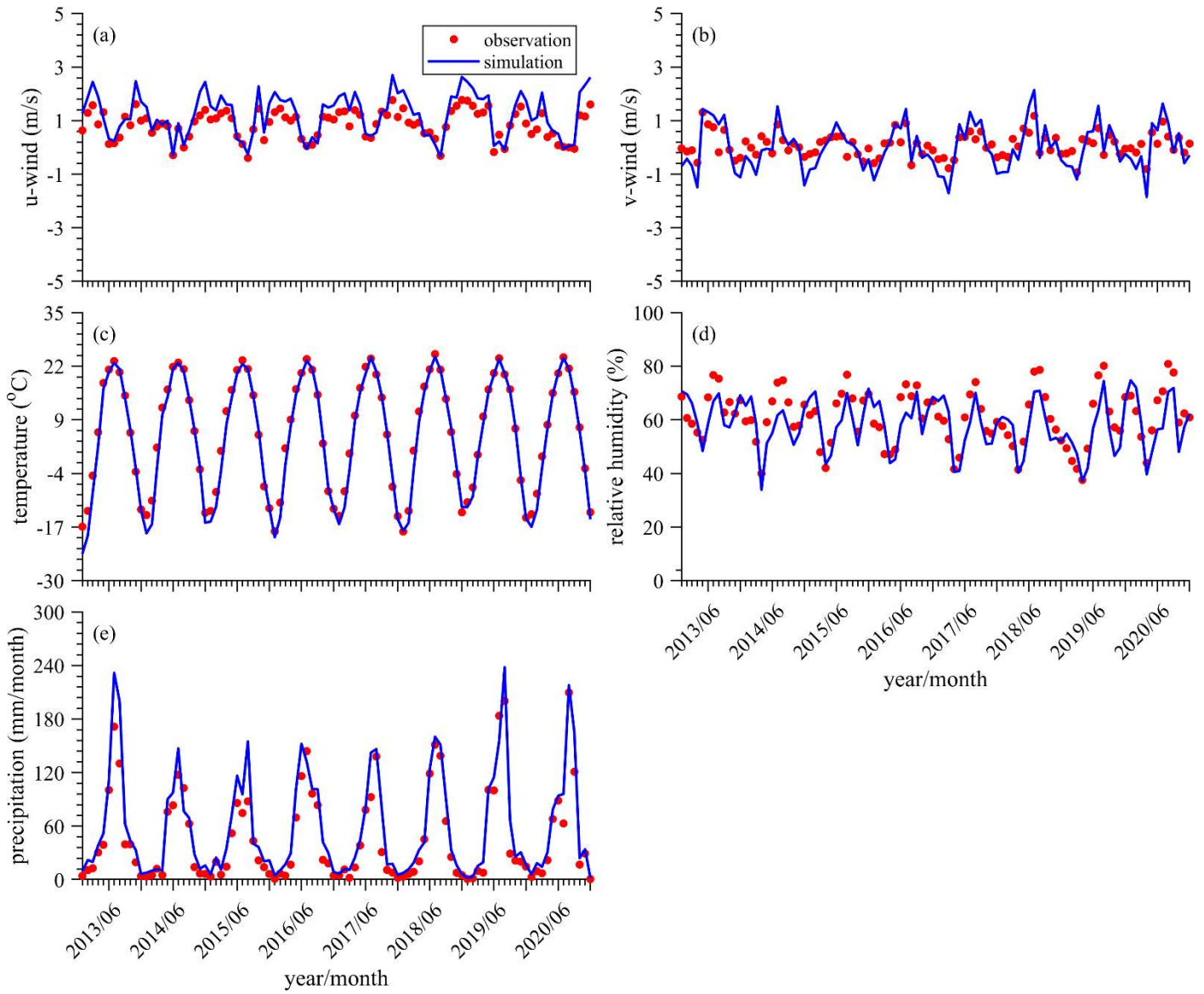
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184 **Figure S17: Comparisons of the calculated emission changes of (a)-NO<sub>x</sub>, (b)-SO<sub>2</sub>, (c)-CO, (d)-PM<sub>2.5</sub>, (e)-PM<sub>10</sub>, and (f)-**  
185 **NMVOCs over different(a) NCP, (b) NE, (c) SE, (d) SW, (e) NW and (f) Central regions of China from 2015 to 2018**  
186 **between CAQIEI and previous inventories.**  
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189 **Figure S18: Timeseries of observed (red dots) and simulated (blue line) monthly values of (a) zonal wind, (b) meridional**  
190 **wind, (c) temperature, (d) relative humidity and (e) precipitation over NCP region from Jan 2013 to Dec 2020.**  
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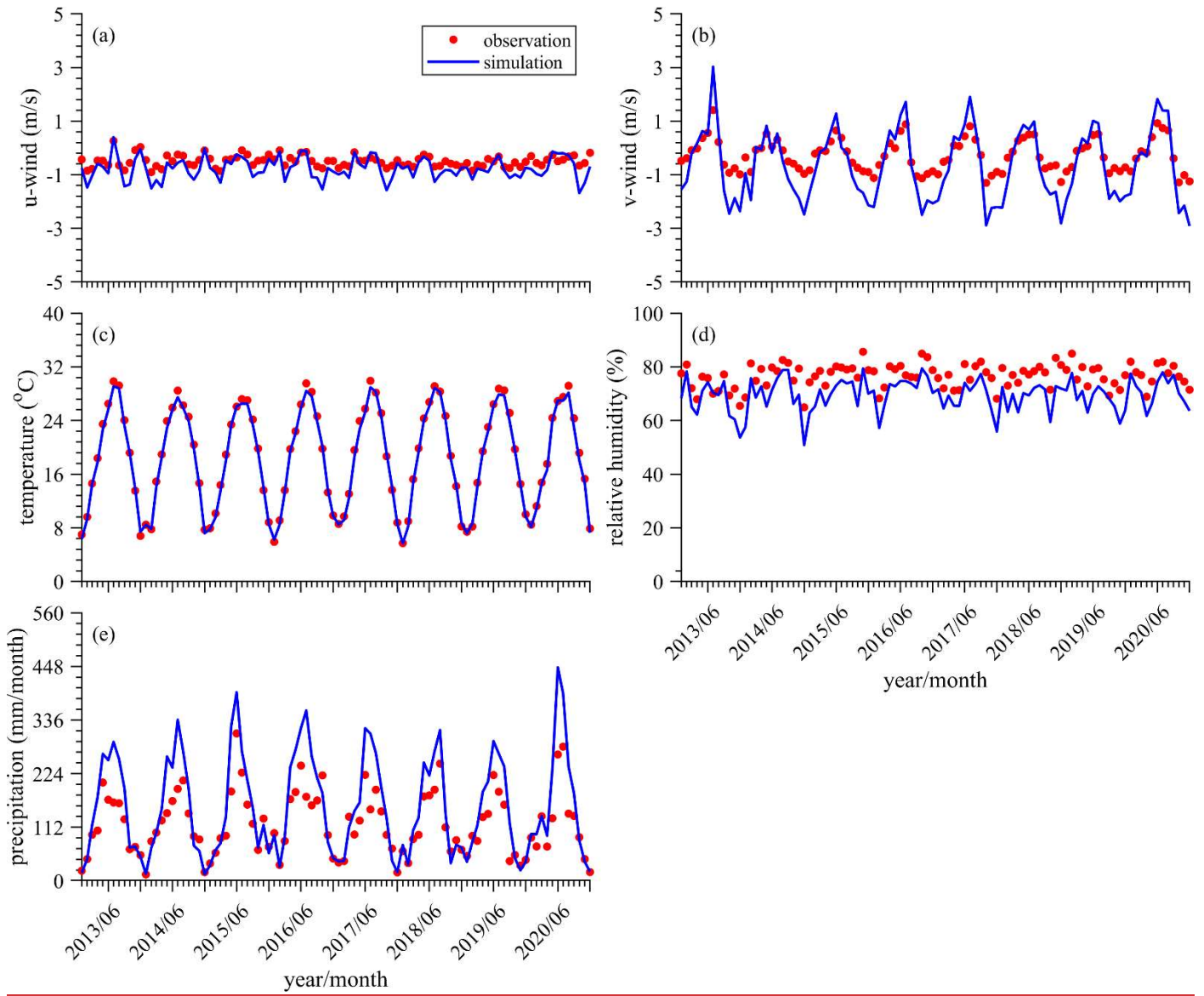
NE



**Figure S19: Same as in Figure S18 but over the NE region.**

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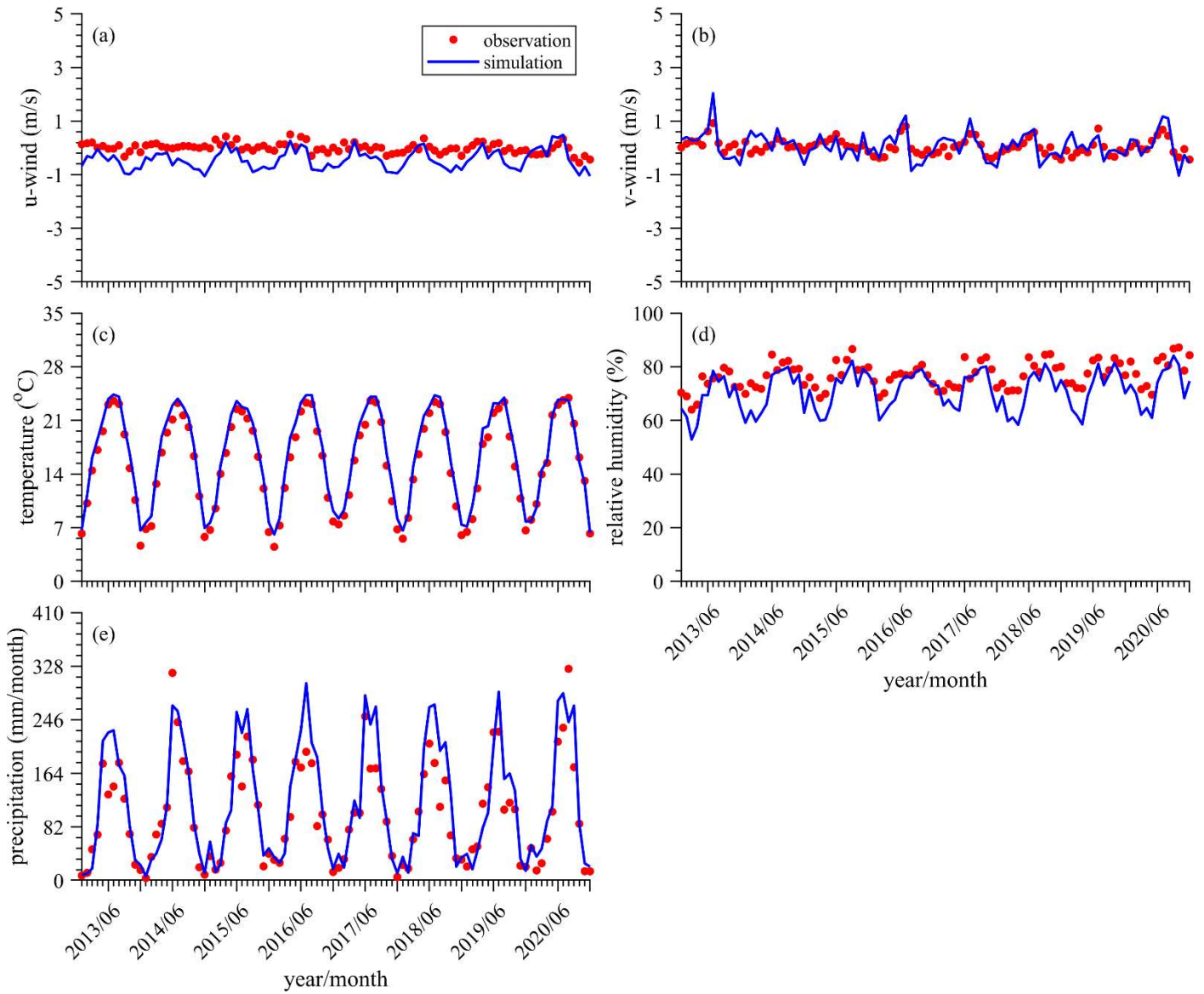
SE



**Figure S20: Same as in Figure S18 but over the SE region.**

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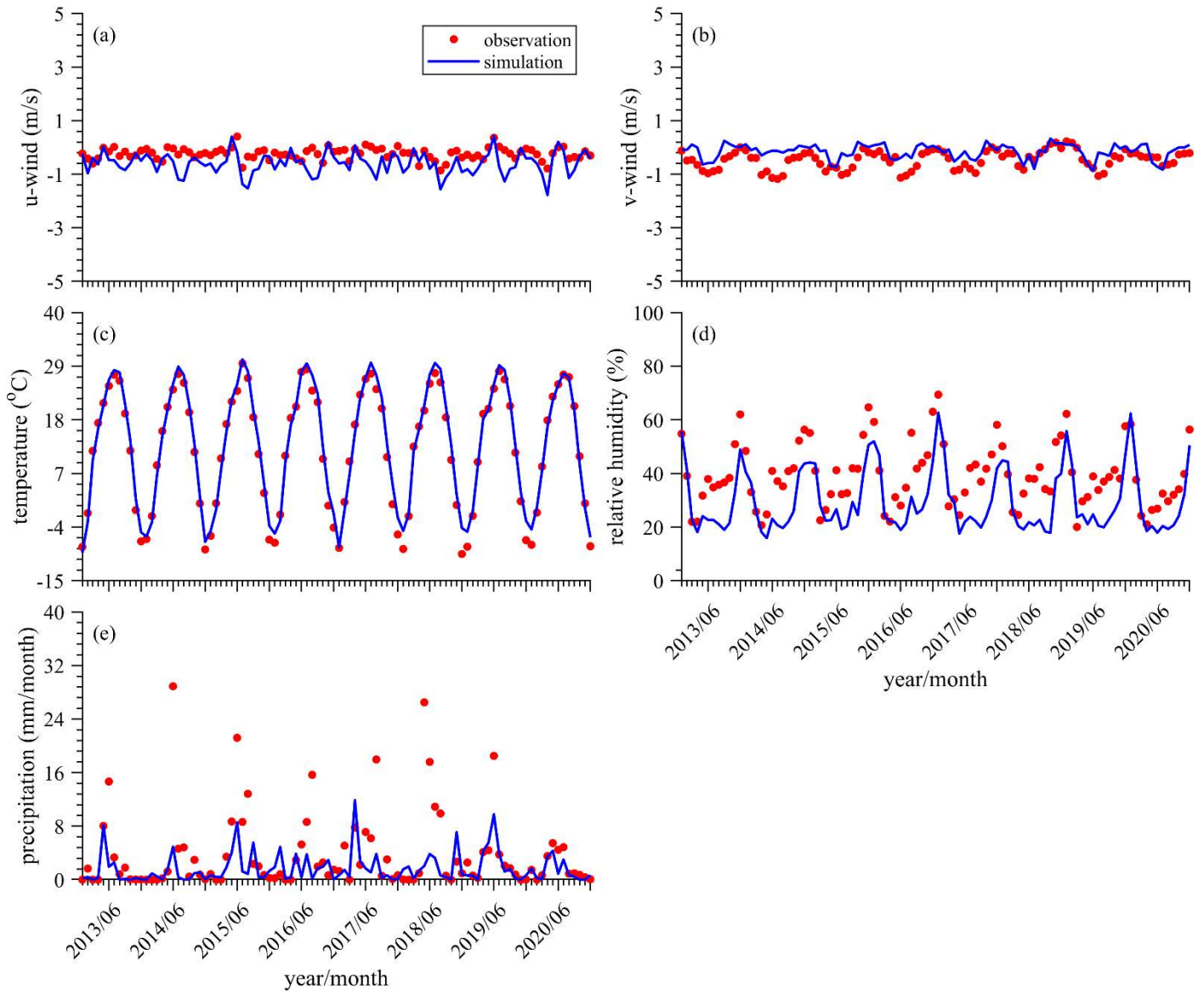
SW



**Figure S21: Same as in Figure S18 but over the SW region.**

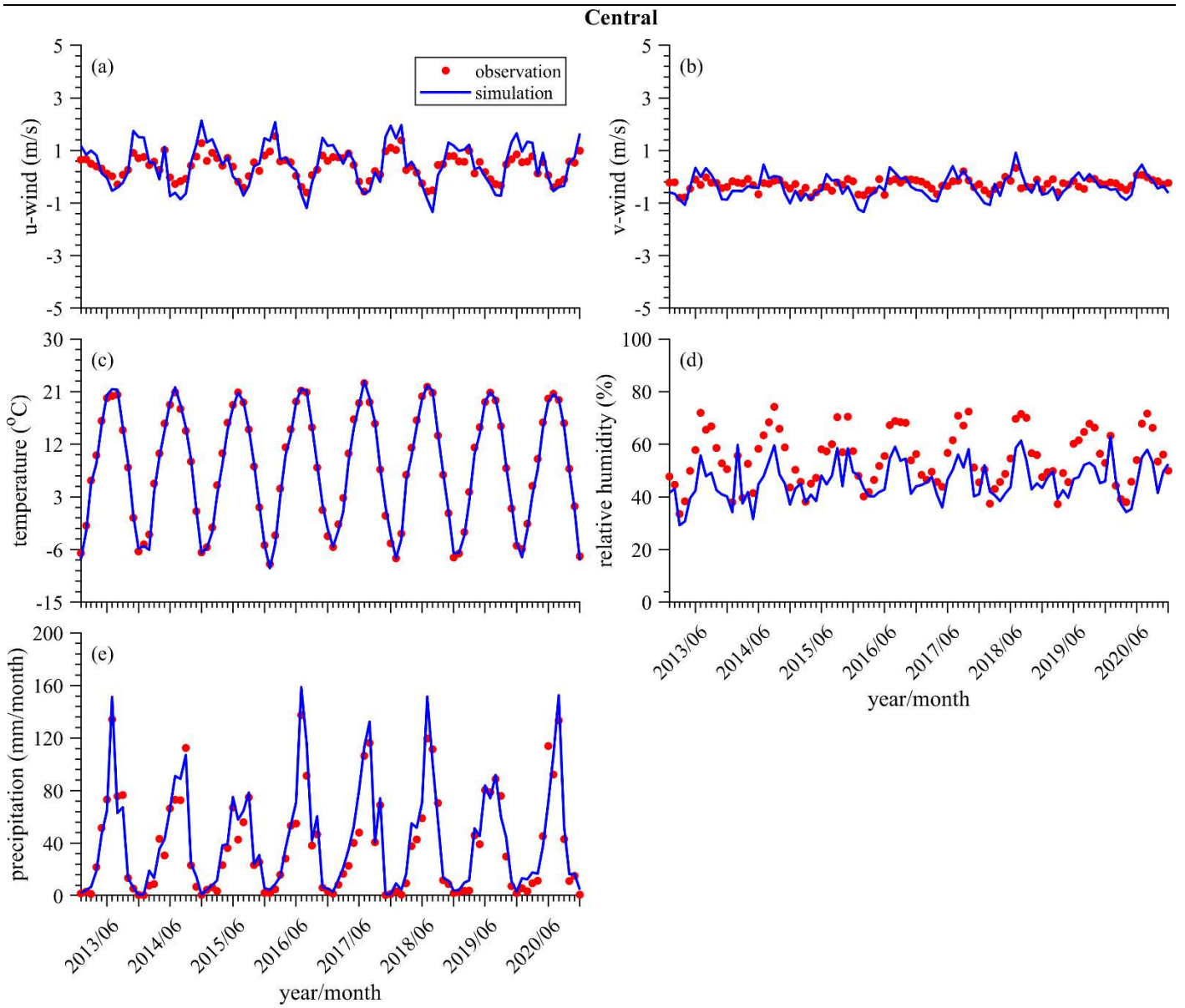
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NW



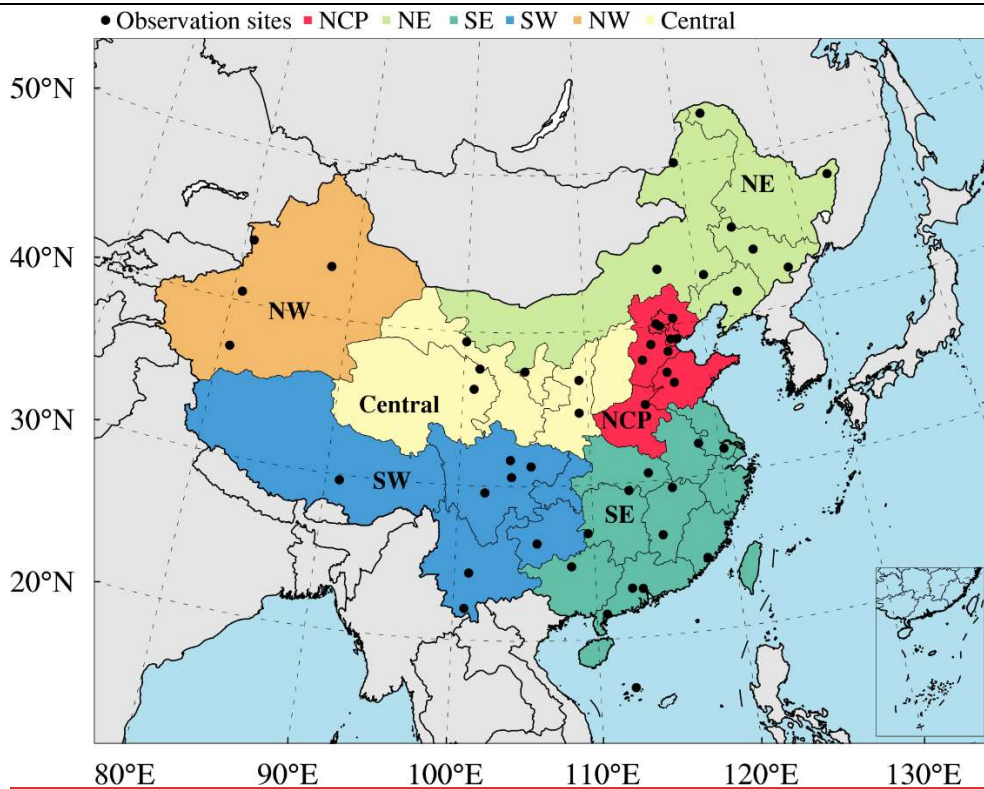
**Figure S22: Same as in Figure S18 but over the NW region.**

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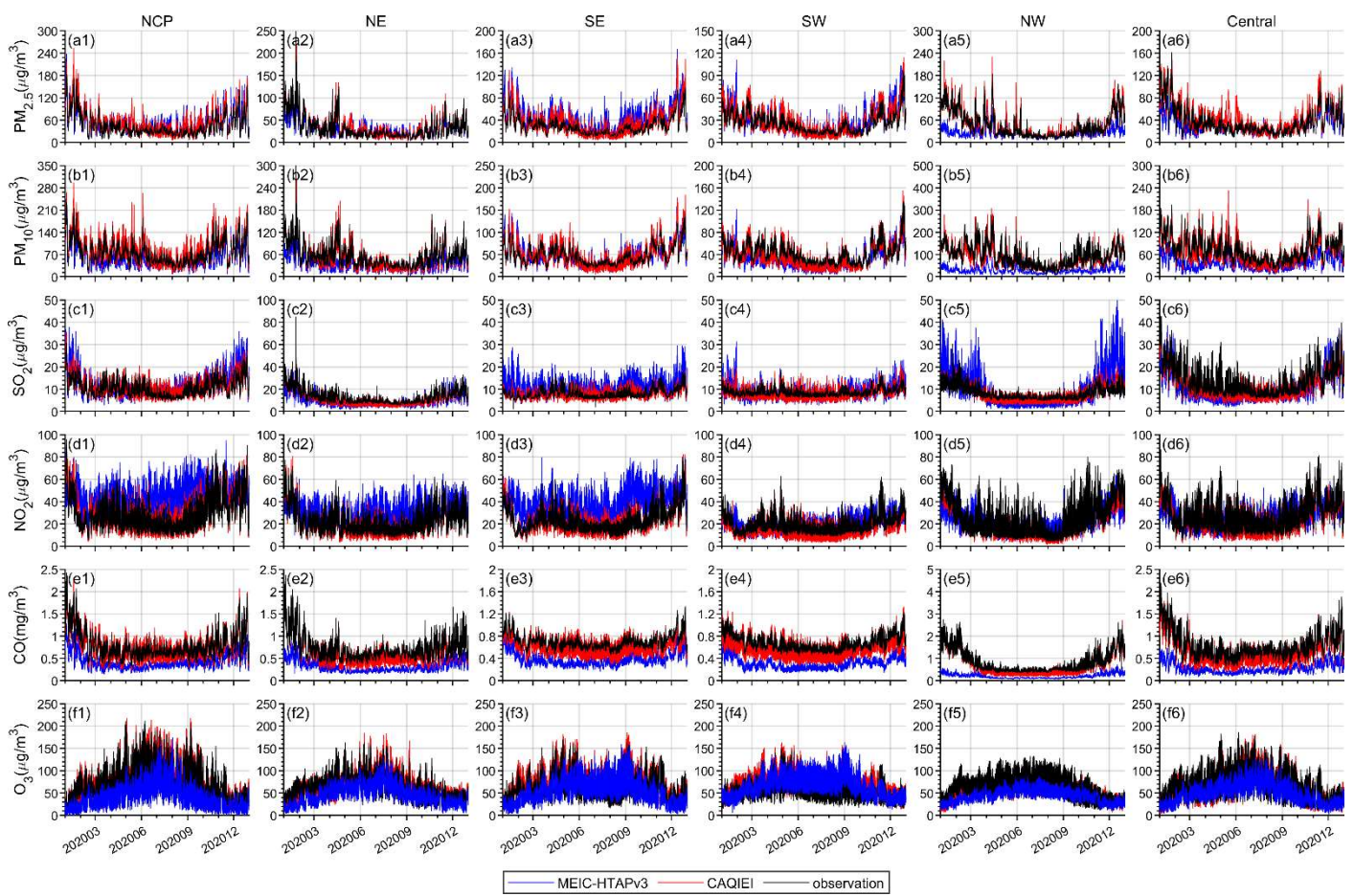


**Figure S23: Same as in Figure S18 but over the Central region.**

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**Figure S24: Spatial distribution of meteorological observation sites used in the evaluation of meteorology simulations over different regions of China**



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**Figure S25: Timeseries of observed (black lines) and simulated concentrations of different air pollutants in China driven by CAQIEI (red lines) and MEIC-HTAPv3 (blue lines) over different regions of China.**