1	A Regional multi-Air Pollutant Assimilation System (RAPAS v1.0)
2	for emission estimates: System development and application
3	Shuzhuang Feng ¹ , Fei Jiang ^{1,2} , Zheng Wu ³ , Hengmao Wang ^{1,2} , Wei He ¹ , Yang Shen ¹ ,
4	Lingyu Zhang ¹ , Yanhua Zheng ¹ , Chenxi Lou ¹ , Ziqiang Jiang ⁴ , Weimin Ju ^{1,2}
5	
6	¹ Jiangsu Provincial Key Laboratory of Geographic Information Science and Technology, International
7	Institute for Earth System Science, Nanjing University, Nanjing, 210023, China
8	² Jiangsu Center for Collaborative Innovation in Geographical Information Resource Development and
9	Application, Nanjing, 210023, China
10	³ Chongqing Institute of Meteorological Sciences, Chongqing, 401147, China
11	⁴ Jiangsu Environmental Monitoring Center, Nanjing, 210019, China
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16	Correspondence to: Fei Jiang (jiangf@nju.edu.cn)
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30 Abstract

Top-down atmospheric inversion infers surface-atmosphere fluxes from spatially 31 distributed observations of atmospheric composition in order to quantify anthropogenic 32 and natural emissions. In this study, we developed a Regional multi-Air Pollutant 33 Assimilation System (RAPAS v1.0) based on the Weather Research and 34 Forecasting/Community Multiscale Air Quality Modelling System (WRF/CMAQ) 35 model, the three-dimensional variational (3DVAR) algorithm, and the ensemble square 36 37 root filter (EnSRF) algorithm. This system can simultaneously assimilate hourly in-situ CO, SO₂, NO₂, PM_{2.5} and PM₁₀ observations to infer gridded emissions of CO, SO₂, 38 NO_x , primary $PM_{2.5}$ (PPM_{2.5}), and coarse PM_{10} (PMC) on a regional scale. In each data 39 assimilation window, we use a "two-step" scheme, in which the emissions are inferred 40 first, and then input into the CMAQ model to simulate initial conditions (IC) of the next 41 window. The posterior emissions are then transferred to the next window as prior 42 emissions, and the original emission inventory is only used in the first window. 43 Additionally, a "super-observation" approach is implemented to decrease the 44 computational costs, observation error correlations, and influence of representative 45 errors. Using this system, we estimated the emissions of CO, SO₂, NO_x, PPM_{2.5}, and 46 PMC in December and July 2016 over China using nationwide surface observations. 47 The results show that compared to the prior emissions (MEIC 2016), the posterior 48 emissions of CO, SO₂, NO_x, PPM_{2.5}, and PMC in December 2016 increased by 129%, 49 20%, 5%, 95%, and 1045%, respectively, and the emission uncertainties decreased by 50 44%, 45%, 34%, 52%, and 56%, respectively. With the inverted emissions, the RMSE 51 of simulated concentrations decreased by 40-56%. Sensitivity tests were conducted 52 53 with different prior emissions, prior uncertainties, and observation errors. The results showed that the "two-step" scheme employed in RAPAS is robust in estimating 54 emissions using nationwide surface observations over China. This study offers a useful 55 tool for accurately quantifying multi-species anthropogenic emissions at large scales 56 and in near real time. 57

58

59 1. Introduction

Owing to rapid economic development and pollution control legislation, there is an 60 increasing demand to provide updated emission estimates, especially in areas where 61 anthropogenic emissions are intensive. Accurately estimating source emission 62 quantities and spatiotemporal changes resulting from various regulations is imperative 63 and valuable for understanding air quality responses and is crucial for providing timely 64 instructions for the design of future emission regulations. However, most inventories 65 were developed based on a bottom-up approach and are usually updated with a delay 66 of a few years owing to the complexity of gathering statistical information on activity 67 levels and sector-specific emission factors (Ding et al., 2015). The large uncertainty 68 associated with the low temporal and spatial resolutions of these datasets also greatly 69 limits the assessment of emission changes. Some studies (Bauwens et al., 2020; Shi and 70 Brasseur, 2020) evaluated emission changes indirectly through concentration 71 measurements; however, air pollution changes are not only dominated by emission 72 changes, but also highly affected by meteorological conditions (Shen et al., 2021). 73

74 Top-down atmospheric inversion infers surface-atmosphere fluxes from spatially 75 distributed observations of atmospheric compositions. Recent efforts have been focused on developing air pollution data assimilation (DA) systems to conduct top-down 76 inversions, which can integrate model and multi-source observational information to 77 constrain emission sources. Two major methods are widely used in those DA systems: 78 4D-variational data assimilation (4DVAR) and ensemble Kalman filter (EnKF). 79 4DVAR provides a global optimal analysis by minimizing a cost function. It shows an 80 implicit flow-dependent background error covariance and can reflect complex 81 82 nonlinear constraint relationships (Lorenc, 2003). Additionally, a weak constraint 4DVAR method can partly account for the model error by defining a systematic error 83 term in a cost function (Derber, 1989). For example, the GEOS-Chem and TM5 4DVAR 84 frameworks have been used to estimate CH₄ (Alexe et al., 2015; Monteil et al., 2013; 85 Schneising et al., 2009; Stanevich et al., 2021; Wecht et al., 2014) and CO₂ fluxes (Basu 86 et al., 2013; Nassar et al., 2011; Wang et al., 2019a) from different satellite retrieval 87

products. Additionally, Jiang et al. (2017) and Stavrakou et al. (2008) also used the 88 4DVAR algorithm to estimate global CO and NO_x emission trends using MOPITT and 89 GOME/SCIAMACHY retrievals, respectively. Using NIES LiDAR observations, 90 Yumimoto et al. (2008) applied the 4DVAR DA to infer dust emissions over eastern 91 Asia and the results agreed well with various satellite data and surface observations. 92 Based on surface observations, Meirink et al. (2008) developed a 4DVAR system to 93 optimize monthly methane emissions, which showed a high degree of consistency in 94 95 posterior emissions and uncertainties when compared with an analogous inversion 96 based on the traditional synthesis approach.

Although considerable progress has been made to reduce large uncertainties in emission 97 98 inventories, the drawback of the 4DVAR method is the additional development of 99 adjoint models, which are technically difficult and cumbersome for complex chemical transport models (Bocquet and Sakov, 2013). Instead, EnKF uses flow-dependent 100 101 background error covariance generated by ensemble simulations to map deviations in 102 concentrations to increments of emissions, which is more flexible and easier to 103 implement. Many previous studies used EnKF techniques to assimilate single- or dual-104 species observations to optimize the corresponding emission species (Chen et al., 2019; Peng et al., 2017; Schwartz et al., 2014; Sekiyama et al., 2010). Miyazaki et al. (2017) 105 improved NO_x emission estimates using multi-constituent satellite observations, and 106 107 further estimated global surface NO_x emissions from 2005 to 2014. Feng et al., (2020b) used surface observations of NO_2 to infer the NO_x emission changes in China during 108 the COVID-19, and quantitatively evaluate the impact of the epidemic on economic 109 activities from the perspective of emission change. Tang et al. (2011) adjusted the 110 emissions of NO_x and VOCs through assimilating surface O₃ observations and achieved 111 an better performance in O_3 forecasts. However, such a revision may encounter the 112 problem of model error compensation rather than a retrieval of physically meaningful 113 quantities, which should be avoided from overfitting for emission inversion purposes 114 (Bocquet, 2012; Navon, 1998; Tang et al., 2011). The EnKF has also been widely 115 applied to optimize emissions of carbon dioxide (Jiang et al., 2021; Liu et al., 2019), 116

carbon monoxide (Feng et al., 2020a; Mizzi et al., 2018), sulfur dioxide (Chen et al.,
2019), ammonia (Kong et al., 2019), etc.

Multi-species data assimilation can efficiently reduce the uncertainty in emission 119 inventories and has led to improvements in air quality forecasting (Ma et al., 2019; 120 121 Miyazaki et al., 2012b) as it offers additional constraints on emission estimates through improvements in related atmospheric fields, chemical reactions, and gas-particle 122 123 transformations (Miyazaki and Eskes, 2013). Barbu et al. (2009) updated sulfur oxide 124 (SO_x) emissions with SO₂ and sulfate aerosol observations and found that the simultaneous assimilation of both species performed better than assimilating them 125 126 separately. Muller and Stavrakou (2005) also found that the simultaneous optimization 127 of the sources of CO and NO_x led to better agreement between simulations and observations compared to the case where only CO observations are used. 128

The deviation in the chemical initial conditions (IC) is an important source of error that 129 affects the accuracy of emission inversion because atmospheric inversion fully 130 131 attributes the biases in simulated and observed concentrations to deviations in emissions (Meirink et al., 2006; Peylin et al., 2005). The biases of concentrations would 132 be compensated by the unreasonable adjustment of pollution emissions without the 133 optimization of ICs (Tang et al., 2013). Simultaneously optimizing chemical ICs and 134 135 emissions has been applied to constrain emissions in many previous studies (Ma et al., 2019; Miyazaki et al., 2012a; Peng et al., 2018). For example, Elbern et al. (2007) 136 adjusted O₃, NO_x, and VOCs ICs, and NO_x and VOCs emissions jointly through 137 assimilating surface O_3 and NO_x observations. Although the forecast skills of O_3 were 138 139 improved, due to the coarse model resolution and the strong nonlinear relationship between O_3 and NO_x , the assimilation of O_3 observation worsened emission inversion 140 and forecast of NOx. Peng et al. (2018) assimilated near-surface observations to 141 simultaneously optimize the ICs and emissions. In the 72-hr forecast evaluation, their 142 resultant emission succeeded in improving SO₂ forecast while having little influence 143 on CO and aerosol forecast and even degrading the forecast of NO_2 . Ma et al. (2019) 144 also found that the DA benefits for forecast almost disappeared after 72 hr using 145

optimized ICs and emissions. Although a large improvement has been achieved, this 146 method has significant limitations in emission inversion as the contributions from the 147 emissions and chemical ICs to the model's biases are difficult to distinguish (Jiang et 148 al., 2017). In addition, the constraints of the chemical ICs with observations in each 149 assimilation window make the emission inversions between the windows independent. 150 This means that if the emission in one window is overestimated or underestimated, it 151 cannot be transferred to the next window for further correction and compensation. 152 153 Considering the importance of emissions in chemical field prediction (Bocquet et al., 2015), the rapid disappearance of the DA benefits seems unrealistic, indicating that 154 simultaneously optimizing chemical ICs and emissions may result in a systematic bias 155 in the inverted emissions (Jiang et al., 2021). 156

Since 2013, China has deployed an air pollution monitoring network that publishes 157 nationwide and real-time hourly surface observations. This dataset provides an 158 opportunity to improve emission estimates using DA. In this study, a regional multi-air 159 160 pollutant assimilation system using 3DVAR and EnKF DA techniques was constructed to simultaneously assimilate various surface observations (e.g. CO, SO₂, NO₂, O₃, 161 PM_{2.5}, and PM₁₀). We adopted a "two-step" method in this system, in which the ICs of 162 each DA window were simulated using the posterior emissions of the previous DA 163 window. The capabilities of RAPAS for reanalysis field generation and emission 164 inversion estimation were also evaluated. The robustness of the system was investigated 165 with different prior inventories, uncertainty settings of prior emissions, and observation 166 errors. The remainder of the paper is organized as follows: Section 2 introduces the DA 167 system and observation data, Section 3 describes the experimental design, Section 4 168 169 presents and discusses the results of the system performance and sensitivity tests, and Section 5 concludes the paper. 170

171 **2. Method and data**

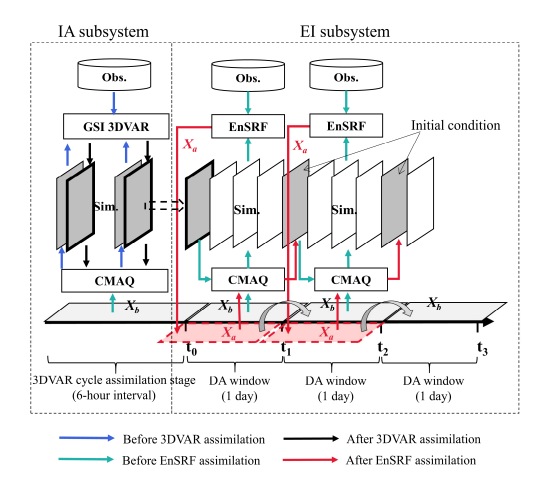
172 **2.1 System description**

2.1.1 Procedure of the assimilation system

A regional air pollutant assimilation system has been preliminarily constructed and 174 successfully applied in our previous studies to optimize the gridded CO and NO_x 175 emissions (Feng et al., 2020a; Feng et al., 2020b). Herein, the system was further 176 extended to simultaneously assimilate multiple species (e.g. CO, SO₂, NO₂, O₃, PM_{2.5}, 177 and PM₁₀) and officially named the Regional multi- Air Pollutant Assimilation System 178 (RAPASv1.0). The RAPAS has three components: a regional chemical transport model 179 (CTM), which is coupled offline and used to simulate the meteorological fields and 180 atmospheric compositions, and the 3DVAR and ensemble square root filter (EnSRF) 181 modules, which are used to optimize chemical ICs (Feng et al., 2018; Jiang et al., 2013b) 182 and anthropogenic emissions (Feng et al., 2020a; Feng et al., 2020b), respectively. 183 3DVAR was introduced considering its excellent performance in our previous study and 184 the lower computational cost during the spin-up period in optimizing ICs. Additionally, 185 the 3DVAR method can obtain a better IC than the EnKF method (Schwartz et al., 2014). 186

187 Based on the above three components, the RAPAS was divided into two subsystems: 188 the IC assimilation (IA) subsystem (CTM plus 3DVAR) and the emission inversion (EI) subsystem (CTM plus EnSRF). As shown in Figure 1, the IA subsystem was first run 189 190 to optimize the chemical ICs (Kleist et al., 2009; Wu et al., 2002) for the subsequent EI subsystem. Distinguish the source type of model-observation mismatch error was not 191 required for the IA subsystem. The EI subsystem runs cyclically with a "two-step" 192 scheme. In the first step, the prior emissions (X^b) are perturbed and input into the CTM 193 model to simulate chemical concentration ensembles. The simulated concentrations of 194 195 the lowest model level were then interpolated to the observation space according to the locations and times of the observations using the nearest-neighbor interpolation method. 196 Prior emissions (X^b) , simulated observations and real observations were entered into 197 the EnSRF module to generate optimized emissions (X^{a}) . In the second step, the 198 optimized emissions were re-entered into the CTM model to generate the ICs of the 199 next DA window. Meanwhile, the optimized emissions were transferred to the next 200 window as prior emissions. Unlike joint adjustment of ICs and emissions ("one-step" 201 scheme) in emission inversion (Chen et al., 2019), the "two-step" scheme needs to run 202

203 the CTM model twice, which is time consuming but can transfer the potential errors of 204 the inverted emissions in one DA window to the next for further correction.



205

Figure 1. Composition and flow chart of RAPAS. \mathbf{x}_{a} and \mathbf{x}_{b} represent the prior and posterior emissions. The 3DVAR assimilation stage lasts five days with data input frequency of six hours and the DA window in the EI subsystem is set to one day.

209 2.1.2 Atmospheric transport model

chemical transport model of the Weather Research 210 The regional and Forecasting/Community Multiscale Air Quality Modelling System (WRF/CMAQ) was 211 212 adopted in this study. CMAQ is a regional 3-D Eulerian atmospheric chemistry and 213 transport model with a "one-atmosphere" design developed by the US Environmental Protection Agency (EPA). It can simultaneously address the complex interactions 214 among multiple pollutants/air quality issues. The CMAQ model was driven by the WRF 215 model, which is a state-of-the-art mesoscale numerical weather prediction system 216

designed for both atmospheric research and meteorological field forecasting. In this 217 study, WRF version 4.0 and CMAQ version 5.0.2 were used. The WRF simulations 218 were performed with a 36-km horizontal resolution on 169×129 grids, covering all of 219 mainland China (Figure 2). This spatial resolution has been widely adopted in regional 220 simulations as it can provide good simulations of spatiotemporal variations in air 221 pollutants (Mueller and Mallard, 2011; Sharma et al. 2016). In the vertical direction, 222 there were 51 sigma levels on the sigma-pressure coordinates extending from the 223 224 surface to 100 hPa. The underlying surface of the urban and built-up land was replaced by the MODIS land cover retrieval of 2016 to adapt to the rapid expansion of 225 urbanization. The CMAO model was run with the same domain but with three grid cells 226 removed from each side of the WRF domain. There were 15 layers in the CMAQ 227 vertical coordinates, which were interpolated from 51 WRF layers. 228

The meteorological initial and lateral boundary conditions were both provided by the 229 Final Operational Global Analysis data of the National Center for Environmental 230 Prediction (NCEP) with a $1^{\circ} \times 1^{\circ}$ resolution at 6-h intervals. The chemical lateral 231 232 boundary conditions and chemical ICs in the IA subsystem originate from background 233 profiles. As mentioned above, in the EI subsystem, the chemical IC in the first window is provided by the IA subsystem and in the following windows, it is forward simulated 234 using optimized emissions from the previous window. Carbon Bond 05 with updated 235 toluene chemistry (CB05tucl) and the 6th generation aerosol module (AERO6) were 236 chosen as the gas-phase and aerosol chemical mechanisms, respectively (Appel et al., 237 2013; Sarwar et al., 2012). The detailed physical and chemical configurations are listed 238 in Table 1. 239

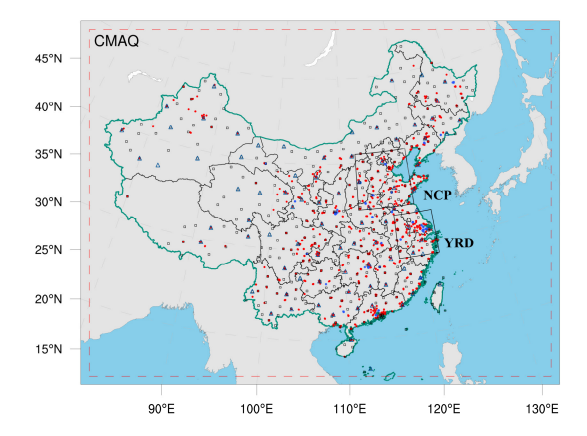


Figure 2. Model domain and observation network. The red dashed frame depicts the CMAQ computational domain; the black squares represent the surface meteorological measurement sites; the navy blue triangles represent the sounding sites; and the red and blue dots represent the air pollution measurement sites. Observations from all sites were assimilated in the 3DVAR subsystem, while observations of city sites where red dots were averaged are used for assimilation and where blue dots were averaged are used for independent evaluation in the EI subsystem; the boxed subregions are the North China Plain (NCP) and Yangtze River Delta (YRD); and the shaded area depicts the topography.

WRF		CMAQ				
Parameter	Scheme	Parameter	Scheme			
Microphysics	WSM6	Horizontal/Vertical advection	yamo/wrf			
Longwave	RRTM	Horizontal/Vertical diffusion	multiscale/acm2			
Shortwave	Goddard	Deposition	m3dry			
Boundary layer	ACM	Chemistry solver	EBI			
Cumulus	Kain-Fritsch	Photolysis	phot_inline			
Land-surface	Noah	Aerosol module	AERO6			
Surface layer	Revised	Cloud module	cloud_acm_ae6			
Urban canopy	No	Gas-phase chemistry	CB05tucl			

255 **Table 1.** Configuration options of WRF/CMAQ

256 2.1.3 3DVAR assimilation algorithm

Grid-point Statistical Interpolation (GSI) developed by the US NCEP was utilized in 257 258 this study. Building on the work of Liu et al. (2011), Jiang et al. (2013b) and Feng et al. (2018), we extended GSI to simultaneously assimilate multiple species (including CO, 259 SO₂, NO₂, O₃, PM_{2.5}, and PM₁₀) and first used individual aerosol species of PM_{2.5} as 260 261 analysis variables within the GSI/WRF/CMAQ framework. Additional work includes the construction of surface air pollutant observation operators, the updating of 262 observation errors, and the statistics of background error covariance for the analysis 263 variables. Moreover, the data interface was modified to read/write the CMAQ 264 output/input file directly, which was easy to implement. 265

In the sense of minimum analysis error variance, the 3DVAR algorithm optimizes the analysis fields with observations by an iterative process to minimize the cost function J(x) defined below:

²⁶⁹
$$J(\mathbf{x}) = \frac{1}{2} (\mathbf{x}_{a} - \mathbf{x}_{b})^{T} \mathbf{B}^{-1} (\mathbf{x}_{a} - \mathbf{x}_{b}) + \frac{1}{2} [H(\mathbf{x}_{a}) - \mathbf{y}]^{T} \mathbf{R}^{-1} [H(\mathbf{x}_{a}) - \mathbf{y}],$$
(1)

where $\mathbf{x}_{\mathbf{a}}$ is a vector of the analysis field, \mathbf{x}_{b} is the background field, \mathbf{y} is the vector of observations, \mathbf{B} and \mathbf{R} are the background and observation error covariance matrices, respectively, representing the relative contributions to the analysis, and H is the observation operator that maps the model variables to the observation space.

The analysis variables were the 3D mass concentrations of the pollution components 274 (e.g. CO and sulfate) at each grid point. Hourly mean surface pollution observations 275 within a one-hour window of the analysis were assimilated. To assimilate the surface 276 pollution observations, model-simulated compositions were first diagnosed at 277 observation locations. For gas concentrations to be directly used as analysis variables, 278 the units need to be converted from ppm and ppb to mg m⁻³ and μ g m⁻³, respectively, to 279 match the observations. The model-simulated PM2.5 and PM10 concentrations at the 280 ground level were diagnosed as follows: 281

282
$$PM_{2.5} = f_i \times PM_i + f_j \times PM_j + f_k \times PM_k = OC + EC + SO_4^{2-} + NO_3^{-} + NH_4^{+} +$$

283
$$SEAS + AP_{2.5}$$
 (2)

284
$$PM_{10} = PM_i + PM_i + PM_k = PM_{2.5} + PMC$$
 (3)

where f_i , f_j , and f_k are the PM_{2.5} fractions of the Aitken, accumulation, and coarse 285 modes, respectively. These ratios are recommended as the concentrations of PM2.5 and 286 fine mode aerosols (i.e. Aitken plus accumulation) can differ because PM_{2.5} particles 287 include small tails from the coarse mode in the CMAQ model (Binkowski and Roselle, 288 289 2003; Jiang et al., 2006). PM_i , PM_j , and PM_k are the mass concentrations of the three modes in the CMAQ model, respectively. Seven aerosol species of PM2.5 (organic 290 carbon (OC), elemental carbon (EC), sulfate (SO_4^{2-}) , nitrate (NO_3^{-}) , ammonium (NH_4^{+}) , 291 sea salt (SEAS), and fine-mode unspeciated aerosols $(AP_{2.5})$) and additional coarse 292 293 PM₁₀ (PMC) were extracted as analysis variables and were updated using the PM_{2.5} and PMC observations. Before calculating equation (1) within the GSI, the analysis 294 variables were bilinearly interpolated in the horizontal direction to the observation 295 locations. 296

Calculating background error covariance (B) is generally costly and difficult when a
high-dimensional numerical model is used. For simplification, B was represented as a

299 product of spatial correlation matrices and standard deviations (SDs).

 $B = DCD^T$ (4)

301
$$\mathbf{C} = \mathbf{C}_{\mathbf{x}} \otimes \mathbf{C}_{\mathbf{y}} \otimes \mathbf{C}_{\mathbf{z}}$$
(5)

where **D** is the background error SD matrix; **C** is the background error correlation matrix; \otimes is the Kronecker product; and C_x , C_y , and C_z denote three onedimensional correlation submatrices in the longitude, latitude, and vertical coordinate directions, respectively. C_x and C_y are assumed to be horizontally isotropic such that they can be represented using a Gaussian function. The correlation between any two points x_i and x_j in the horizontal direction is expressed as follows:

308
$$c(x_i, x_j) = e^{-\frac{(x_i - x_j)^2}{2L^2}}$$
 (6)

where *L* is the horizontal correlation scale estimated using the proxy of the background error (Figure 3). The vertical correlation matrix C_z is directly estimated from the model background field as C_z is only an $n_z \times n_z$ (here, $n_z=15$) matrix.

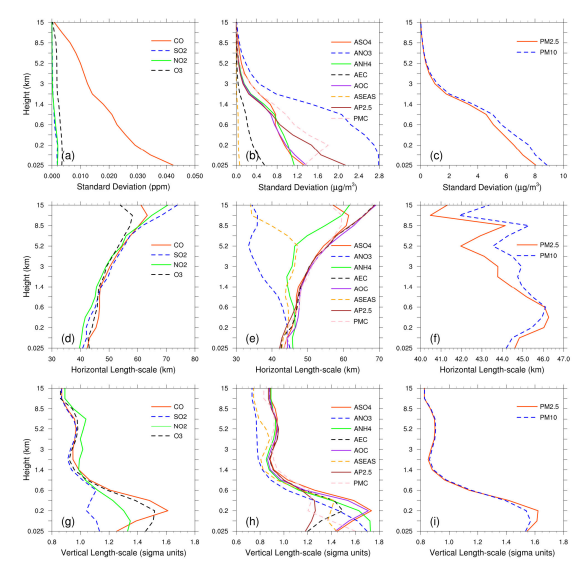


Figure 3. Vertical profiles of standard deviations (top, μg m⁻³), horizontal (middle, km)
and vertical (bottom, km) length scales for CO, SO₂, NO₂, O₃, sulfate, nitrate,
ammonium, EC, OC, sea salt, unspeciated aerosols (AP2.5), PMC, PM_{2.5} and PM₁₀.

312

To estimate these matrices, the "NMC" method was used to compute B for each variable 316 by taking the differences between forecasts of different lengths valid at the same time 317 (Parrish and Derber, 1992; Rabier et al., 1998). Differences between the 24- and 12-h 318 WRF/CMAQ forecasts of 60 pairs (two pairs per day) of analysis variables valid at 319 either 0000 or 1200 UTC over November 2016 were used. The horizontal and vertical 320 length scales of the correlation matrices were estimated using recursive filters (Purser 321 et al., 2003). The vertical distribution of the background error SDs, which varies with 322 height and species, is shown in Figure 3. The vertical profile of the background error 323

SDs corresponds to the vertical concentration distribution. This means that higher 324 concentrations tend to have larger background error SDs (e.g., CO and nitrate). These 325 SDs exhibit a common reduction as the height increases, especially at the top of the 326 boundary layer. The horizontal correlation of the background error determines the 327 propagation of observation information in this direction, whereas the vertical 328 correlation determines the vertical extension of such increments. For gaseous pollutants 329 and most individual aerosol components, the horizontal length scales increased with 330 height, whereas for the total particulate matter (i.e. PM_{2.5}, PM₁₀), the scales increased 331 with height in the boundary layer and decreased with height in the free troposphere. 332 The ground-level scale generally spread 40–45 km for all control variables. The vertical 333 length scale of most species first increased and then decreased with height, which may 334 be related to vertical mixing (Kahnert, 2008) and stack emissions at approximately 200 335 m height. 336

337 2.1.4 EnKF assimilation algorithm

In EnKF, the time-dependent uncertainties of the state variables are estimated using a Monte Carlo approach through an ensemble. Uncertainty can be propagated using linear or nonlinear dynamic models (flow-dependent background error covariance) by simply implementing ensemble simulations. The EnSRF algorithm introduced by Bierman (1977) and Maybeck (1979) was used to constrain pollution emissions in this study. EnSRF is a deterministic EnKF that obviates the need to perturb observations, which has a higher computational efficiency and a better performance (Sun et al., 2009).

The perturbation of the prior emissions represents the uncertainty. We implemented additive emission adjustment methods, which were calculated using the following function:

348

$$X_{i}^{b} = X_{0}^{b} + \delta X_{i}^{b}, i = 1, 2, ..., N$$
(7)

where **b** is the background (prior) state, *i* is the identifier of the perturbed samples, and N is the ensemble size, which was set to 40 considering the trade-off between computational cost and inversion accuracy (Figure S1). In contrast to the estimation of

parameters based on the augmentation of the conventional state vector (e.g. 352 concentrations) with the parameter variables, X only comprises emissions in this study 353 (similarly hereafter). δX_i^b is the randomly perturbed samples added to the prior 354 emissions X_0^b to produce ensemble samples of the inputs X_i^b . δX_i^b is drawn from 355 Gaussian distributions with a mean of zero and standard deviation of the prior emission 356 uncertainty in each grid. The state variables of the emissions include CO, SO₂, NO_x, 357 primary PM_{2.5} (PPM_{2.5}) and PMC. We used variable localization to update the analysis, 358 which means that the covariance among different state variables was not considered, 359 and the emission of one species was constrained only by its corresponding air pollutant 360 observation. This method has been widely used in chemical data assimilation systems 361 362 to avoid spurious correlations between species (Ma et al., 2019; Miyazaki et al., 2012b).

After obtaining an ensemble of state vectors (prior emissions), ensemble runs of the CMAQ model were conducted to propagate the errors in the model with each ensemble sample of state vectors. Combined with the observational vector y, the state vector $\overline{X^b}$ was updated by minimizing the analysis variance.

$$\overline{X^a} = \overline{X^b} + \mathbf{K}(\mathbf{y} - H\overline{X^b}) \tag{8}$$

³⁶⁸
$$\mathbf{K} = \mathbf{P}^{\mathbf{b}} \mathbf{H}^{T} (\mathbf{H} \mathbf{P}^{\mathbf{b}} \mathbf{H}^{T} + \mathbf{R})^{-1}$$
(9)

369
$$\boldsymbol{P}^{\boldsymbol{b}} = \frac{1}{N-1} \sum_{i=1}^{N} (\boldsymbol{X}_{i}^{\boldsymbol{b}} - \overline{\boldsymbol{X}^{\boldsymbol{b}}}) (\boldsymbol{X}_{i}^{\boldsymbol{b}} - \overline{\boldsymbol{X}^{\boldsymbol{b}}})^{T}$$
(10)

$$\delta X_i^a = \delta X_i^b - \widetilde{K} H \delta X_i^b \tag{11}$$

³⁷¹ While employing sequential assimilation and independent observations, \tilde{K} is ³⁷² calculated as follows:

373
$$\widetilde{K} = \left(1 + \sqrt{\frac{R}{(HP^bH^T + R)}}\right)^{-1} \mathbf{K}$$
(12)

where $\overline{X^b}$ is the mean of the ensemble samples X_i^b ; H is the observation operator that maps the model space to the observation space, consisting of the model integration

process converting emissions into concentrations and spatial interpolation matching the 376 model concentration to the locations of the observations; $y - H\overline{X^b}$ reflects the 377 differences between the simulated and observed concentrations: P^{b} is the ensemble-378 estimated background (a priori) error covariance; K is the Kalman gain matrix of the 379 ensemble mean depending on the background error covariance P^b and the observation 380 error covariance R, representing the relative contributions to analysis; and \tilde{K} is the 381 Kalman gain matrix of the ensemble perturbation, which is used to calculate emission 382 perturbations after inversions δX_i^a . The ensemble mean $\overline{X^a}$ of the analyzed state was 383 considered the best estimate of the emissions. 384

When large volumes of site observations are at a much higher resolution than the model 385 grid spacing, many correlated or fully consistent model-data mismatch errors can 386 appear in one cluster, resulting in excessive adjustments and deteriorated model 387 performance (Houtekamer and Mitchell, 2001). To reduce the horizontal observation 388 error correlations and influence of representativeness errors, a "super-observation" 389 390 approach combining multiple noisy observations located within the same grid and assimilation window was developed based on optimal estimation theory (Miyazaki et 391 al., 2012a). Previous studies demonstrated the necessity for data-thinning and 392 dealiasing errors (Feng et al., 2020b; Zhang et al., 2009a). The super-observation y_{new} , 393 super-observation error r_{new} , and corresponding simulation $x_{new,i}$ of the *i*th sample 394 are calculated as follows: 395

396
$$1/r_{new}^2 = \sum_{j=1}^m 1/r_j^2$$
 (13)

397
$$y_{new} = \sum_{j=1}^{m} w_j y_j / \sum_{j=1}^{m} w_j$$
(14)

398
$$x_{new,i} = \sum_{j=1}^{m} w_j \, x_{ij} / \sum_{j=1}^{m} w_j \tag{15}$$

where *j* is the identifier of *m* observations within a super-observation grid; r_j is the observational error of the actual *j*th observation y_j ; x_{ij} is the simulated concentration 401 using the *i*th prior emission sample corresponding to the *j*th observation; and $w_j =$ 402 $\frac{1}{r_j^2}$ is the weighting factor. The super-observation error decreased as the number of 403 observations used within a super-observation increased. This method was used in our 404 previous inversions using surface-based (Feng et al., 2020b) and satellite-based (Jiang 405 et al., 2021) observations.

406 In this study, the DA window was set to one day because the model requires a longer time to integrate the emission information into the concentration ensembles (Ma et al., 407 2019). Due to the "super-observation" approach, only one assimilation is needed per 408 grid cell in one assimilation window. In addition, owing to the complexity of hourly 409 410 emissions, it is difficult to simulate hourly concentrations that match the observations well. Although a longer DA window would allow more observations to constrain the 411 emission change of one grid, the spurious correlation signals of EnKF would attenuate 412 the observation information over time (Bruhwiler et al., 2005; Jiang et al., 2021). Kang 413 414 et al. (2012) conducted OSSEs and demonstrated that owing to the transport errors and increased spurious correlation, a longer DA window (e.g. 3 weeks) would cause the 415 analysis system to blur essential emission information away from the observation. 416 Therefore, daily mean simulations and observations were used in the EnSRF algorithm 417 418 and daily emissions were optimized in this system.

419 EnKF is subject to spurious correlations because of the limited number of ensembles 420 when it is applied in high-dimensional atmospheric models, which can cause rank 421 deficiencies in the estimated background error covariance and filter divergence and 422 further degrade analyses and forecasts (Wang et al., 2020). Covariance localization is 423 performed to reduce spurious correlations caused by a finite ensemble size 424 (Houtekamer and Mitchell, 2001). Covariance localization preserves the meaningful 425 impact of observations on state variables within a certain distance (cutoff radius) but 426 limits the detrimental impact of observations on remote state variables. The localization 427 function of Gaspari and Cohn (Gaspari and Cohn, 1999) is used in this system, which

428 is a piecewise continuous fifth-order polynomial approximation of a normal distribution. 429 The optimal localization scale is related to the ensemble size, assimilation window, 430 dynamic system, and lifetime of the chemical species in the atmosphere. CO, SO2 and 431 $PM_{2.5}$ are rather stable in the atmosphere, with a lifetime of more than one day. 432 According to the average wind speed (3.3 m/s, Table 4) and length of the DA window, 433 the localization scales of CO, SO₂ and PM_{2.5} were set to 300 km. In addition, the 434 localization scales of NO₂, which is rather reactive and has a lifetime of approximately 435 10 hours in winter (de Foy et al., 2015), and PMC, which mainly comes from local 436 sources and has a short residence time in the atmosphere owing to the rapid deposition 437 rate (Clements et al., 2014; Clements et al., 2016; Hinds, 1982), were set to 150 and 438 250 km, respectively.

439 **2.2 Prior emissions and uncertainties**

Anthropogenic emissions over China were obtained from the 2016 Multi-resolution 440 Emission Inventory for China (MEIC 2016) (Zheng et al., 2018), while those over the 441 442 other regions of East Asia were obtained from the mosaic Asian anthropogenic emission inventory (MIX) (Li et al., 2017). The spatial resolutions of the MEIC and MIX 443 inventories were both $0.25^{\circ} \times 0.25^{\circ}$ and they are downscaled to match the model grid 444 spacing of 36 km. The spatial distributions of CO, SO₂, NO_x, PPM_{2.5}, and PMC 445 emissions are shown in Figure 11. The daily emission inventory, which was 446 arithmetically averaged from the combined monthly emission inventory, was directly 447 used in the EI subsystem and was employed as the prior emission of the first DA 448 window in the EI subsystem (Figure 1). During the simulations, daily emissions were 449 further converted to hourly emissions. All species emitted from area sources were 450 451 converted to hourly emissions using the same diurnal profile (Figure S2) and for the point source, we assumed that there was no diurnal change. MEIC 2012 was used as an 452 alternative a priori over China to investigate the impact of different prior emissions on 453 optimized emissions. The Model of Emissions of Gases and Aerosols from Nature 454 (MEGAN) (Guenther et al., 2012) was used to calculate time-dependent biogenic 455 emissions, which was driven by the WRF model. Biomass burning emissions were not 456

included because they have little impact across China during the study period (Zhang 457 et al., 2020). 458

During the inversion cycles, inverted emissions of different members converge 459 gradually, and the ensemble-estimated error covariance matrix is likely to be 460 underestimated. To avoid this, considering the compensation of model errors and 461 comparable emission uncertainties from one day to the next, we imposed the same 462 463 uncertainty on emissions at each DA window. As mentioned above, the optimized 464 emissions of the current DA window were transferred to the next DA window as prior emissions. The technology-based emission inventory developed by Zhang et al. (2009b), 465 using the same method as MEIC, showed that the emissions of PMC and PPM_{2.5} had 466 467 the largest uncertainties, followed by CO, and finally SO₂ and NO_x. Therefore, the uncertainties of PMC, PPM_{2.5}, CO, SO₂, and NO_x in this study were set as 40%, 40%, 468 30%, 25%, and 25%, respectively. However, previous studies have shown that inversely 469 estimated CO and PMC emissions can exceed 100% higher than the bottom-up 470 471 emissions (MEIC) in certain areas (Feng et al., 2020b; Ma et al., 2019). Therefore, according to the extent of underestimation, we set an uncertainty of 100% for both the 472 473 CO and PMC emissions at the beginning of the three DA windows to quickly converge the emissions. Mean emission analysis is generally minimally sensitive to the 474 uncertainty setting in the assimilation cycle method (Feng et al., 2020; Gurney et al., 475 2004; Miyazaki et al., 2012a) as the inversion errors of the current window can be 476 transferred to the next window for further optimization (Section 4.3). 477

478

2.3 Observation data and errors

479 Hourly averaged surface CO, SO₂, NO₂, O₃, PM_{2.5}, and PM₁₀ observations from 1504 national control air quality stations were assimilated into this system, which were 480 obtained from the Ministry of Ecology and Environment of the People's Republic of 481 China (http://106.37.208.233:20035/, last access: 25 June 2020). These sites are 482 distributed over most of central and eastern China and become denser near metropolitan 483 areas (see Figure 2). To ensure data quality, value-range checks were performed to 484 eliminate unrealistic or unrepresentative observations and only the observations within 485

486 the subjectively selected threshold range were assimilated (Table 2). In additionally, a time-continuity check was performed to eliminate gross outliers and sudden anomalies 487 using the function of $max(|O(t) - O(t \pm 1)|) \le f(t)$, where O(t) and $O(t \pm 1)$ 488 represent observations at time t and $t \pm 1$, respectively, and $f(t) = T_a + T_b \times O_t$. 489 490 This means that the concentration difference between time t and time t+1 and t-1 should be less than f(t). T_b was fixed at 0.15 and the section of T_a is given in Table 2, 491 492 which was determined empirically according to the time series change of concentration at each site. To avoid potential cross-correlations, we assimilated PM_{2.5} and PMC. 493 Additionally, in the EI subsystem, the observations within each city were averaged to 494 reduce the data density, reduce the error correlation, and increase spatial representation 495 496 (Houtekamer and Mitchell, 2001; Houtekamer and Zhang, 2016). Finally, 336 city sites were available across mainland China, in which data from 311 cities were selected for 497 498 assimilation and the remaining 25 were selected for independent validation (Figure 2). In the IA subsystem, owing to the small horizontal correlation scale (Figure 3), all site 499 observations were assimilated to provide a good IC for the next emission inversion to 500 obtain more extensive observation constraints. 501

502 The observation error covariance matrix (**R**) includes both the measurement and 503 representation errors. The measurement error ε_0 is defined as follows:

504 $\varepsilon_0 = ermax + ermin \times \Pi_0$ (16)

where *ermax* is the base error and Π_0 denotes the observed concentration. These parameters for different species are listed in Table 2 and were determined according to Chen et al. (2019), Feng et al. (2018), and Jiang et al. (2013b).

508 The representative error depends on the model resolution and characteristics of the 509 observation locations, which were calculated using the equations of Elbern et al. (2007), 510 defined as follows:

511
$$\varepsilon_r = \gamma \varepsilon_0 \sqrt{\Delta l/L}$$
 (17)

512 where γ is a tunable parameter (here, $\gamma=0.5$), Δl is the grid spacing (36 km), and L

513 is the radius (3 km for simplification) of the influence area of the observation. The total 514 observation error (r) was defined as follows:

$$r = \sqrt{\varepsilon_0^2 + \varepsilon_r^2} \tag{18}$$

Parameter	CO mg m ⁻³	SO ₂ µg m ⁻³	NO ₂ μg m ⁻³	O ₃ µg m ⁻³	PM _{2.5} μg m ⁻³	PMC μg m ⁻³
value-range	0.1-12	1-800	1-250	1-250	1-800	1-900
time-continuity (T_a)	2.5	160	70	80	180	180
ermax	0.05	1	1	1	1.5	1.5
ermin	0.5%	0.5%	0.5%	0.5%	0.75%	0.75%

516 **Table 2**. Parameters of quality control and measurement error

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515

518 **3 Experimental design**

RAPAS inversions were conducted according to the procedure and settings described 519 520 in Section 2. December is one of the months with the most severe air pollution, whereas July is one of the least polluted months in China. Therefore, this study mainly tested 521 the performance of the RAPAS system over these two months. For December, the IA 522 subsystem was run from 26 November to 31, 2016, with a 6-hour interval cycling 523 524 assimilation to optimize ICs (ICDA). A better IC at 0000 UTC on 1 December could be obtained by a five-day high-frequency cycling assimilation and atmospheric mixing. 525 The EI subsystem was then run for December 2016 with a one-day assimilation window 526 to optimize emissions (EMDA). In July, the system operated identically to that of 527 December. It should be noted that owing to the stronger atmospheric oxidation, the 528 lifetime of NO₂ in July was significantly shorter than that in December; thus, we 529 adopted a smaller localization scale for NO₂ (80 km). Both assimilation experiments 530 used the combined prior emission inventories of 2016, as described in Section 2.2, and 531 532 the emission base year coincided with the research stage. An Observing Systems Simulation Experiment (OSSE) was conducted to evaluate the performance of the 533

RAPAS system, which has been widely used in previous assimilation systems 534 development (Daley, 1997). In the OSSE experiment, we used the MEIC 2016 535 inventory as a "true" emission and reduced by 30% over mainland China as a prior 536 emission. The simulations were performed using the "true" emission and sampled 537 according to the locations and times of the real observations used as artificial 538 observations. The observation errors were the same as those in EMDA. To evaluate the 539 IC improvements from the IA subsystem, an experiment without 3DVAR (NODA) was 540 conducted with the same meteorological fields and physical and chemistry 541 parameterization settings as those of the ICDA. To evaluate the posterior emissions of 542 the EI subsystem, two parallel forward modelling experiments were performed for 543 December 2016: a control experiment (CEP) with prior (MEIC 2016) emissions and a 544 validation experiment (VEP) with posterior emissions. Both experiments used the same 545 IC at 0000 UTC on December 01 generated through the IA subsystem. The only 546 difference between CEP and VEP were emissions. Table 3 summarizes the different 547 emission inversion experiments conducted in this study. 548

549 To investigate the robustness of our system, seven sensitivity tests (from EMS1 to 550 EMS7; see Table 3) were performed. These experiments were all based on EMDA. EMS1 used MEIC 2012 as the original prior emission in China, aiming to investigate 551 the impact of different prior inventories on the estimates of emissions. The other 552 experiments (EMS2–5) aimed to test the impact of different prior uncertainty settings, 553 in which the prior uncertainties were reduced by -50% and -25%, and increased by 25% 554 and 50%, respectively. EMS6 aimed to evaluate the impact of observation errors on 555 emission estimates, in which all observation errors are magnified twice. EMS7 aimed 556 557 to evaluate the impact of IC optimization of the first window on emission estimates, in which the ICs were taken from a five-day spin-up simulation. Eight forward modelling 558 experiments (VEP1, VEP2, ..., VEP7) were also performed with the posterior 559 emissions of EMS1 to EMS7 to evaluate their performance. 560

561

Ехр. Туре	Exp. Name	Period	IC of the first DA Window	ICs of the subsequent DA window	Emission
Assimilation	EMDA	1–31 December	0000 UTC on December 1, taken from ICDA	Forecast with posterior emissions in the previous window	MEIC 2016 for December (the first DA window), optimized emissions of the previous window (other DA windows)
	OSSE	1–31 December	Same as EMDA	Same as EMDA	Same as EMDA but with a decrease of 30% for CO, SO ₂ , NO _x , PPM _{2.5} , and PMC
	EMS1	1–31 December	Same as EMDA	Same as EMDA	Same as EMDA but for EMIC 2012
	EMS2-5	1–31 December	Same as EMDA	Same as EMDA	Same as EMDA but with a \pm 25% or \pm 50% of default uncertainty
Sensitivity	EMS6	1–31 December	Same as EMDA	Same as EMDA	Same as EMDA but with a +100% of default observation errors
	EMS7	1–31 December	0000 UTC on December 1, taken from ICNO	Same as EMDA	Same as EMDA

562 Table 3. Emission inversion and sensitivity experiments conducted in this study

563

564 **4 Results**

565 4.1 Evaluations

566 4.1.1 Simulated meteorological fields

567 In the RAPAS system, the inversion approach attributes all biases between the 568 simulated and observed concentrations to emissions. Meteorological fields dominate 569 the physical and chemical processes of air pollutants in the atmosphere, and thus their

simulation accuracy would significantly affect the estimates of emissions in this study. 570 To quantitatively evaluate the performance of the WRF simulations, the mean bias 571 (BIAS), root mean square error (RMSE), and correlation coefficient (CORR) were 572 calculated against the surface meteorological observations measured at 400 stations and 573 the planetary boundary layer height (PBLH) was calculated using the sounding data at 574 92 sites. Surface observations were obtained from the National Climate Data Center 575 integrated surface database (http://www.ncdc.noaa.gov/oa/ncdc.html, last access: 25 576 577 October 2021) and sounding data were obtained from the website of the University of Wyoming (http://weather.uwyo.edu/upperair/sounding.html, last access: 10 March 578 2022). The sounding data had a 12 hour interval. The observed PBLH was calculated 579 using sound data via the bulk Richardson number method (Richardson et al., 2013). 580 581 The spatial distribution of meteorological stations is shown in Figure 2. The simulated temperature at 2 m (T2), relative humidity at 2 m (RH2), wind speed at 10 m (WS10), 582 and PBLH from 26 November to 31 December 2016 were evaluated against the 583 observations. Table 4 summarizes the statistical results of the evaluation of the 584 585 simulated meteorological parameters. Overall, T2, RH2 and PBLH were slightly underestimated, with biases of -0.1 °C, -3.8%, and -41.1 m, respectively. CORRs were 586 approximately 0.98 for T2, 0.94 for RH2, and 0.90 for PBLH, showing good 587 consistency between the observations and simulations. WS10 was overestimated, with 588 a bias of 0.7 m/s and an RMSE of 0.8 m/s, but were better than the simulations from 589 many previous studies (Chen et al., 2016; Jiang et al., 2012a; Jiang et al., 2012b). 590 Therefore, the WRF can generally reproduce meteorological conditions sufficiently in 591 terms of their temporal variation and magnitude over China, which is adequate for our 592 593 inversion estimation.

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Table 4. Statistics comparing the simulated and observed 10-m wind speed (WS10), 2m temperature (T2), and 2-m relative humidity (RH2), and planetary boundary layer
height (PBLH).

Variable Met.	No. of sites	Mean Obs.	Mean Sim.	BIAS	RMSE	CORR
WS10 (m/s)	400	2.6	3.3	0.7	0.8	0.72
T2 (°C)	400	2.9	2.8	-0.1	0.7	0.98
RH2 (%)	400	66.3	62.6	-3.8	5.2	0.94
PBLH (m)	92	267.5	226.4	-41.1	50.4	0.90

* BIAS, mean bias; RMSE, root mean square error; CORR, correlation coefficient

602 4.1.2 Initial conditions

Figure 4 shows an evaluation of the analyzed concentrations of the six species against 603 604 surface observations. For comparison, the evaluations of the simulations without 3DVAR (NODA) are also shown in Figure 4. The simulations of the NODA experiment 605 (red dots) are scattered on both sides of the central line, as large systematic biases 606 607 remain across many measurement sites. Conversely, the ICDA experiment (blue dots) showed a much better agreement with the observations than those from NODA. The 608 statistics show that there are large systematic biases in the NODA simulations, with 609 large RMSEs and small CORRs for all species, particularly for CO and PMC. After the 610 611 assimilation of surface observations, the RMSE of CO decreased to 0.7 mg m⁻³, and those of SO₂, NO₂, O₃, PM_{2.5}, and PMC decrease to 22.0, 12.0, 9.6, 20.5, and 19.6 µg 612 m⁻³, respectively, with respective reductions of 50.0%, 73.1%, 61.0%, 64.7%, 69.5%, 613 and 60.8% compared to those of the NODA (Table 5). The CORRs of ICDA increased 614 by 290.0%, 291.3%, 55.4%, 87.2%, 130.0%, and 214.8% to 0.78, 0.90, 0.87, 0.88, 0.92, 615 and 0.85, respectively. These statistics indicate that the ICs of the ground level 616 improved significantly. However, owing to the lack of observations, we still do not 617 know the simulation bias in the upper-middle boundary layer. Although concentrations 618 at high altitudes can be constrained by ground-based observations through vertical 619

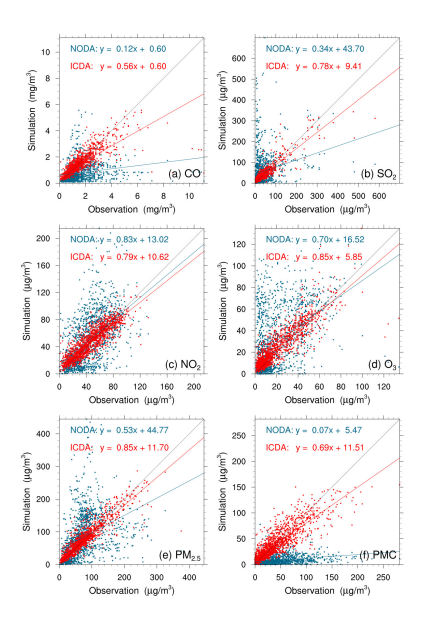


Figure 4. Scatter plots of simulated versus observed (a) CO, (b) SO₂, (c) NO₂, (d) O₃,
(e) PM_{2.5}, and (f) PMC mass concentrations at 0000 UTC on December 1 initializations
from the background (red) and analysis (blue) fields.

Species	Exp. Name	Mean Obs.	Mean Sim.	BIAS	RMSE	CORR
СО	NODA	1.5	0.8	-0.7	1.4	0.20
	ICDA	1.5	1.5	-0.1	0.7	0.78
SO_2	NODA	36.3	56.0	19.7	81.7	0.23
302	ICDA	50.5	37.8	1.5	22.0	0.90
NO ₂	NODA	45.8	51.1	5.3	30.8	0.56
1102	ICDA		47.0	1.1	12.0	0.87
O_3	NODA	20.5	30.8	10.4	27.2	0.47
03	ICDA		23.3	2.8	9.6	0.88
PM _{2.5}	NODA	70.9	82.2	11.3	67.3	0.40
F 1 V1 2.5	ICDA	/0.9	71.8	0.9	20.5	0.92
PMC	NODA	43.5	8.5	-35.0	50.0	0.27
r MC	ICDA	43.3	41.6	-1.9	19.6	0.85

Table 5. Comparisons of the surface CO, SO₂, NO₂, O₃, PM_{2.5}, and PMC mass concentrations from the control and assimilation experiment against observations aggregated over all analysis times. CO unit: mg m⁻³; others units: μ g m⁻³.

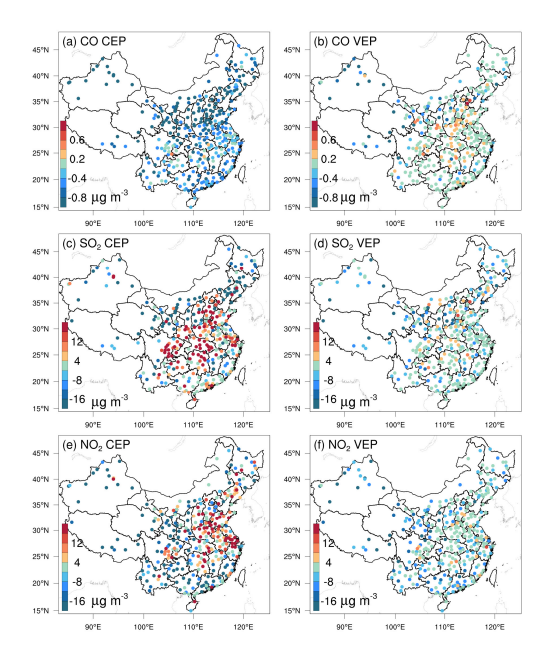
* BIAS, mean bias; RMSE, root mean square error; CORR, correlation coefficient

634 4.1.3 Posterior emissions

635 Owing to the mismatched spatial scales, it is difficult to directly evaluate the optimized emissions against observations. Generally, we indirectly validated the optimized 636 emissions by comparing the forward simulated concentrations using the posterior 637 emissions against atmospheric measurements (e.g., Jiang et al., 2014; Jin et al., 2018; 638 Peters et al., 2007). Figure 5 shows the spatial distributions of the mean biases between 639 the gaseous pollutants simulated using prior and posterior emissions and assimilated 640 observations. In the CEPs, for each species, the distribution of biases was similar to the 641 642 increments in background fields constrained through 3DVAR, as shown in Figure S3. 643 For example, almost all sites had large negative biases for CO, while for SO₂ and NO₂, positive biases were mainly distributed over the North China Plain (NCP), Yangtze 644 River Delta (YRD), Sichuan Basin (SCB), and Central China and negative biases were 645 distributed over remaining areas. After constraining with observations, the biases of all 646

three gaseous air pollutants were significantly reduced at most sites. For CO, the biases 647 at 62% of the sites decreased to absolute values less than 0.2 mg m⁻³ and for SO₂ and 648 NO₂, the biases at 52% and 47% of the sites were within $\pm 4 \ \mu g \ m^{-3}$. However, large 649 negative biases were still observed in western China, indicating that the uncertainties 650 of the posterior emissions are still large in western China, which may be attributed to 651 the large biases in prior emissions and the relatively limited observations. Overall, the 652 statistics show that there are different levels of improvement at the 311 assimilation 653 sites of 92%, 85%, and 85% for CO, SO₂, and NO₂, respectively. The small number of 654 sites with worse performance may be related to over-adjusted emissions by EI or 655 contradictory adjustments caused by opposite biases in adjacent areas. 656

Table 6 lists the statistical results of the evaluations averaged over the whole mainland 657 of China. For CO, the mean bias was -0.8 mg m⁻³ with the prior emissions, while it 658 substantially reduced to -0.1 mg m⁻³ (reduction rate of 89.6%) when simulating with 659 the posterior emissions. Additionally, the RMSE decreased by 48.1% from 1.08 to 0.56 660 mg m⁻³, and the CORR increased by 76.1% from 0.46 to 0.81. For SO₂ and NO₂, the 661 662 regional mean biases slightly increased as the positive/negative biases among different sites might be offset. However, the RMSEs decreased to 17.7 and 12.3 µg m⁻³, 663 respectively, which were 58.3% and 50.8% lower than those of CEPs, and the CORRs 664 increased by 125.6% and 35.4%, both reaching up to 0.88, indicating that EI 665 significantly improved the NO_x and SO₂ emission estimates. 666

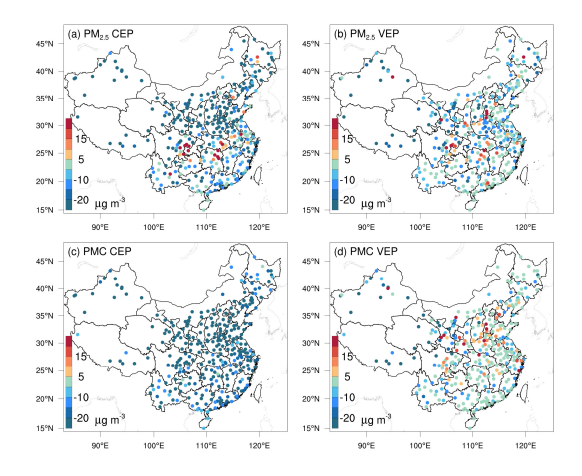


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Figure 5. Spatial distribution of the BIAS of the simulated (a, b) CO, (c, d) SO₂, and (e, f) NO₂ with prior (left, CEP) and posterior (right, VEP) emissions. CO unit: mg m⁻ 3 ; SO₂ and NO₂ units: μ g m⁻³.

Figure 6 shows the spatial distributions of the mean biases of simulated $PM_{2.5}$ and PMCevaluated against assimilated observations. Similarly, the CEP simulations did not perform well. There were widespread underestimations across the country, with mean biases of -24.0 and -32.4 µg m⁻³. After data assimilation, the performance of the VEP simulations significantly improved. The biases decreased by 72.1% and 90.4% to -6.7

and -3.1 μ g m⁻³, the RMSEs decreased by 41.2% and 40.7% to 29.6 and 24.6 μ g m⁻³, 676 and the CORRs increased by 35.9% and 176.0% to 0.87 and 0.69 for PM_{2.5} and PMC, 677 respectively. Overall, 89.6% and 97.2% of the assimilation sites were improved for 678 PM_{2.5} and PMC, respectively. However, compared with the results for the three gaseous 679 pollutants, there were sites with large biases scattered throughout the entire domain. In 680 addition to the potential over-adjusted or contradictory adjustments of emissions as in 681 the three gas species, the sites with large biases may be related to the complex 682 precursors and complex homogeneous and heterogeneous chemical reactions and 683 transformation processes of secondary PM_{2.5}, and the fact that we did not simulate the 684 time variation of dust blowing caused by wind speed for PMC owing to the lack of land 685 cover data that is compatible with the CMAQ dust module and agricultural activity data 686 to identify dust source regions. 687





689

Figure 6. Same as in Figure 5 but for PM_{2.5} and PMC.

Figures 7 and 8 show the spatial distributions of the biases calculated against 690 independent observations for the five species. With posterior emissions, the decreasing 691 ratios of RMSEs ranged from 26.7%-42.0% and the CORRs increased by 13.7-59.0% 692 to 0.62–0.87. Overall, the biases at the independent sites are similar or slightly worse 693 than those at the assimilated sites, which is reasonable as the closer the independent 694 sites are to the assimilated site, the more constraints of observation information can be 695 obtained and the more significant the improvements in the optimized state variables of 696 697 the model. For example, generally, the transmission distance of NO₂ is relatively short and remote cities with small emission correlations to the cities with assimilated 698 observations are relatively less constrained, resulting in only a 26.7% decrease in the 699 RMSE. 700

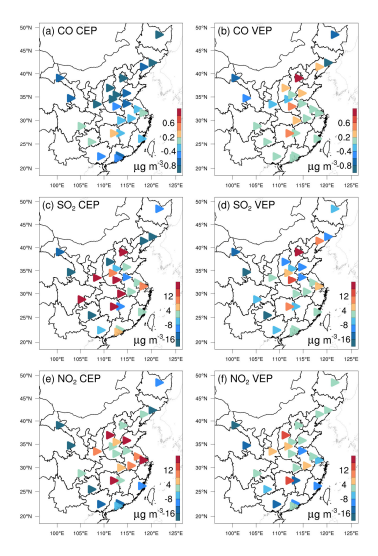




Figure 7. As in Figure 5 but for the independent validation.

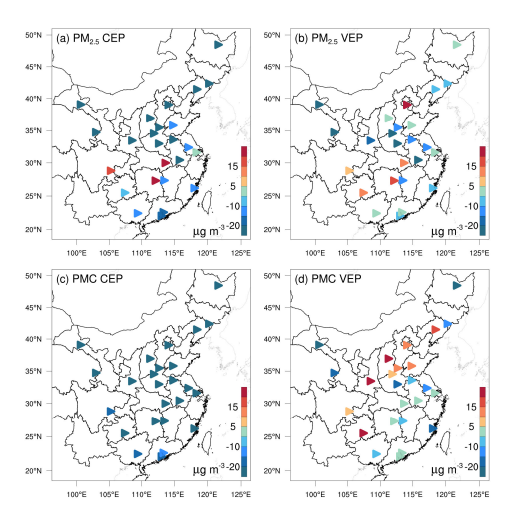




Figure 8. As in Figure 6 but for the independent validation.

Comparing our results with those of previous studies, Tang et al. (2013) inverted CO 705 emissions over Beijing and the surrounding areas and obtained comparable 706 improvements (Table 6) in the RMSE (37-48% vs. 30-51%) and CORR (both studies 707 ~ 0.81); however, we decreased the biases by 90–97%, which is much greater than their 708 48-64% reductions. Additionally, Chen et al. (2019) showed that the RMSE of 709 simulated SO₂ with updated SO₂ emissions decreased by 4.2-52.2% for different 710 regions, and the CORR only increased to 0.69 at most. These improvements are smaller 711 than those obtained in this study, which may be due to the insufficient adjustment of 712 emissions caused by the underestimated ensemble spread through the inflation method. 713 The better performance in this study may be related to our inversion process, which 714 causes the optimized emissions of the current DA window to propagate to the next DA 715 window for further correction. 716

717 **Table 6**. Statistics comparing the pollution concentrations from the simulations with

718 prior (CEP) and posterior (VEP) emissions against assimilated and independent

Species	Mean	Mean Sim.		BIAS		RMSE		CORR	
	Obs.	CEP	VEP	CEP	VEP	CEP	VEP	CEP	VEP
	Against assimilated observations								
СО	1.43	0.66	1.36	-0.77	-0.08	1.08	0.56	0.46	0.81
SO_2	32.5	34.4	28.4	1.9	-4.1	42.4	17.7	0.39	0.88
NO ₂	43.8	40.8	39.0	-2.9	-4.8	25.0	12.3	0.65	0.88
PM _{2.5}	77.0	53.1	70.3	-24.0	-6.7	50.3	29.6	0.64	0.87
PMC	40.5	8.1	37.5	-32.4	-3.1	41.5	24.6	0.25	0.69
		Agains	st indepe	endent ol	bservatio	ons			
СО	1.54	0.79	1.52	-0.75	-0.02	1.15	0.72	0.59	0.82
SO ₂	40.6	39.2	37.3	-1.3	-3.2	44.3	27.2	0.57	0.87
NO ₂	50.2	50.0	47.5	-0.3	-2.7	21.7	15.9	0.73	0.83
PM _{2.5}	91.5	64.6	84.1	-26.9	-7.4	64.1	37.2	0.62	0.87
РМС	42.0	9.2	40.4	-32.8	-1.6	39.3	26.6	0.39	0.62

observations, respectively. CO unit: mg m⁻³; others units: μ g m⁻³.

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* BIAS, mean bias; RMSE, root mean square error; CORR, correlation coefficient

721 **4.1.4 Uncertainty reduction**

The uncertainty reduction rate (UR) is an important quantity to evaluate the performance of RAPAS and the effectiveness of *in-situ* observations (Chevallier et al., 2007; Jiang et al., 2021; Takagi et al., 2011). Following Jiang et al. (2021), the UR was calculated as

726
$$UR = (1 - \frac{\sigma_{posterior}}{\sigma_{prior}}) \times 100$$
(19)

where $\sigma_{posterior}$ and σ_{prior} are the posterior and prior uncertainties, respectively, calculated using the standard deviations of the prior and posterior perturbations (Text S2). Table 7 shows the URs averaged in each province and mainland China. URs varied with species as they are closely related to the magnitude settings of prior uncertainties (Jiang et al., 2021). The URs of PPM_{2.5} and PMC were the most effective while the UR of NO_x emissions was the lowest. For mainland China overall, uncertainties were reduced by 44.4%, 45.0%, 34.3%, 51.8%, and 56.1% for CO, SO₂, NO_x, PPM_{2.5}, and 734 PMC, respectively. For one species, URs varied across provinces. URs are usually related to observation coverage, which means that the more observation constraints 735 there are, the more URs decrease. Additionally, URs may also be related to emission 736 distributions. Generally, URs were more significant in the provinces where 737 observations and emissions were both relatively concentrated (e.g. Tibet), while they 738 were much lower where the emissions were scattered or relatively uniform, but the 739 observations were only in large cities, even if there were many more observations than 740 741 in other provinces.

Table 7. Time-averaged posterior emission uncertainty reduction (%) indicated by the
standard deviation reduction of total emissions per province calculated by prior and
posterior ensembles.

Province	со	SO ₂	NO _x	PPM _{2.5}	РМС	
Mainland	44.4	45.0	34.3	51.8	56.1	
Shanghai	16.9	16.7	20.8	24.7	18.5	
Jiangsu	17.7	25.3	29.3	34.1	52.3	
Zhejiang	24.7	13.3	17.9	42.4	31.4	-70
Anhui	20.1	52.7	39.1	58.1	40.9	
Shandong	32.1	30.0	20.3	53.7	26.7	-65
Beijing	28.2	6.2	37.0	43.3	31.4	
Tianjin	20.0	7.0	21.4	41.3	17.8	60
Hebei	29.5	40.2	28.8	56.0	30.3	
Shanxi	38.4	37.9	22.5	55.3	35.0	-55
Neimenggu	30.1	45.8	40.4	37.6	52.8	50
Henan	27.4	16.1	21.9	53.7	30.8	50
Hunan	36.0	27.7	34.4	16.9	41.6	-45
Hubei	30.8	16.6	26.0	46.4	46.5	45
Jiangxi	20.9	28.4	29.4	47.0	46.7	-40
Guangdong	31.2	14.9	41.1	53.1	46.4	40
Guangxi	22.6	13.9	42.5	48.1	55.2	-35
Fujian	9.9	8.1	31.9	31.6	49.2	55
Hainan	0.6	0.5	4.5	0.7	23.3	-30
Liaoning	35.6	34.6	19.0	33.9	54.0	
Heilongjiang	29.9	27.7	17.4	42.0	65.2	-25
Jilin	27.9	44.5	18.7	42.0	42.8	
Shaanxi	41.3	13.2	29.8	47.9	43.1	-20
Gansu	24.8	36.1	33.7	46.3	56.4	
Xinjiang	38.3	27.9	20.2	46.3	66.5	-15
Qinghai	53.9	25.8	27.3	46.0	57.9	
Ningxia	47.0	36.6	17.6	38.0	30.1	-10
Sichuan	29.4	25.0	39.5	61.1	46.5	
Chongqing	5.7	8.2	8.8	12.7	13.8	-5
Guizhou	14.4	16.4	26.6	40.3	38.2	
Yunnan	38.3	29.9	31.4	40.1	55.9	-0
Tibet	30.2	0.5	52.8	67.3	73.2	

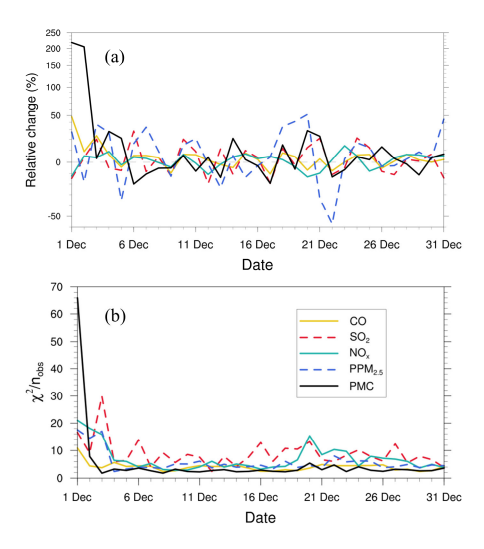
746 4.1.5 Evaluation using chi-squared statistics

To diagnose the performance of the EnKF analysis, chi-squared (χ^2) statistics were calculated, which are generally used to test whether the prior ensemble mean RMSE with respect to the observations is consistent with the prior "total spread" (square root of the sum of ensemble variance and observation error variance). Following Zhang et al. (2015), for the *t*-th window, χ^2 is defined as:

752
$$\chi_t^2 = (\mathbf{y} - \mathbf{H}\overline{\mathbf{X}}\overline{\mathbf{b}})^T (\mathbf{H}\mathbf{P}^b\mathbf{H}^T + \mathbf{R})^{-1} (\mathbf{y} - \mathbf{H}\overline{\mathbf{X}}\overline{\mathbf{b}})$$
(20)

Figure 9 shows the time series of the relative changes between the prior and posterior 753 emissions and the χ^2 statistics. There were relatively large adjustments in emissions in 754 the first three windows, especially for the PMC. Subsequently, the five species reached 755 a more optimal state with successive emission inversion cycles. The χ^2 statistics showed 756 similar variation characteristics as the daily changes in emissions. The χ^2 value was 757 slightly greater than 1, indicating that the uncertainties from the error covariance 758 759 statistics did not fully account for the error in the ensemble simulations. A similar result was reported by Chen et al. (2019). Further investigations should be conducted to 760 generate larger spreads by accounting for the influence of model errors. As we imposed 761 the same uncertainty of prior emissions at each DA window to partially compensate for 762 the influence of model errors, χ^2 statistics showed small fluctuations, indicating that the 763 system updated emissions consistently and stably. 764

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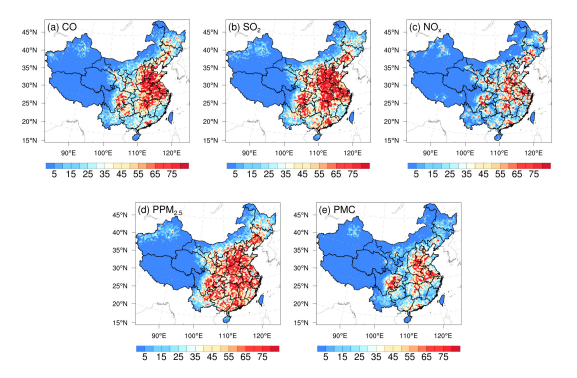


766

Figure 9. Relative changes (a) in posterior emission estimates of CO, SO₂, NO_x, PPM_{2.5}, and PMC and χ^2 statistics (b) of these state vectors in each window.

769 **4.1.6 Evaluation using OSSE**

Figure 10 shows the spatial distribution of the error reduction in the posterior emissions 770 of the five species. After inversion, in most areas, the emission errors were reduced by 771 772 more than 80%, especially in the central and eastern regions with dense observation sites, while in remote areas far away from cities, due to the sparse observation sites, the 773 emission errors were still not well adjusted. Overall, the error reduction rates of CO, 774 SO₂, NO_x, PPM_{2.5}, and PMC were 78.4%, 86.1%, 78.8%, 77.6%, and 72.0%, 775 respectively, indicating that with the in-situ observations in China, RAPAS can 776 significantly reduce emission errors and thus showed good performance in emission 777 estimates. 778



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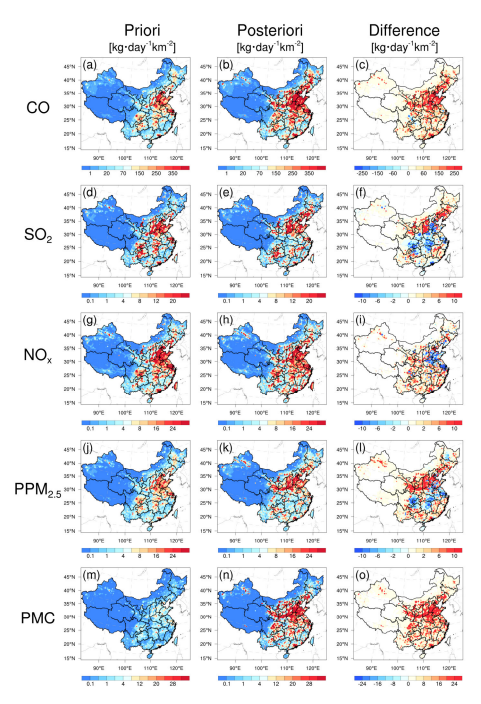
Figure 10. Spatial distribution of the error reduction (%) of posterior emissions in theOSSE.

782 4.2 Inverted emissions

Figure 11 shows the spatial distribution of temporally averaged prior and posterior 783 emissions and their differences in emissions in December 2016. It should be noted that 784 emissions outside China were masked; as the observation sites were limited to China in 785 this study, there was a slight change in the emissions outside China. Higher emissions 786 were mainly concentrated in central and eastern China, especially in the NCP, YRD, 787 and Pearl River Delta, and lower emissions occurred across Northwest and Southern 788 China. Compared with the prior emissions, posterior CO emissions were considerably 789 790 increased across most areas of mainland China, especially in northern China, with an overall increase of 129%. A notable underestimation of prior emissions was also 791 792 confirmed by inversion estimations (Feng et al., 2020b; Tang et al., 2013; Wu et al., 2020) and model evaluations (Kong et al., 2019b) in previous studies. For SO₂, the 793 emissions increased mainly in Northeast China, Shanxi, Ningxia, Gansu, Fujian, 794 Jiangxi, and Yunnan provinces. In SCB, Central China, YRD, and part of the NCP, 795

emissions were significantly reduced. The national total SO₂ emissions increased by 796 20%. For NO_x, although the increment of national total emissions was small 797 798 (approximately 5%), there were large deviations. The emissions in NCP and YRD were reduced, whereas the emissions in most cities in other regions increased. The changes 799 in the emission of PPM_{2.5} were similar to those of SO₂. Compared with the prior 800 801 emissions, the posterior PPM_{2.5} emissions decreased over central China, SCB, and YRD, whereas those in southern and northern China increased, especially in Shanxi, Shaanxi, 802 Gansu, and southern Hebei provinces. Overall, the relative increase was 95%. For PMC, 803 the posterior emissions were increased over all of mainland China, with a national mean 804 relative increase of 1045%. Larger emission increments mainly occurred in areas with 805 significant anthropogenic emissions of CO and PPM_{2.5}, indicating that the large 806 underestimation of PMC emissions in the prior inventory may be mainly attributed to 807 the underestimations of anthropogenic activities. The absence of natural dust is another 808 reason, as the wind-blown dust scheme was not applied in this study. Overall, PM10 809 emissions (PPM_{2.5}+PMC) increased by 318%. If we assume that all the increments in 810 811 PM₁₀ emissions are from natural dust, that means the contribution of natural dust accounted for 75% of total PM₁₀ emissions, which is consistent with the source 812 apportionment of PM₁₀ of 75% in Changsha in Central China (Li et al., 2010). Large 813 814 PMC emission increments were also reported by Ma et al. (2019).

Detailed estimations of posterior emissions and relative changes compared to prior 815 emissions in each province and mainland China are given in Table S1. The evaluation 816 817 results for July showed that the emission uncertainty could still be significantly reduced and the performance of the system in July was comparable to that in December (Table 818 S2). Additionally, the seasonal variation in emissions was well reflected (Figures S4 819 and S5), which means that our system performed well at different times of the year. 820 Note that the differences, excluding PMC, between the prior and posterior emissions 821 mainly reflect the deficiencies of the prior emissions as the times of the prior emissions 822 and observations were consistent in this study. 823



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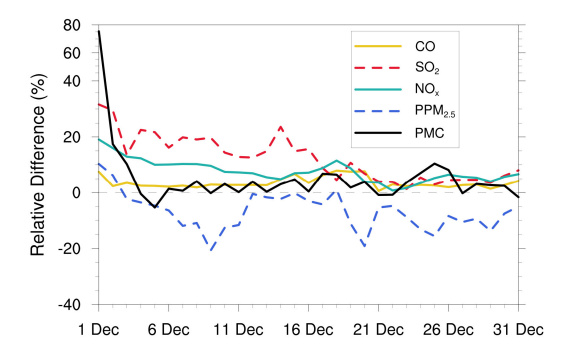
Figure 11. Spatial distribution of the time-averaged prior emissions (left column, MEIC
2016), posterior emissions (middle column), and differences (right column, posterior
minus prior).

828 4.3 Sensitivity tests

829 4.3.1 Impact of prior inventories

830 Various prior inventories have shown considerable differences in space allocation and

emission magnitudes. Inversion results can be sensitive to a priori emissions if the 831 observations are insufficient (Gurney et al., 2004; He et al., 2018). MEIC 2012 was 832 833 used as an alternative a priori in EMS1 to investigate the impact of different prior emissions on posterior emissions. Figure 12 shows the time series of the relative 834 differences in the daily posterior emissions of the five species between the EMDA (base) 835 and EMS1 experiments. Overall, the differences between the two posterior emissions 836 gradually decreased over time. At the beginning, the differences in the CO, SO₂, NO_x, 837 PPM_{2.5}, and PMC between the two inventories (i.e. MEIC 2012 vs. MEIC 2016) were 838 17.5%, 114.5%, 30.8%, 46.0%, and 72.0%, respectively, compared to 2.5%, 4.5%, 839 4.5%, -8.9%, and 3.0% in the last ten days. In addition, the species with larger emission 840 differences at the beginning took a longer time (i.e. more DA steps) to achieve 841 convergence. The quick convergence of PMC emissions was attributed to the large prior 842 uncertainty of 100% used in the first three DA windows. In contrast to the other species, 843 there were significant negative deviations in PPM_{2.5} emissions between the two 844 experiments. This may be due to the positive deviations in the precursors of PM_{2.5} (i.e., 845 846 SO_2 and NO_x), which lead to a larger amount of secondary production. The PPM_{2.5} emissions will be reduced to balance the total PM2.5. We compared the PM2.5 847 concentrations simulated by the two optimized inventories and found that they were 848 almost the same (Figure S6). Overall, this indicates that observations in China were 849 sufficient to infer emissions and that our system was robust. Meanwhile, the monthly 850 posterior emissions shown in Section 4.2 were still underestimated to a certain extent. 851



852

Figure 12. Relative differences in CO, SO₂, NO_x, PPM_{2.5}, and PMC emissions (%, the
ratio of absolute difference to EMDA) between the EMDA and EMS1 experiments.

4.3.2 Impact of prior uncertainties settings

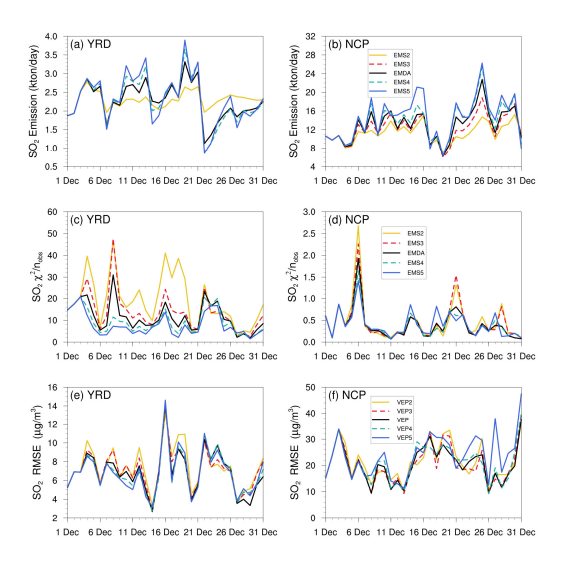
The uncertainty of prior emissions determines how closely the analysis is weighted 857 towards the background and observations; however, information about prior 858 uncertainties is generally not readily available. To evaluate the possible influence of 859 prior uncertainties on the optimized emissions, we increased/reduced the uncertainties 860 after three days of cycling, namely starting at 0000 UTC, 3 December, by 25% and 50 % 861 in EMS2 (-50%), EMS3 (-25%), EMS4 (+25%), and EMS5 (+50%), respectively. Table 862 8 summarizes the emission changes with different prior uncertainty settings in the 863 864 EMS2–5 experiments. To better understand the response of the system to the emission uncertainty settings, Figure 13 illustrates the time series of SO₂ emission changes, Chi-865 square statistics, and RMSEs of simulated SO₂ with emissions updated in the EMDA 866 and EMS2-5 experiments over the YRD and NCP (Figure 2). Compared with the 867 EMDA, when the uncertainties decreased (increased), the emissions of the five species 868 decreased (increased) accordingly. This is because the posterior emissions of the five 869 species were larger than the prior emissions and, as shown in Figure 13a-d, larger 870

uncertainty will lead to faster convergence, resulting in larger posterior emissions. It 871 can also be seen from Figure 13 that a faster convergence will reduce the RMSE of the 872 simulated concentration with the posterior emissions in the early stage of the 873 experiment; however, in the later stage of the experiment, there were no significant 874 differences in the RMSE and Chi-square statistics among the different experiments. 875 However, day-to-day changes in emissions also cause slight fluctuations. In addition, 876 when greater uncertainties are set, the day-to-day changes in emissions are more drastic, 877 resulting in a larger RMSE, as shown in the NCP. Moreover, the significant day-to-day 878 variations in the estimated emissions may not be in line with the actual situation. Owing 879 to the spatial-temporal inhomogeneity of emissions, the differences in Chi-square 880 statistics between the YRD and NCP show that it may be necessary to apply different a 881 priori uncertainties according to different regions (Chen et al., 2019). Therefore, when 882 using an EnKF system for emission estimation, error setting must be carefully executed. 883 Overall, the uncertainties chosen in EMDA aim to minimize the deviation of the 884 concentration fields and maintain the stability of the inversion. 885

Table 8. Relative differences in CO, SO₂, NO_x, PPM_{2.5} and PMC emissions (%, the ratio of absolute difference to EMDA) between the EMDA and EMS2-5 experiments.

Species	EMS2	EMS3	EMS4	EMS5
СО	-8.6	-4	3	5.2
SO_2	-14	-5.7	3.6	6.8
NO _x	-6.5	-3	2.8	4.5
PPM _{2.5}	-16.5	-7.8	4.6	8.7
PMC	-18.5	-8.2	7.3	13.1

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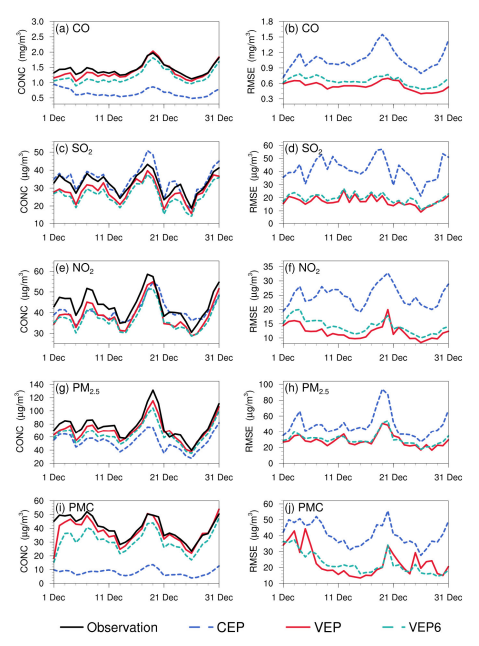
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Figure 13. Time-series of SO₂ emission changes, Chi-square statistics, and RMSE of
simulated SO₂ with updated SO₂ emissions in the EMDA and EMS2-5 experiments
over the YRD and NCP.

893 **4.3.3 Impact of observation error settings**

Observation errors are another factor that determine the relative weights of the observations and background in the analysis. A proper estimate of the observation error is important for filter performance; however, observation errors are generally not provided with datasets. The observation error is usually set to a fixed value (Ma et al., 2019), specific proportion of the observation value (Tang et al., 2013), or value calculated by combining measurement error with representative error as used in this study. Generally, the performance of data assimilation is sensitive to the specification

of the observation error (Tang et al., 2013). A sensitivity experiment (EMS6) with 901 doubled observation error was conducted to evaluate the influence of observation error 902 on the optimized emissions. Overall, the spatial distribution of emissions after 903 optimization was almost the same as that of the EMDA experiment but with a lower 904 increment (Figure S7), resulting in a weaker estimate of the national total emissions for 905 each species. This is because that the observation error inflates and the system becomes 906 more certain of the prior emission, and reduces the effect of observation information. 907 908 Figure 14 shows the time series of simulated and observed daily concentrations and their RMSEs verified against the assimilated sites. The simulations in VEP6 usually 909 performed worse, with larger biases and RMSEs than those of VEP (Figures S8 and S9), 910 especially in western and southern China, where posterior emissions were significantly 911 underestimated. These results generally corresponded to sluggish emission changes and 912 large Chi-square statistics (Figure S10), suggesting that an observation error that is too 913 large may substantially impact the estimated emissions. 914

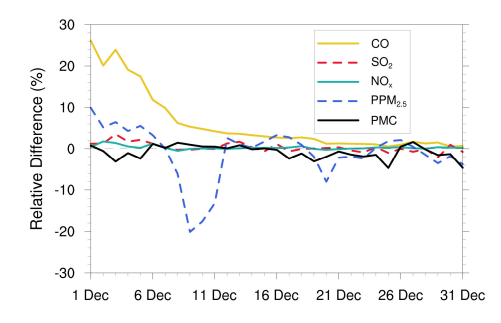


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916 Figure 14. Time series of the daily concentrations (CONC, left) and root mean square
917 error (RMSE, right) obtained from CEP, VEP, and VEP6. The simulations were verified
918 against the assimilated sites.

919 4.3.4 Impact of the IC optimization of the first window

920 Several studies indicate large emission discrepancies resulting from IC errors (Jiang et 921 al., 2013a; Miyazaki et al., 2017; Tang et al., 2013), which means that if the IC is not 922 optimized, the errors of concentrations would be compensated for through the 923 adjustment of emissions. To evaluate the impact of IC optimization of the first window

on the emission inversions, an EMS7 experiment without the IA step was conducted. 924 Figure 15 shows the time series of the relative differences in the daily posterior 925 emissions of the five species between the EMDA and EMS7 experiments. It can be 926 observed that IC optimization had a significant impact on the emission inversions of 927 long-lived species (i.e. CO). The overall difference in the inverted CO emissions 928 between the two experiments was approximately 5.3% but can reach 26.1% in the first 929 few windows. For the short-lived species, IC optimization had little impact on the 930 emissions; for example, the average emission differences of SO₂, NO_x, and PMC in the 931 two experiments were 0.3%, 0.3%, and 0.9%, respectively. For PPM_{2.5}, the average 932 emission difference is affected not only by primary emissions, but also by the complex 933 chemistry of its precursors. Therefore, the difference between the two experiments 934 fluctuated, with overall difference of 2%. Notably, with the gradual disappearance of 935 936 the benefit of IC assimilation, the two experiments reached a unified state after several windows. For CO, the impact of IA on emission inversion lasted approximately half a 937 month. These results indicate that removing the bias of the IC of the first DA window 938 939 is essential for the subsequent inverse analysis (Jiang et al., 2017).



940

Figure 15. Relative differences in CO, SO₂, NO_x, PPM_{2.5}, and PMC emissions (%, the ratio of absolute difference to EMDA) between the EMDA and EMS7.

943 4.4 Discussion

Optimal state estimation using an EnKF relies on the assumption of an unbiased 944 Gaussian prior error, which is not guaranteed in such highly nonlinear and large bias 945 systems. In this study, some pollutants (e.g. CO, PMC) have very large simulated biases; 946 947 thus, if a small uncertainty is adopted, the emission bias cannot be fully reduced. If a very large uncertainty is adopted, then the degree of freedom of adjustment is too large 948 949 and the inverted daily emissions will fluctuate abnormally. Therefore, we only set a larger prior uncertainty in the first three windows, adopting a moderate uncertainty in 950 the following windows and used a "two-step" inversion scheme and cyclic iteration to 951 gradually correct the emission errors. Figure 9a shows the time series of the relative 952 differences between prior and posterior emissions in each window. There were 953 relatively large adjustments for the emissions in the first three windows, especially for 954 PMC, but the adjustment ranges of the five species after the first three windows were 955 within the uncertainty range (e.g. \pm 25%), indicating that with this scheme, the EnKF 956 957 method used in this system had a good performance in emission inversion.

Model-data mismatch errors are from both the emissions and the inherent model errors 958 arising from the model structure, discretization, parameterizations, and biases in the 959 simulated meteorological fields. Neglecting model errors would attribute all 960 uncertainties to emissions and lead to considerable bias in the estimated emissions. In 961 the version of the CMAQ model used in this study, there are no heterogeneous reactions 962 (Quan et al., 2015; Wang et al., 2017), the parameterization scheme for the formation 963 of secondary organic aerosols (SOA) is imperfect (Carlton et al., 2008; Jiang et al., 964 965 2012; Yang et al., 2019), no feedback between chemistry and meteorology was considered, and we used an idea profile for chemical lateral boundary conditions. All 966 the above problems can lead to underestimated concentrations of pollutants, which in 967 turn require more emissions to compensate, leading to overestimation of emissions. In 968 addition, previous studies showed that ammonia emissions in the MEIC inventory are 969 underestimated (Kong et al., 2019b; Paulot et al., 2014; Zhang et al., 2018). Owing to 970 lack of ammonia observations, our system does not include emission estimates of 971

ammonia, which means that the concentration of ammonium aerosol was 972 underestimated in this system, also resulting in an overestimation of the PPM_{2.5} 973 974 emission. Wind-blown dust was also not simulated; thus, the PMC emission inverted in this system come from anthropogenic activities and natural sources. Although some of 975 these shortcomings can be solved by updating the CTM model, there will still be errors 976 in each parameterization and process. In general, a parameter estimation method was 977 used to reduce the model errors, in which some uncertain parameters were included in 978 the augmented state vector and optimized synchronously based on the available 979 observations (Brandhorst et al., 2017; Evensen, 2009). However, it is difficult to 980 identify the key uncertain parameters of different species in different models, which 981 generally comes not only from the complex atmospheric chemical model but also from 982 hundreds of model inputs (Tang et al., 2013). Another method is bias correction, which 983 984 treats the model error as a bias term and includes it in an augmented state vector (Brandhorst et al., 2017; De Lannoy et al., 2007; Keppenne et al., 2005). In addition, 985 the weak-constraint 4DVAR method can be used to reduce model errors, which adds a 986 987 correction term in the model integration to account for the different sources of model error (Sasaki, 1970). Although the reliable diagnosis of model error remains a challenge 988 (Laloyaux et al., 2020), it should be considered in an assimilation system. In the future, 989 we will consider model errors in our system to obtain better emission estimates. 990

Independent variable localization was adopted to avoid potential spurious correlations 991 across different species in this study. However, the transmission scales for different 992 993 species in different regions differ, and a more accurate localization range can be obtained through backward trajectory analysis. In addition, O₃ observations were not 994 995 assimilated to improve NO_x and VOC emissions using cross-species information. O_3 concentration and NO_x (VOC) emissions were positively correlated in the NO_x (VOC)-996 limited region and negatively correlated in the VOC (NO_x)-limited region (Tang et al., 997 2011; Wang et al., 2019b). Hamer et al. (2015) successfully used O₃ observations to 998 estimate NO_x and VOC emissions within the 4DVAR framework within an ideal model. 999 However, the NO_x emissions are often point or line sources, which are all small 1000

compared to the model resolution. With a coarse spatial resolution, the model cannot 1001 1002 accurately simulate the relationships between O₃ and its precursors. When assimilating O₃ observations to infer NO_x or VOC emissions, the inaccurate relationships simulated 1003 by model would worsen the inversion of NO_x emissions (Inness et al., 2015). In general, 1004 improving the model resolution can improve the detailed simulation and provide better 1005 prior information on O_3 -NO_x-VOC, but it is still difficult to determine whether the 1006 condition is NO_x-limited or VOC-limited in the real atmosphere using prior emissions 1007 1008 (Liu and Shi, 2021). Elbern et al. (2007) emphasized that assimilating O_3 to correct NO_x or VOC emissions must follow the EKMA framework derived based on observations, 1009 otherwise, even if the resolution is improved to sufficiently solve point and line sources, 1010 precursor emissions may be still adjusted in an opposite direction. This can be 1011 1012 demonstrated in our OSSE experiment at high resolution of 3 km (Figure S11). In this study, the spatial resolutions of the prior emission inventory (i.e., MEIC) is $0.25^{\circ} \times$ 1013 0.25°, which is appropriate for modeling at regional scales (Zheng et al., 2017). With 1014 this emission inventory, it is unable to accurately simulate the O_3 -NO_x-VOC 1015 1016 relationships. Therefore, to avoid the impact of inaccurate O_3 -NO_x relationship on emission inversion, in our system, we did not assimilate O₃, but directly assimilate NO₂ 1017 to optimize the NO_x emissions. This work will be followed by an ongoing study using 1018 the available VOC observations. 1019

1020 Although we do not assimilate O₃ observation, model resolution still has some influence on inversion results. In our previous study (Feng et al., 2022), we have inferred the NO_x 1021 emissions over YRD in China using NO₂ observations, which has a spatial resolution 1022 of 12 km. The study period, assimilated observations, and inversion settings are the 1023 1024 same as this study. We compared the posterior emissions of YRD between this study 1025 and Feng et al. (2022). The results showed that there was similar spatial distribution of 1026 posterior emissions inferred using the two resolutions (36 km vs 12 km) (Figure S12), but the total NO_x emission in YRD inferred using 36 km resolution was about 8.8% 1027 1028 higher than that inferred using 12 km resolution. The differences are mainly caused by meteorological differences at different resolutions. This indicates that coarse model 1029

resolution may lead to some overestimation of the inverted emissions. In addition, as shown previously, the concentrations after DA were evidently underestimated in western China, indicating that the inverted emissions over these regions still have large uncertainties because of the sparsity of observations, which are spatially insufficient for sampling the inhomogeneity of emissions. Therefore, further investigations with the joint assimilation of multisource observations (e.g. satellite) are underway.

 NO_x is mainly emitted by transportation (Li et al., 2017), which can reflect the level of 1036 economic activity to a certain extent. Weekly emission changes were explored to verify 1037 the performance of the system in depicting emission changes (Figure S13). Although 1038 the "weekend effect" of emissions in China is not significant (Wang et al., 2014; Wang 1039 et al., 2015), the posterior NO_x emission changes are in good agreement with the 1040 1041 observations. In our previous studies (Feng et al., 2020a; Feng et al., 2020b), this system was successfully applied to optimize NO_x and CO emissions. The inverted emission 1042 changes were also in line with the epidemic control time points. Additionally, the 1043 1044 emission changes can reflect the emission migration from developed or urban areas to 1045 developing or surrounding areas in recent years, which is consistent with the emission 1046 control strategies in China. Although the system did not consider the model error, 1047 resulting in a certain difference between the posterior and actual emissions, the spatiotemporal changes in posterior emissions were relatively reasonable and can be 1048 1049 used to monitor emission changes and inform emission regulations.

1050 **5 Summary and conclusions**

In this study, we developed a Regional multi-Air Pollutant Assimilation System (RAPASv1.0) based on the WRF/CMAQ model, 3DVAR algorithm, and EnKF algorithm. RAPAS can quantitatively optimize gridded emissions of CO, SO₂, NO_x, PPM_{2.5}, and PMC on a regional scale by simultaneously assimilating hourly *in-situ* measurements of CO, SO₂, NO₂, PM_{2.5}, and PM₁₀. This system includes two subsystems: IA subsystem and EI subsystem, which optimize chemical ICs and infer anthropogenic emissions. Taking the 2016 MEIC in December as a priori, the emissions of CO, SO₂, NO_x, PPM_{2.5}, and PMC in December 2016 were inferred by assimilating the corresponding nationwide observations over China. The optimized ICs and posterior emissions were examined against assimilated and independent observations through parallel forward simulation experiments with and without DA. Sensitivity tests were performed to investigate the impact of different inversion processes, prior emissions, prior uncertainties, and observation errors on emission estimates.

1065 RAPAS showed a good performance in assimilating surface *in-situ* observations, with the calculated emission uncertainties reduced by 44.4%, 45.0%, 34.3%, 51.8%, and 1066 56.1% for CO, SO₂, NO_x, PPM_{2.5}, and PMC, respectively. It can also significantly 1067 1068 improve the simulations; the RMSEs of the simulated concentrations with posterior 1069 emissions decreased by 40.1-56.3% and the CORRs increased from 0.26-0.66 to 0.69-0.87 for different species. The OSSE experiment showed that the errors of posterior CO, 1070 SO₂, NO_x, PPM_{2.5}, and PMC could be reduced by 78.4%, 86.1%, 78.8%, 77.6%, and 1071 1072 72.0%, respectively. Overall, compared with the prior emissions (MEIC 2016), the 1073 posterior emissions increased by 129%, 20%, 5%, and 95% for CO, SO₂, NO_x, and 1074 PPM_{2.5}, respectively. The posterior PMC emissions, which included anthropogenic and 1075 natural dust contributions, increased by 1045%. Sensitivity tests with different prior inventories showed that the observations in China were sufficient to infer emission and 1076 1077 that our system was less dependent on prior inventories. Additionally, sensitivity tests with different prior uncertainties indicated that when the posterior emissions were 1078 emissions, the emissions decreased/increased with larger than 1079 the prior decreases/increases in uncertainties because of the different convergence rates. These 1080 1081 results demonstrate the advantage of the two-step method in emission inversion in that 1082 the inversion errors of the last window can be transferred to the current window for 1083 further optimization and robustness of the emissions estimated from RAPAS using nationwide observations over China. It should be noted that the system usually responds 1084 1085 slowly to too small a priori uncertainties or too large observation errors, which may result in large errors in the estimated emissions. 1086

In summary, the comprehensive evaluation and sensitivity tests revealed that RAPAS could serve as a useful tool for accurately quantifying the spatial and temporal changes in multi-species emissions at regional scales and near-real time, which will be helpful for air pollution control in China and other regions around the world with dense ground observation networks.

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1093 **Code and data availability**

1094 The codes of RAPAS v1.0 are available at https://doi.org/10.5281/zenodo.5566225.

1095 The WRF model code is open-source code and can be obtained from the WRF Model

1096 User's Page (https://www2.mmm.ucar.edu/wrf/users, last access: 25 April 2021). The

1097 CMAQ model is available through an open license as well (https://www.epa.gov/cmaq,

last access: 25 April 2021). The observational and emission data used in this study are

available at https://doi.org/10.5281/zenodo.4718290 (Feng and Jiang, 2021).

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1101 Author contribution

SF, FJ, ZW and ZJ developed RAPAS v1.0. SF and FJ designed the research. SF performed model simulations, analyzed data, and prepared the paper with contributions from all co-authors. FJ supervised the model development project and assisted in conceptualization and writing. HW, WH, YS, LZ, YZ, CL, and WJ contributed to the discussion and improvement of the paper.

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1108 **Competing interests**

1109 The authors declare that they have no conflict of interest.

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1112 Acknowledgements

This work is supported by the National Key R&D Program of China (Grant No. 2020YFA0607504), the National Natural Science Foundation of China (Grant No. 41907378), and the Nanjing University Innovation and Creative Program for Ph.D. candidate (Grant No. CXCY19-60). We are grateful to the High Performance Computing Center (HPCC) of Nanjing University for doing the numerical calculations in this paper on its blade cluster system, and thank the MEIC team for providing the prior anthropogenic emissions (http://www.meicmodel.org/).

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1121 **References**

- Appel, K. W., Pouliot, G. A., Simon, H., Sarwar, G., Pye, H. O. T., Napelenok, S. L., Akhtar, F., and
 Roselle, S. J.: Evaluation of dust and trace metal estimates from the Community Multiscale Air
 Quality (CMAQ) model version 5.0, Geoscientific Model Development, 6, 883-899,
 10.5194/gmd-6-883-2013, 2013.
- Alexe, M., Bergamaschi, P., Segers, A., Detmers, R., Butz, A., Hasekamp, O., Guerlet, S., Parker,
 R., Boesch, H., Frankenberg, C., Scheepmaker, R. A., Dlugokencky, E., Sweeney, C., Wofsy,
 S. C., and Kort, E. A.: Inverse modelling of CH4 emissions for 2010-2011 using different
 satellite retrieval products from GOSAT and SCIAMACHY, Atmospheric Chemistry and
 Physics, 15, 113-133, 2015.
- Barbu, A. L., Segers, A. J., Schaap, M., Heemink, A. W., and Builtjes, P. J. H.: A multi-component
 data assimilation experiment directed to sulphur dioxide and sulphate over Europe,
 Atmospheric Environment, 43, 1622-1631, 2009.
- Bocquet, M.: Parameter-field estimation for atmospheric dispersion: application to the Chernobyl
 accident using 4D-Var, Quarterly Journal of the Royal Meteorological Society, 138, 664-681,
 2012.
- Bocquet, M., Elbern, H., Eskes, H., Hirtl, M., Žabkar, R., Carmichael, G. R., Flemming, J., Inness,
 A., Pagowski, M., Pérez Camaño, J. L., Saide, P. E., San Jose, R., Sofiev, M., Vira, J., Baklanov,
 A., Carnevale, C., Grell, G., and Seigneur, C.: Data assimilation in atmospheric chemistry
 models: current status and future prospects for coupled chemistry meteorology models,
 Atmospheric Chemistry and Physics, 15, 5325-5358, 2015.
- Bocquet, M. and Sakov, P.: Joint state and parameter estimation with an iterative ensemble Kalman
 smoother, Nonlinear Processes in Geophysics, 20, 803-818, 2013.
- Basu, S., Guerlet, S., Butz, A., Houweling, S., Hasekamp, O., Aben, I., Krummel, P., Steele, P.,
 Langenfelds, R., Torn, M., Biraud, S., Stephens, B., Andrews, A., and Worthy, D.: Global CO2
 fluxes estimated from GOSAT retrievals of total column CO2, Atmospheric Chemistry and

1147 Physics, 13, 8695-8717, 2013.

- Bauwens, M., Compernolle, S., Stavrakou, T., Müller, J.-F., van Gent, J., Eskes, H., Levelt, P. F.,
 van der A, R., Veefkind, J. P., Vlietinck, J., Yu, H., and Zehner, C.: Impact of Coronavirus
 Outbreak on NO2 Pollution Assessed Using TROPOMI and OMI Observations, 47,
 e2020GL087978, 10.1029/2020gl087978, 2020.
- 1152 Bierman: Factorization methods for Discrete Sequential estimation, Academic Press, 1977.
- Binkowski, F. S. and Roselle, S. J.: Models-3 community multiscale air quality (CMAQ) model
 aerosol component 1. Model description, Journal of Geophysical Research-Atmospheres, 108,
 10.1029/2001jd001409, 2003.
- Brandhorst, N., Erdal, D., and Neuweiler, I.: Soil moisture prediction with the ensemble Kalman
 filter: Handling uncertainty of soil hydraulic parameters, Advances in Water Resources, 110,
 360-370, 2017.
- Bruhwiler, L. M. P., Michalak, A. M., Peters, W., Baker, D. F., and Tans, P.: An improved Kalman
 Smoother for atmospheric inversions, Atmos. Chem. Phys., 5, 2691-2702, 10.5194/acp-52691-2005, 2005.
- Carlton, A. G., Turpin, B. J., Altieri, K. E., Seitzinger, S. P., Mathur, R., Roselle, S. J., and Weber,
 R. J.: CMAQ Model Performance Enhanced When In-Cloud Secondary Organic Aerosol is
 Included: Comparisons of Organic Carbon Predictions with Measurements, Environmental
 Science & Technology, 42, 8798-8802, 2008
- Chen, D., Liu, Z., Ban, J., and Chen, M.: The 2015 and 2016 wintertime air pollution in China: SO2
 emission changes derived from a WRF-Chem/EnKF coupled data assimilation system,
 Atmospheric Chemistry and Physics, 19, 8619-8650, 10.5194/acp-19-8619-2019, 2019.
- Chen, D., Liu, Z., Fast, J., and Ban, J.: Simulations of sulfate-nitrate-ammonium (SNA) aerosols
 during the extreme haze events over northern China in October 2014, Atmospheric Chemistry
 and Physics, 16, 10707-10724, 10.5194/acp-16-10707-2016, 2016.
- Chevallier, F., Bréon, F.-M., and Rayner, P. J.: Contribution of the Orbiting Carbon Observatory to
 the estimation of CO2 sources and sinks: Theoretical study in a variational data assimilation
 framework, 112, 10.1029/2006JD007375, 2007.
- Clements, A. L., Fraser, M. P., Upadhyay, N., Herckes, P., Sundblom, M., Lantz, J., and Solomon,
 P. A.: Chemical characterization of coarse particulate matter in the Desert Southwest Pinal
 County Arizona, USA, Atmospheric Pollution Research, 5, 52-61, 10.5094/apr.2014.007, 2014.
- Clements, N., Hannigan, M. P., Miller, S. L., Peel, J. L., and Milford, J. B.: Comparisons of urban and rural PM10-2.5 and PM2.5 mass concentrations and semi-volatile fractions in northeastern Colorado, Atmospheric Chemistry and Physics, 16, 7469-7484, 10.5194/acp-16-7469-2016, 2016.
- Daley, R.: Atmospheric Data Assimilation (gtSpecial IssueltData Assimilation in Meteology and
 Oceanography: Theory and Practice), Journal of the Meteorological Society of Japan. Ser. II,
 75, 319-329, 1997.

- 1185 Derber, J. C.: A VARIATIONAL CONTINUOUS ASSIMILATION TECHNIQUE, Monthly
 1186 Weather Review, 117, 2437-2446, 1989.
- de Foy, B., Lu, Z., Streets, D. G., Lamsal, L. N., and Duncan, B. N.: Estimates of power plant NOx
 emissions and lifetimes from OMI NO2 satellite retrievals, Atmospheric Environment, 116, 111, 10.1016/j.atmosenv.2015.05.056, 2015.
- De Lannoy, G. J. M., Houser, P. R., Pauwels, V. R. N., and Verhoest, N. E. C.: State and bias
 estimation for soil moisture profiles by an ensemble Kalman filter: Effect of assimilation depth
 and frequency, 43, 2007.
- Ding, J., van der A, R. J., Mijling, B., Levelt, P. F., and Hao, N.: NOx emission estimates during the
 2014 Youth Olympic Games in Nanjing, Atmospheric Chemistry and Physics, 15, 9399-9412,
 10.5194/acp-15-9399-2015, 2015.
- Elbern, H., Strunk, A., Schmidt, H., and Talagrand, O.: Emission rate and chemical state estimation
 by 4-dimensional variational inversion, Atmospheric Chemistry and Physics, 7, 3749-3769,
 10.5194/acp-7-3749-2007, 2007.
- Evensen, G.: The Ensemble Kalman Filter for Combined State and Parameter Estimation MONTE
 CARLO TECHNIQUES FOR DATA ASSIMILATION IN LARGE SYSTEMS, Ieee Control
 Systems Magazine, 29, 83-104, 10.1109/mcs.2009.932223, 2009.
- Feng, S., Jiang, F., Jiang, Z., Wang, H., Cai, Z., and Zhang, L.: Impact of 3DVAR assimilation of
 surface PM2.5 observations on PM2.5 forecasts over China during wintertime, Atmospheric
 Environment, 187, 34-49, 10.1016/j.atmosenv.2018.05.049, 2018.
- Feng, S., Jiang, F., Wang, H., Shen, Y., Zheng, Y., Zhang, L., Lou, C., and Ju, W.: Anthropogenic
 emissions estimated using surface observations and their impacts on PM2.5 source
 apportionment over the Yangtze River Delta, China, Science of The Total Environment, 828,
 154522, 2022
- Feng, S., Jiang, F., Wu, Z., Wang, H., Ju, W., and Wang, H.: CO Emissions Inferred From Surface
 CO Observations Over China in December 2013 and 2017, Journal of Geophysical ResearchAtmospheres, 125, 10.1029/2019jd031808, 2020a.
- Feng, S., Jiang, F., Wang, H., Wang, H., Ju, W., Shen, Y., Zheng, Y., Wu, Z., and Ding, A.: NOx
 Emission Changes Over China During the COVID-19 Epidemic Inferred From Surface NO2
 Observations, Geophysical Research Letters, 47, 10.1029/2020gl090080, 2020b.
- Feng, S. and Jiang, F.: Anthropogenic air pollutant emissions over China inferred by Regional multi Air Pollutant Assimilation System (RAPAS v1.0), Zenodo, 10.5281/zenodo.4718290, 2021.
- Gaspari, G. and Cohn, S. E.: Construction of correlation functions in two and three dimensions,
 Quarterly Journal of the Royal Meteorological Society, 125, 723-757, 10.1256/smsqj.55416,
 1999.
- Guenther, A. B., Jiang, X., Heald, C. L., Sakulyanontvittaya, T., Duhl, T., Emmons, L. K., and Wang,
 X.: The Model of Emissions of Gases and Aerosols from Nature version 2.1 (MEGAN2.1): an
 extended and updated framework for modeling biogenic emissions, Geoscientific Model

- 1223 Development, 5, 1471-1492, 10.5194/gmd-5-1471-2012, 2012.
- Gurney, K. R., Law, R. M., Denning, A. S., Rayner, P. J., Pak, B. C., Baker, D., Bousquet, P.,
 Bruhwiler, L., Chen, Y. H., Ciais, P., Fung, I. Y., Heimann, M., John, J., Maki, T., Maksyutov,
 S., Peylin, P., Prather, M., and Taguchi, S.: Transcom 3 inversion intercomparison: Model mean
 results for the estimation of seasonal carbon sources and sinks, Global Biogeochemical Cycles,
 18, 10.1029/2003gb002111, 2004.
- He, W., van der Velde, I. R., Andrews, A. E., Sweeney, C., Miller, J., Tans, P., van der Laan-Luijkx,
 I. T., Nehrkorn, T., Mountain, M., Ju, W., Peters, W., and Chen, H.: CTDAS-Lagrange v1.0: a
 high-resolution data assimilation system for regional carbon dioxide observations,
 Geoscientific Model Development, 11, 3515-3536, 10.5194/gmd-11-3515-2018, 2018.
- Hinds, W.C.: Aerosol Technology: Properties, Behavior, and Measurement of Airborne Particles.
 New York: John Wiley, 1982.
- 1235Houtekamer, P. L. and Mitchell, H. L.: A sequential ensemble Kalman filter for atmospheric data1236assimilation, Monthly Weather Review, 129, 123-137, 10.1175/1520-12370493(2001)129<0123:asekff>2.0.co;2, 2001.
- Houtekamer, P. L. and Zhang, F.: Review of the Ensemble Kalman Filter for Atmospheric Data
 Assimilation, Monthly Weather Review, 144, 4489-4532, 10.1175/mwr-d-15-0440.1, 2016.
- Inness, A., Blechschmidt, A. M., Bouarar, I., Chabrillat, S., Crepulja, M., Engelen, R. J., Eskes, H.,
 Flemming, J., Gaudel, A., Hendrick, F., Huijnen, V., Jones, L., Kapsomenakis, J., Katragkou,
 E., Keppens, A., Langerock, B., de Maziere, M., Melas, D., Parrington, M., Peuch, V. H.,
 Razinger, M., Richter, A., Schultz, M. G., Suttie, M., Thouret, V., Vrekoussis, M., Wagner, A.,
 and Zerefos, C.: Data assimilation of satellite-retrieved ozone, carbon monoxide and nitrogen
 dioxide with ECMWF's Composition-IFS, Atmospheric Chemistry and Physics, 15, 5275-5303,
 2015.
- Jiang, F., Liu, Q., Huang, X., Wang, T., Zhuang, B., and Xie, M.: Regional modeling of secondary
 organic aerosol over China using WRF/Chem, Journal of Aerosol Science, 43, 57-73,
 10.1016/j.jaerosci.2011.09.003, 2012a.
- Jiang, F., Zhou, P., Liu, Q., Wang, T., Zhuang, B., and Wang, X.: Modeling tropospheric ozone
 formation over East China in springtime, Journal of Atmospheric Chemistry, 69, 303-319,
 10.1007/s10874-012-9244-3, 2012b.
- Jiang, F., Wang, H. M., Chen, J. M., Machida, T., Zhou, L. X., Ju, W. M., Matsueda, H., and Sawa,
 Y.: Carbon balance of China constrained by CONTRAIL aircraft CO2 measurements,
 Atmospheric Chemistry and Physics, 14, 10133-10144, 10.5194/acp-14-10133-2014, 2014.
- Jiang, F., Wang, H., Chen, J. M., Ju, W., Tian, X., Feng, S., Li, G., Chen, Z., Zhang, S., Lu, X., Liu,
 J., Wang, H., Wang, J., He, W., and Wu, M.: Regional CO2 fluxes from 2010 to 2015 inferred
 from GOSAT XCO2 retrievals using a new version of the Global Carbon Assimilation System,
 Atmos. Chem. Phys., 21, 1963-1985, 10.5194/acp-21-1963-2021, 2021.
- Jiang, W., Smyth, S., Giroux, E., Roth, H., and Yin, D.: Differences between CMAQ fine mode
 particle and PM2.5 concentrations and their impact on model performance evaluation in the

1262 1263	lower Fraser valley, Atmospheric Environment, 40, 4973-4985, 10.1016/j.atmosenv.2005.10.069, 2006.
1264	Jiang, Z., Jones, D. B. A., Worden, H. M., Deeter, M. N., Henze, D. K., Worden, J., Bowman, K. W.,
1265	Brenninkmeijer, C. A. M., and Schuck, T. J.: Impact of model errors in convective transport on
1266	CO source estimates inferred from MOPITT CO retrievals, Journal Of Geophysical Research-
1267	Atmospheres, 118, 2073-2083, 2013a.
1268	Jiang, Z., Liu, Z., Wang, T., Schwartz, C. S., Lin, HC., and Jiang, F.: Probing into the impact of
1269	3DVAR assimilation of surface PM10 observations over China using process analysis, Journal
1270	of Geophysical Research: Atmospheres, 118, 6738-6749, 10.1002/jgrd.50495, 2013b.
1271	Jiang, Z., Worden, J. R., Worden, H., Deeter, M., Jones, D. B. A., Arellano, A. F., and Henze, D. K.:
1272	A 15-year record of CO emissions constrained by MOPITT CO observations, Atmospheric
1273	Chemistry And Physics, 17, 4565-4583, 10.5194/acp-17-4565-2017, 2017.
1274	Jin, J., Lin, H. X., Heemink, A., and Segers, A.: Spatially varying parameter estimation for dust
1275	emissions using reduced-tangent-linearization 4DVar, Atmospheric Environment, 187, 358-
1276	373, 10.1016/j.atmosenv.2018.05.060, 2018.
1277 1278	Kahnert, M.: Variational data analysis of aerosol species in a regional CTM: background error covariance constraint and aerosol optical observation operators, Tellus B, 60, 2008.
1279 1280	Kang, JS., Kalnay, E., Miyoshi, T., Liu, J., and Fung, I.: Estimation of surface carbon fluxes with an advanced data assimilation methodology, 117, 10.1029/2012JD018259, 2012.
1281	Keppenne, C. L., Rienecker, M. M., Kurkowski, N. P., and Adamec, D. A.: Ensemble Kalman filter
1282	assimilation of temperature and altimeter data with bias correction and application to seasonal
1283	prediction, Nonlin. Processes Geophys., 12, 491-503, 2005.
1284	Kleist, D. T., Parrish, D. F., Derber, J. C., Treadon, R., Wu, WS., and Lord, S.: Introduction of the
1285	GSI into the NCEP Global Data Assimilation System, Weather and Forecasting, 24, 1691-1705,
1286	10.1175/2009waf2222201.1, 2009.
1287	Kong, L., Tang, X., Zhu, J., Wang, Z., Pan, Y., Wu, H., Wu, L., Wu, Q., He, Y., Tian, S., Xie, Y., Liu,
1288	Z., Sui, W., Han, L., and Carmichael, G.: Improved Inversion of Monthly Ammonia Emissions
1289	in China Based on the Chinese Ammonia Monitoring Network and Ensemble Kalman Filter,
1290	Environmental Science & Technology, 53, 12529-12538, 10.1021/acsest.9b02701, 2019a.
1291 1292 1293 1294 1295 1296	 Kong, L., Tang, X., Zhu, J., Wang, Z., Fu, J. S., Wang, X., Itahashi, S., Yamaji, K., Nagashima, T., Lee, H. J., Kim, C. H., Lin, C. Y., Chen, L., Zhang, M., Tao, Z., Li, J., Kajino, M., Liao, H., Sudo, K., Wang, Y., Pan, Y., Tang, G., Li, M., Wu, Q., Ge, B., and Carmichael, G. R.: Evaluation and uncertainty investigation of the NO2, CO and NH3 modeling over China under the framework of MICS-Asia III, Atmos. Chem. Phys. Discuss., 2019, 1-33, 10.5194/acp-2018-1158, 2019b.
1297 1298	Laloyaux, P., Bonavita, M., Chrust, M., and Gürol, S.: Exploring the potential and limitations of weak-constraint 4D-Var, Quarterly Journal of the Royal Meteorological Society, 146, 4067-

4082, 2020

- Li, J.-d., Deng, Q.-h., Lu, C., and Huang, B.-l.: Chemical compositions and source apportionment
 of atmospheric PM10 in suburban area of Changsha, China, Journal of Central South
 University of Technology, 17, 509-515, 2010.
- Li, M., Zhang, Q., Kurokawa, J.-i., Woo, J.-H., He, K., Lu, Z., Ohara, T., Song, Y., Streets, D. G.,
 Carmichael, G. R., Cheng, Y., Hong, C., Huo, H., Jiang, X., Kang, S., Liu, F., Su, H., and Zheng,
 B.: MIX: a mosaic Asian anthropogenic emission inventory under the international
 collaboration framework of the MICS-Asia and HTAP, Atmospheric Chemistry and Physics,
 17, 935-963, 10.5194/acp-17-935-2017, 2017.
- Liu, C. and Shi, K.: A review on methodology in O3-NOx-VOC sensitivity study, Environmental
 Pollution, 291, 118249, 2021.
- Liu, Y., Kalnay, E., Zeng, N., Asrar, G., Chen, Z., and Jia, B.: Estimating surface carbon fluxes based
 on a local ensemble transform Kalman filter with a short assimilation window and a long
 observation window: an observing system simulation experiment test in GEOS-Chem 10.1,
 Geoscientific Model Development, 12, 2899-2914, 2019.
- Liu, Z., Liu, Q., Lin, H.-C., Schwartz, C. S., Lee, Y.-H., and Wang, T.: Three-dimensional variational assimilation of MODIS aerosol optical depth: Implementation and application to a dust storm
 over East Asia, Journal of Geophysical Research: Atmospheres, 116, n/a-n/a, 10.1029/2011jd016159, 2011.
- Lorenc, A. C.: Modelling of error covariances by 4D-Var data assimilation, Quarterly Journal of the
 Royal Meteorological Society, 129, 3167-3182, 2003.
- Hamer, P. D., Bowman, K. W., Henze, D. K., Attie, J. L., and Marecal, V.: The impact of observing
 characteristics on the ability to predict ozone under varying polluted photochemical regimes,
 Atmospheric Chemistry and Physics, 15, 10645-10667, 2015.
- Ma, C., Wang, T., Mizzi, A. P., Anderson, J. L., Zhuang, B., Xie, M., and Wu, R.: Multiconstituent
 Data Assimilation With WRF-Chem/DART: Potential for Adjusting Anthropogenic Emissions
 and Improving Air Quality Forecasts Over Eastern China, 124, 7393-7412,
 10.1029/2019jd030421, 2019.
- Meirink, J. F., Bergamaschi, P., and Krol, M. C.: Four-dimensional variational data assimilation for
 inverse modelling of atmospheric methane emissions: method and comparison with synthesis
 inversion, Atmospheric Chemistry and Physics, 8, 6341-6353, 2008.
- Meirink, J. F., Eskes, H. J., and Goede, A. P. H.: Sensitivity analysis of methane emissions derived
 from SCIAMACHY observations through inverse modelling, Atmospheric Chemistry and
 Physics, 6, 1275-1292, 10.5194/acp-6-1275-2006, 2006.
- 1333 Maybeck: Stochastic Models, Estimation and Control Academic Press, 1979.
- Miyazaki, K. and Eskes, H.: Constraints on surface NOx emissions by assimilating satellite
 observations of multiple species, Geophysical Research Letters, 40, 4745-4750,
 10.1002/grl.50894, 2013.
- 1337 Miyazaki, K., Eskes, H. J., and Sudo, K.: Global NOx emission estimates derived from an

- assimilation of OMI tropospheric NO2 columns, Atmospheric Chemistry and Physics, 12,
 2263-2288, 10.5194/acp-12-2263-2012, 2012a.
- Miyazaki, K., Eskes, H. J., Sudo, K., Takigawa, M., van Weele, M., and Boersma, K. F.:
 Simultaneous assimilation of satellite NO2, O-3, CO, and HNO3 data for the analysis of
 tropospheric chemical composition and emissions, Atmospheric Chemistry and Physics, 12,
 9545-9579, 10.5194/acp-12-9545-2012, 2012b.
- Miyazaki, K., Eskes, H., Sudo, K., Boersma, K. F., Bowman, K., and Kanaya, Y.: Decadal changes
 in global surface NOx emissions from multi-constituent satellite data assimilation,
 Atmospheric Chemistry and Physics, 17, 807-837, 2017.
- Mizzi, A. P., Edwards, D. P., and Anderson, J. L.: Assimilating compact phase space retrievals
 (CPSRs): comparison with independent observations (MOZAIC in situ and IASI retrievals)
 and extension to assimilation of truncated retrieval profiles, Geoscientific Model Development,
 1350 11, 3727-3745, 2018.
- Monteil, G., Houweling, S., Butz, A., Guerlet, S., Schepers, D., Hasekamp, O., Frankenberg, C.,
 Scheepmaker, R., Aben, I., and Rockmann, T.: Comparison of CH4 inversions based on 15
 months of GOSAT and SCIAMACHY observations, Journal of Geophysical ResearchAtmospheres, 118, 11807-11823, 2013.
- Muller, J. F. and Stavrakou, T.: Inversion of CO and NOx emissions using the adjoint of the
 IMAGES model, Atmospheric Chemistry and Physics, 5, 1157-1186, 2005.
- Nassar, R., Jones, D. B. A., Kulawik, S. S., Worden, J. R., Bowman, K. W., Andres, R. J.,
 Suntharalingam, P., Chen, J. M., Brenninkmeijer, C. A. M., Schuck, T. J., Conway, T. J., and
 Worthy, D. E.: Inverse modeling of CO2 sources and sinks using satellite observations of CO2
 from TES and surface flask measurements, Atmospheric Chemistry and Physics, 11, 60296047, 2011.
- Navon, I. M.: Practical and theoretical aspects of adjoint parameter estimation and identifiability in
 meteorology and oceanography, Dynamics of Atmospheres and Oceans, 27, 55-79, 1998.
- Parrish, D. F. and Derber, J. C.: The National Meteorological Center's spectral statisticalinterpolation analysis system, Monthly Weather Review, 120, 1747-1763, 10.1175/15200493(1992)120<1747:tnmcss>2.0.co;2, 1992.
- Paulot, F., Jacob, D. J., Pinder, R. W., Bash, J. O., Travis, K., and Henze, D. K.: Ammonia emissions
 in the United States, European Union, and China derived by high-resolution inversion of
 ammonium wet deposition data: Interpretation with a new agricultural emissions inventory
 (MASAGE_NH3), Journal of Geophysical Research-Atmospheres, 119, 4343-4364, 2014.
- Peng, Z., Liu, Z., Chen, D., and Ban, J.: Improving PM<sub>2.5</sub> forecast over
 China by the joint adjustment of initial conditions and source emissions with an ensemble
 Kalman filter, Atmospheric Chemistry and Physics, 17, 4837-4855, 10.5194/acp-17-48372017, 2017.
- Peng, Z., Lei, L., Liu, Z., Su, J., Ding, A., Ban, J., Chen, D., Kou, X., and Chu, K.: The impact of
 multi-species surface chemical observation assimilation on air quality forecasts in China,

- 1377 Atmospheric Chemistry and Physics, 18, 10.5194/acp-18-17387-2018, 2018.
- Peters, W., Jacobson, A. R., Sweeney, C., Andrews, A. E., Conway, T. J., Masarie, K., Miller, J. B.,
 Bruhwiler, L. M. P., Petron, G., Hirsch, A. I., Worthy, D. E. J., van der Werf, G. R., Randerson,
 J. T., Wennberg, P. O., Krol, M. C., and Tans, P. P.: An atmospheric perspective on North
 American carbon dioxide exchange: CarbonTracker, Proceedings of the National Academy of
 Sciences of the United States of America, 104, 18925-18930, 10.1073/pnas.0708986104, 2007.
- Peylin, P., Rayner, P. J., Bousquet, P., Carouge, C., Hourdin, F., Heinrich, P., Ciais, P., and
 contributors, A.: Daily CO2 flux estimates over Europe from continuous atmospheric
 measurements: 1, inverse methodology, Atmospheric Chemistry and Physics, 5, 3173-3186,
 10.5194/acp-5-3173-2005, 2005.
- Purser, R. J., Wu, W. S., Parrish, D. F., and Roberts, N. M.: Numerical aspects of the application of recursive filters to variational statistical analysis. Part I: Spatially homogeneous and isotropic
 Gaussian covariances, Monthly Weather Review, 131, 1524-1535, 10.1175//1520-0493(2003)131<1524:naotao>2.0.co;2, 2003.
- Quan, J., Liu, Q., Li, X., Gao, Y., Jia, X., Sheng, J., Liu, Y., 2015. Effect of heterogeneous aqueous
 reactions on the secondary formation of inorganic aerosols during haze events. Atmospheric
 Environment 122, 306-312.
- Rabier, F., McNally, A., Andersson, E., Courtier, P., Unden, P., Eyre, J., Hollingsworth, A., and
 Bouttier, F.: The ECMWF implementation of three-dimensional variational assimilation (3DVar). II: Structure functions, Quarterly Journal Of the Royal Meteorological Society, 124,
 1809-1829, 10.1256/smsqj.55002, 1998.
- Reichle, R. H., McLaughlin, D. B., and Entekhabi, D.: Hydrologic data assimilation with the
 ensemble Kalman filter, Monthly Weather Review, 130, 103-114, 2002.
- 1400 Richardson, H., Basu, S., and Holtslag, A. A. M.: Improving Stable Boundary-Layer Height
 1401 Estimation Using a Stability-Dependent Critical Bulk Richardson Number, Boundary-Layer
 1402 Meteorology, 148, 93-109, 2013.
- Ruiz, J. and Pulido, M.: Parameter Estimation Using Ensemble-Based Data Assimilation in the
 Presence of Model Error, Monthly Weather Review, 143, 1568-1582, 2015.
- Sarwar, G., Simon, H., Bhave, P., and Yarwood, G.: Examining the impact of heterogeneous nitryl
 chloride production on air quality across the United States, Atmospheric Chemistry and
 Physics, 12, 6455-6473, 10.5194/acp-12-6455-2012, 2012.
- Sasaki, Y.: SOME BASIC FORMALISMS IN NUMERICAL VARIATIONAL ANALYSIS,
 Monthly Weather Review, 98, 875-&, 1970.
- Schneising, O., Buchwitz, M., Burrows, J. P., Bovensmann, H., Bergamaschi, P., and Peters, W.:
 Three years of greenhouse gas column-averaged dry air mole fractions retrieved from satellite
 Part 2: Methane, Atmospheric Chemistry and Physics, 9, 443-465, 2009.
- Schwartz, C. S., Liu, Z., Lin, H.-C., and Cetola, J. D.: Assimilating aerosol observations with a
 "hybrid" variational-ensemble data assimilation system, Journal Of Geophysical Research-

- 1415 Atmospheres, 119, 4043-4069, 10.1002/2013jd020937, 2014.
- Sekiyama, T. T., Tanaka, T. Y., Shimizu, A., and Miyoshi, T.: Data assimilation of CALIPSO aerosol
 observations, Atmospheric Chemistry and Physics, 10, 39-49, 10.5194/acp-10-39-2010, 2010.
- Shen, Y., Jiang, F., Feng, S., Zheng, Y., Cai, Z., and Lyu, X.: Impact of weather and emission changes
 on NO2 concentrations in China during 2014–2019, Environmental Pollution, 269, 116163,
 10.1016/j.envpol.2020.116163, 2021.
- Shi, X. and Brasseur, G. P.: The Response in Air Quality to the Reduction of Chinese Economic
 Activities During the COVID-19 Outbreak, 47, e2020GL088070, 10.1029/2020gl088070,
 2020.
- Stanevich, I., Jones, D. B. A., Strong, K., Keller, M., Henze, D. K., Parker, R. J., Boesch, H., Wunch,
 D., Notholt, J., Petri, C., Warneke, T., Sussmann, R., Schneider, M., Hase, F., Kivi, R.,
 Deutscher, N. M., Velazco, V. A., Walker, K. A., and Deng, F.: Characterizing model errors in
 chemical transport modeling of methane: using GOSAT XCH4 data with weak-constraint fourdimensional variational data assimilation, Atmospheric Chemistry and Physics, 21, 9545-9572,
 2021.
- Stavrakou, T., Müller, J.-F., Boersma, K. F., De Smedt, I., and van der A, R. J.: Assessing the
 distribution and growth rates of NOx emission sources by inverting a 10-year record of NO2
 satellite columns, 35, 10.1029/2008gl033521, 2008.
- Sun, A. Y., Morris, A., and Mohanty, S.: Comparison of deterministic ensemble Kalman filters for
 assimilating hydrogeological data, Advances in Water Resources, 32, 280-292,
 10.1016/j.advwatres.2008.11.006, 2009.
- Takagi, H., Saeki, T., Oda, T., Saito, M., Valsala, V., Belikov, D., Saito, R., Yoshida, Y., Morino, I.,
 Uchino, O., Andres, R. J., Yokota, T., and Maksyutov, S.: On the Benefit of GOSAT
 Observations to the Estimation of Regional CO₂ Fluxes, SOLA, 7, 161-164,
 10.2151/sola.2011-041, 2011.
- Tang, X., Zhu, J., Wang, Z. F., and Gbaguidi, A.: Improvement of ozone forecast over Beijing based
 on ensemble Kalman filter with simultaneous adjustment of initial conditions and emissions,
 Atmospheric Chemistry And Physics, 11, 12901-12916, 10.5194/acp-11-12901-2011, 2011.
- Tang, X., Zhu, J., Wang, Z. F., Wang, M., Gbaguidi, A., Li, J., Shao, M., Tang, G. Q., and Ji, D. S.:
 Inversion of CO emissions over Beijing and its surrounding areas with ensemble Kalman filter,
 Atmospheric Environment, 81, 676-686, 10.1016/j.atmosenv.2013.08.051, 2013.
- Wang, C., Lei, L., Tan, Z.-M., and Chu, K.: Adaptive Localization for Tropical Cyclones With
 Satellite Radiances in an Ensemble Kalman Filter, Frontiers in Earth Science, 8,
 10.3389/feart.2020.00039, 2020.
- Wang, H., Jiang, F., Wang, J., Ju, W., and Chen, J. M.: Terrestrial ecosystem carbon flux estimated
 using GOSAT and OCO-2 XCO2 retrievals, Atmospheric Chemistry and Physics, 19, 1206712082, 2019a.
- 1452 Wang, N., Lyu, X., Deng, X., Huang, X., Jiang, F., and Ding, A.: Aggravating O3 pollution due to

- 1453 NOx emission control in eastern China, Science of The Total Environment, 677, 732-744,1454 2019b.
- Wang, Y. H., Hu, B., Ji, D. S., Liu, Z. R., Tang, G. Q., Xin, J. Y., Zhang, H. X., Song, T., Wang, L.
 L., Gao, W. K., Wang, X. K., and Wang, Y. S.: Ozone weekend effects in the Beijing-TianjinHebei metropolitan area, China, Atmospheric Chemistry and Physics, 14, 2419-2429, 2014.
- Wang, Z., Li, Y., Dong, X., Sun, R., Sun, N., and Pan, L.: Analysis on weekend effect of air
 pollutants in urban atmosphere of Beijing, Journal of University of Chinese Academy of
 Sciences, 32, 843-850, 2015.
- Wang, Z., Wang, W., Tham, Y.J., Li, Q., Wang, H., Wen, L., Wang, X., Wang, T., 2017. Fast
 heterogeneous N2O5 uptake and CINO2 production in power plant and industrial plumes
 observed in the nocturnal residual layer over the North China Plain. Atmospheric Chemistry
 and Physics 17, 12361-12378.
- Wecht, K. J., Jacob, D. J., Sulprizio, M. P., Santoni, G. W., Wofsy, S. C., Parker, R., Boesch, H., and
 Worden, J.: Spatially resolving methane emissions in California: constraints from the CalNex
 aircraft campaign and from present (GOSAT, TES) and future (TROPOMI, geostationary)
 satellite observations, Atmospheric Chemistry and Physics, 14, 8173-8184, 2014.
- Wu, H., Tang, X., Wang, Z., Wu, L., Li, J., Wang, W., Yang, W., and Zhu, J.: High-spatiotemporalresolution inverse estimation of CO and NOx emission reductions during emission control
 periods with a modified ensemble Kalman filter, Atmospheric Environment, 236,
 10.1016/j.atmosenv.2020.117631, 2020.
- Wu, W. S., Purser, R. J., and Parrish, D. F.: Three-dimensional variational analysis with spatially
 inhomogeneous covariances, Monthly Weather Review, 130, 2905-2916, 10.1175/15200493(2002)130<2905:tdvaws>2.0.co;2, 2002.
- Yang, W., Li, J., Wang, W., Li, J., Ge, M., Sun, Y., Chen, X., Ge, B., Tong, S., Wang, Q., and Wang,
 Z.: Investigating secondary organic aerosol formation pathways in China during 2014,
 Atmospheric Environment, 213, 133-147, 2019.
- Yumimoto, K., Uno, I., Sugimoto, N., Shimizu, A., Liu, Z., and Winker, D. M.: Adjoint inversion
 modeling of Asian dust emission using lidar observations, Atmospheric Chemistry and Physics,
 8, 2869-2884, 2008.
- Zhang, F., Weng, Y., Sippel, J. A., Meng, Z., and Bishop, C. H.: Cloud-Resolving Hurricane
 Initialization and Prediction through Assimilation of Doppler Radar Observations with an
 Ensemble Kalman Filter, Monthly Weather Review, 137, 2105-2125, 10.1175/2009mwr2645.1,
 2009a.
- Zhang, L., Chen, Y., Zhao, Y., Henze, D. K., Zhu, L., Song, Y., Paulot, F., Liu, X., Pan, Y., Lin, Y.,
 and Huang, B.: Agricultural ammonia emissions in China: reconciling bottom-up and top-down
 estimates, Atmospheric Chemistry and Physics, 18, 339-355, 2018.
- Zhang, Q., Streets, D. G., Carmichael, G. R., He, K. B., Huo, H., Kannari, A., Klimont, Z., Park, I.
 S., Reddy, S., Fu, J. S., Chen, D., Duan, L., Lei, Y., Wang, L. T., and Yao, Z. L.: Asian emissions
 in 2006 for the NASA INTEX-B mission, Atmospheric Chemistry and Physics, 9, 5131-5153,

- 1492 10.5194/acp-9-5131-2009, 2009b.
- Zhang, S., Zheng, X., Chen, J. M., Chen, Z., Dan, B., Yi, X., Wang, L., and Wu, G.: A global carbon
 assimilation system using a modified ensemble Kalman filter, Geosci. Model Dev., 8, 805-816,
 10.5194/gmd-8-805-2015, 2015.
- Zhang, X., Liu, J., Han, H., Zhang, Y., Jiang, Z., Wang, H., Meng, L., Li, Y. C., and Liu, Y.: SatelliteObserved Variations and Trends in Carbon Monoxide over Asia and Their Sensitivities to
 Biomass Burning, Remote Sensing, 12, 10.3390/rs12050830, 2020.
- Zheng, B., Tong, D., Li, M., Liu, F., Hong, C., Geng, G., Li, H., Li, X., Peng, L., Qi, J., Yan, L.,
 Zhang, Y., Zhao, H., Zheng, Y., He, K., and Zhang, Q.: Trends in China's anthropogenic
 emissions since 2010 as the consequence of clean air actions, Atmospheric Chemistry And
 Physics, 18, 14095-14111, 10.5194/acp-18-14095-2018, 2018.
- Zheng, B., Zhang, Q., Tong, D., Chen, C., Hong, C., Li, M., Geng, G., Lei, Y., Huo, H., and He, K.:
 Resolution dependence of uncertainties in gridded emission inventories: a case study in Hebei,
 China, Atmospheric Chemistry and Physics, 17, 921-933, 2017.