



# NAQPMS-PDAF v2.0: A Novel Hybrid Nonlinear Data Assimilation System for Improved Simulation of PM<sub>2.5</sub> Chemical Components

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13 Abstract. PM2.5, a complex mixture with diverse chemical components, exerts significant impacts on the environment, human 14 health, and climate change. However, precisely describing spatiotemporal variations of PM2.5 chemical components remains a 15 difficulty. In our earlier work, we developed an aerosol extinction coefficient data assimilation (DA) system (NAQPMS-PDAF 16 v1.0) that is suboptimal for chemical components. This paper introduces a novel hybrid nonlinear chemical DA system 17 (NAQPMS-PDAF v2.0) to accurately interpret key chemical components (SO42-, NO3-, NH4+, OC, and EC). NAQPMS-PDAF 18 v2.0 improves upon v1.0 by effectively handing and balancing stability and nonlinearity in chemical DA, which is achieved 19 by incorporating the non-Gaussian-distribution ensemble perturbation and hybrid Localized Kalman-Nonlinear Ensemble 20 Transform Filter with an adaptive forgetting factor for the first time. The dependence tests demonstrate that NAQPMS-PDAF 21 v2.0 provides excellent DA results with a minimal ensemble size of 10, surpassing previous reports and v1.0. A one-month 22 DA experiment shows that the analysis field generated by NAQPMS-PDAF v2.0 is in good agreement with observations, 23 especially reducing the underestimation of NH4<sup>+</sup> and NO3<sup>-</sup> and the overestimation of SO4<sup>2-</sup>, OC, and EC. In particular, the 24 CORR values for NO3, OC, and EC are above 0.96, and R<sup>2</sup> values are above 0.93. NAQPMS-PDAF v2.0 also demonstrates 25 superior spatiotemporal interpretation, with most DA sites showing improvements of over 50%-200% in CORR and over 50%-26 90% in RMSE for the five chemical components. Compared to the poor performance in global reanalysis dataset (CORR: 27 0.42-0.55, RMSE: 4.51-12.27 µg/m3) and NAQPMS-PDAF v1.0 (CORR: 0.35-0.98, RMSE: 2.46-15.50 µg/m3), NAQPMS-28 PDAF v2.0 has the highest CORR of 0.86-0.99 and the lowest RMSE of 0.14-3.18 µg/m<sup>3</sup>. The uncertainties in ensemble DA 29 are also examined, further highlighting the potential of NAQPMS-PDAF v2.0 for advancing aerosol chemical component 30 studies.





#### 31 1 Introduction

32 PM2.5 is a complex mixture of various chemical fractions, mainly including sulfate (SO42-), nitrate (NO3-), ammonium (NH4+), 33 organic carbon (OC), and elemental carbon (EC), which diversely influences the atmospheric environment (Khanna et al., 34 2018), human health (Bell et al., 2007; Schlesinger, 2007; Li et al., 2022a; Alves et al., 2023), and climate change (Schult et 35 al., 1997; Park et al., 2014; Wilcox et al., 2016). However, current detection technologies, such as field observation with in-36 situ sampling and chemical analysis (Zhang et al., 2015; Ming et al., 2017), remote-sensing inversion (Nishizawa et al., 2008; 37 Nishizawa et al., 2011; Nishizawa et al., 2017), and machine learning (Lin et al., 2022; Su Lee et al., 2023) are insufficient in 38 interpreting PM<sub>2.5</sub> chemical components due to the spatiotemporal discontinuity and limited chemical species. Although 39 atmospheric chemistry transport models (CTMs) (Wang et al., 2014; Wang et al., 2015; Jia et al., 2017; Yang et al., 2019; Li 40 et al., 2020; Lv et al., 2020) are commonly used to characterize spatiotemporal distribution of multiple chemical species, CTMs 41 are associated with uncertainties in initial-boundary conditions, physiochemical mechanisms, emission inventories, and 42 meteorological fields (Sax and Isakov, 2003; Mallet and Sportisse, 2006; Rodriguez et al., 2007; Chang et al., 2015; Miao et 43 al., 2020; Xie et al., 2022), resulting in biases relative to real situation.

44

45 Data assimilation (DA) offers a solution to integrate the multi-source observations, CTMs, and their uncertainties effectively 46 to enhance the simulation and forecasting capabilities of CTMs. Variational methods (3D-Var/4D-Var) (Talagrand and Courtier, 47 1987), Ensemble Kalman Filter (EnKF) (Evensen, 1994; Evensen, 2003), EnKF-variants (EnKFs) (Bishop et al., 2001; Tippett 48 et al., 2003; Hunt et al., 2007; Nerger et al., 2012), and hybrid EnKF-Var methods (Hamill and Snyder, 2000; Schwartz et al., 49 2014) are most widely applied in DA. However, variational methods have a flow-independent Background Error Covariance 50 (BEC) with the assumption of isotropic, static, and uniform characteristics, and they need to develop the tangent linear adjoint 51 model, which is difficult to practice for complex models. Although EnKFs and hybrid EnKF-Var methods have a flow-52 dependent BEC, they are sensitive to inadequate ensemble sampling and have high computational costs. Importantly, these 53 methods cannot address model nonlinearity and non-Gaussian error distribution, yielding suboptimal results for DA in highly 54 nonlinear CTMs.

55

Currently, nonlinear filters, such as Particle Filter (PF) (Gordon et al., 1993) and Nonlinear Ensemble Transform Filter (NETF) (Tödter and Ahrens, 2015), have been proposed to approximate the complete posterior probability distribution of model states and provide a better representation of non-Gaussian information based on Monte Carlo random sampling and Bayesian theory. However, PF is unstable and susceptible to filter degeneration compared to EnKFs. In a recent study, Nerger (2022) proposed the hybrid Kalman-Nonlinear Ensemble Transform Filter (KNETF) to achieve excellent DA performance in the Lorenz-63 and Lorenz-96 model with a smaller ensemble size, which combines the stability of EnKFs and the nonlinearity of NETF (Nerger,





62 2022). However, to the author's knowledge, this algorithm has not been applied to the chemical DA of CTMs.

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64 Studies on chemical DA involve the assimilation of aerosol optical properties, such as aerosol optical depth (AOD) and 65 extinction coefficient (EXT), and the particulate matters (PMs), such as the mass concentrations of PM<sub>2.5</sub> and PM<sub>10</sub>. The 66 commonly AOD observations for DA include OMI-AOD (Ali et al., 2013), MODIS-AOD (Zhang et al., 2008; Huneeus et al., 67 2012; Huneeus et al., 2013; Rubin and Collins, 2014; Lynch et al., 2016; Werner et al., 2019; Kumar et al., 2020), AERONET-68 AOD (Schutgens et al., 2010; Li et al., 2016), Sun-Sky Photometer-Multiband AOD (Chang et al., 2021), GOCI-AOD (Saide 69 et al., 2014; Luo et al., 2020; Kim et al., 2021), and Fengyun/Himawari8-AOD (Bao et al., 2019; Jin et al., 2019; Xia et al., 70 2019; Xia et al., 2020). These studies indicated that AOD observations can enhance the accuracy of aerosol simulation and 71 forecast. Compared to AOD, EXT DA effectively improves the interpretation of aerosol vertical distribution (Zhang et al., 72 2014; Cheng et al., 2019; Wang et al., 2022). Additionally, the simultaneous DA of aerosol optical properties and PMs is widely 73 applied in aerosol studies (Tang et al., 2015; Chai et al., 2017). According to our literature review (Yang et al., 2023), there is 74 currently no DA study on aerosol chemical components due to the limited DA influence of PMs and AOD on chemical 75 compositions (Chang et al., 2021) and the limited chemical observations with an extensive spatial range. Moreover, the aerosol 76 chemical components exhibit nonlinearity and a non-Gaussian distribution (Ha, 2022), while current main-stream algorithms, 77 such as variational methods or EnKFs, are suboptimal for chemical component DA.

78

79 In our previous work, we developed an aerosol vertical DA system (NAQPMS-PDAF v1.0) based on EnKFs to improve the 80 simulation of the extinction coefficient vertical profile (Wang et al., 2022). In this study, we present a novel hybrid nonlinear 81 DA system (NAQPMS-PDAF v2.0) towards various PM2.5 chemical components through online integration of Parallel Data 82 Assimilation Framework (PDAF, version 2.1, released on February 21st, 2023), Observation Module Infrastructure (OMI) and 83 Nested Air Quality Prediction Model System (NAQPMS). We collected 1-month hourly surface observations of five PM2.5 84 chemical components (NH4+, SO42-, NO3-, OC, and EC) over Northern China and surrounding areas. We utilized the hybrid 85 Localized Kalman-Nonlinear Ensemble Transform Filter (LKNETF) to generate a high-resolution and high-accuracy 86 reanalysis dataset of PM2.5 chemical components for the first time. Notably, the ensemble members in NAQPMS-PDAF v2.0 87 are generated by perturbing emission species based on their uncertainties and non-Gaussian distribution assumption. Section 88 2 briefly introduces NAQPMS and PDAF v2.1 with OMI, respectively, and details the development of NAQPMS-PDAF v2.0, 89 including system structure, configuration, ensemble generation, and LKNETF algorithm. The data used in this study and 90 experimental settings are also described in Section 2. Section 3 presents the DA results, including evaluating dependencies, 91 performance, and external comparisons. Besides, Section 3 discusses the ensemble DA uncertainty. Section 4 summarizes the 92 conclusions and outlook.





#### 93 2 Method and data

#### 94 **2.1 NAQPMS**

The Nested Air Quality Prediction Modeling System (NAQPMS), developed by the Institute of Atmospheric Physics (IAP), Chinese Academy of Sciences (CAS), is used to provide background fields of key aerosol chemical components in this study. NAQPMS is capable of characterizing the three-dimensional spatiotemporal distribution of various atmospheric compositions at global and regional scales through multiple physicochemical processes (shown in Table S1) and has been widely used in atmospheric pollution and chemistry research, such as O<sub>3</sub> pollution, haze episodes (Wang et al., 2014; Du et al., 2021), regional transport (Wang et al., 2017; Wang et al., 2019), source identification (Li et al., 2022b), air quality simulation at global scale (Ye et al., 2021) and at urban-street scale (Wang et al., 2023), and acid deposition (Ge et al., 2014).

# 102 **2.2 PDAF v2.1 with OMI**

103 The Parallel Data Assimilation Framework (PDAF, https://pdaf.awi.de/trac/wiki) is an open-source and high-expandability 104 software developed by the Alfred Wegener Institute (AWI) in Germany to integrate observations, numerical models, and 105 assimilation systems for DA tasks, widely applied in meteorology, oceanography, land surface and atmospheric chemistry 106 (Kurtz et al., 2016; Nerger et al., 2020; Mingari et al., 2022; Strebel et al., 2022; Wang et al., 2022; Yu et al., 2022). The initial 107 version of PDAF (PDAF v1.0) was released in 2004. It has undergone continuous improvements and updates, with major 108 updates including the introduction of Ensemble Transform Kalman Filter (ETKF) and its localized variant (LETKF) in version 109 1.6, the implementation of PDAF-OMI (Observation Module Infrastructure) in version 1.16, the integration of 3D-Var methods 110 in version 2.0, and the incorporation of the hybrid KNETF and its localized variant (LKNETF) for the first time in version 2.1, 111 which was released in 2023 to handle the complex DA situations, such as the nonlinearity of system and non-Gaussian error 112 distribution of model state. Notably, the version of PDAF coupled in NAQPMS-PDAF v1.0 is PDAF v1.15 (released in 2019), 113 implying that NAQPMS-PDAF v1.0 has more limited applicability and functionality. In this work, the PDAF v2.1 is coupled 114 in NAQPMS-PDAF v2.0.

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PDAF has two modes, namely offline and online mode. For the offline mode, PDAF and the model perform separately without coupling, which is easy to write code. For the online mode, PDAF is coupled with the model, and model calculation and data assimilation perform continuously. Compared to the offline mode, the online coupling has several advantages. Firstly, the initialization process of PDAF and the model only needs to be executed once instead of twice independently. Secondly, the model integration result can be directly passed to PDAF for data assimilation. Additionally, the assimilation result of PDAF can be directly passed to the model for the next model integration. This eliminates the need for intermediate steps and improves efficiency. Thirdly, the online mode is controlled by a main program, which allows for efficient use of several processors in





- 123 the high-performance computing cluster. Conversely, in the offline mode, the PDAF and the model are managed by distinct 124 programs, often with a reduced number of processors available for each program. Therefore, the online-mode PDAF is used
- 125 in this study.
- 126
- 127 PDAF-OMI, an extension of PDAF, provides I/O interfaces for multi-type observations, simplifying user observation handling 128 by offering generic PDAF-OMI core routines and independent user-supplied routines for each observational type. The user-129 supplied routines, namely init dim obs/init dim obs l, obs op, and localize covar, are responsible for reading and writing 130 multi-type observations, applying corresponding observation operators, and performing covariance localization, respectively. 131 The modules for all observation types are integrated into the callback obs pdafomi, allowing free combinations between 132 different observation types without interference and facilitating the collaborative DA for various aerosol chemical components. 133 PDAF-OMI was not applied in NAQPMS-PDAF v1.0. Consequently, NAQPMS-PDAF v1.0 cannot switch between different 134 observational type combinations, and users need to define complete routines for each observation type for the DA process, 135 resulting in more tedious code writing and higher computational costs in NAQPMS-PDAF v1.0.
- 136 2.3 NAQPMS-PDAF v2.0

# 137 2.3.1 Structure of NAQPMS-PDAF v2.0

Figure 1 illustrates the structure and main workflow of NAQPMS-PDAF v2.0. The observational part involves multi-type observations and PDAF-OMI. PDAF-OMI enables the simultaneous access and scheduling of multi-type and multi-source data through observational indices, which allows for flexible combination. The ensemble forecast/background fields are generated by perturbing emission species (see Sect. 2.3.3) and NAQPMS calculations (the green part in Fig. 1). Then chemical DA is performed by a novel hybrid localized nonlinear DA algorithm (LKNETF, see Sect. 2.3.4) with an adaptive hybrid weight and an adaptive forgetting factor to generate analysis/initial fields for the next realization.







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146 NAQPMS-PDAF v2.0 implements an online coupling between NAQPMS and PDAF v2.1 with OMI, utilizing a level-2 147 parallel computational framework. The online coupling ensures the continuous operation of model forecasts and assimilation 148 analysis at each time step, achieved by directly integrating PDAF routines into the prototype code of NAQPMS (In Fig. 1 right 149 part, the blue represents NAQPMS main routines, while the yellow represents PDAF main routines). The level-2 parallel 150 computational framework, which utilizes the Message Passing Interface standard (MPI), facilitates concurrent processing and 151 data exchange among multiple ensemble members and parallel computation among model state matrixes within each ensemble 152 member, enhancing the efficiency of ensemble analysis and numerical model computations. The description of level-2 parallel 153 implementation was detailed in our previous work (Wang et al. 2022). The workflow of NAQPMS-PDAF v2.0 is outlined as 154 follows:

155 Step 1. init\_system module initializes NAQPMS, such as defining all model state variables, allocating numerical matrixes,

- 156 configuring parameters, I/O of meteorological fields, and emission input.
- 157 Step 2. init\_parallel module initializes MPI (MPI\_COMM\_WORLD) and model communicator (MPI\_COMM\_MODEL),
- 158 their number of processes, and the rank of a process, followed by *init\_parallel\_pdaf*, which initializes MPI communicators for
- the model tasks, filter tasks and the coupling between model and filter tasks.
- 160 Step 3. initialize module initializes the target field (such as PM<sub>2.5</sub> chemical components), such as their spatiotemporal
- 161 dimensions (longitude, latitude, and time steps) and variable dimensions.
- 162 Step 4. init\_pdaf module initializes PDAF variables, such as the local state dimension, global state dimension, and settings for
- 163 analysis steps.





164 Step 5. Perform the time loop of forecast and analysis. The convert field module is employed to match the matrix storage rule 165 of the target field between NAQPMS and PDAF to ensure compatibility. The *field2var* module collects the analysis field/initial 166 field and establishes a relationship between the initial field/analysis field and sub-variables in NAQPMS. Subsequently, the 167 analysis field values are allocated to the corresponding NAQPMS sub-variables, and then the NAQPMS processes module 168 performs the forecast. After that, the var2field module, the inverse of the field2var module, assigns the NAQPMS sub-variables 169 to the forecast field/background field. Finally, the assimilate pdaf module assimilates the target field with observations to 170 generate an analysis field for the next iteration. 171 Step 6. post-processing is responsible for finalizing NAPQMS-PDAF, data analysis, and DA evaluation.

#### 172 **2.3.2** Configures

173 The meteorological field for NAQPMS is provided by the Weather Research and Forecasting model version 4.0 (WRFV4.0, 174 https://www.mmm.ucar.edu/models/wrf). The initial-boundary conditions for WRF are obtained from NCEP GDAS Final 175 Analysis (https://rda.ucar.edu/datasets/ds083.3/), with a horizontal resolution of 0.25°×0.25° and the temporal resolution of 6 176 hours, produced by the Global Data Assimilation System (GDAS). The land use data for WRF was updated by USGS's 177 MCD12Q1 v006 in 2019 (https://lpdaac.usgs.gov/products/mcd12q1v006/) with 20 categories. Three nested model domains 178 are conducted with the horizontal resolutions of 45 km in the East Asia region (domain1), 15 km in most areas of China except 179 for the western area (domain2), and 5 km in the Northern China region (domain3, target research region). WRF and NAQPMS 180 have 40 vertical layers with 27 layers within 2 km. The parameterization schemes for physical processes in WRF are shown 181 in Table S2. The boundary condition input for NAQPMS is provided by the global chemistry transport Model for OZone And 182 Related chemical Tracers version 2.4 (MOZART V2.4) (Horowitz et al., 2003). The anthropogenic emissions for NAQPMS 183 are from Tsinghua University's 2016 Multi-resolution Emissions Inventory for China (MEIC, http://www.meicmodel.org/) 184 with a spatial resolution of 0.25°×0.25°, including residential sources, transportation sources, agricultural sources, industrial 185 sources, and power plant sources. The computational platform is the high-performance supercomputer subsystem cluster with 186 320 computation nodes, a total of 12,800 processors, and about 153 TB memory at the Big Data Cloud Service Infrastructure 187 Platform (BDCSIP), which meets the demand for high-performance parallel computing of NAQPMS-PDAF v2.0.

188

The model state variables include  $NH_{4^+}$ ,  $SO_{4^{2^-}}$ ,  $NO_{3^-}$ , OC, EC,  $Na^+$ , Brown carbon, soil  $PM_{2.5}$ , soil  $PM_{10}$ , sea salt, fine dust, coarse dust,  $SO_2$ ,  $NO_2$  and RH. As shown in Fig. 2, the model state has a 4-dimensional (4-D) structure, with longitudinal dimension (ix, 300 grids), latitudinal dimension (iy, 249 grids), variable dimension (ivar, 15), and vertical dimension (iz, 40 layers) in that order. The 4-D model state with 15 variables is converted to a 2-D state matrix in PDAF, the number of grids in the horizontal axis direction is ix, and the number of grids in the vertical axis direction is  $iy^*ivar^*iz$ . Notably, the coordinate index of the 2-D state matrix contains 3-D information for each variable to implement the horizontal and vertical domain





195 localization separately, because the horizontal and vertical resolutions are not uniform. This structure has two advantages. First, 196 the parallel cutting of the horizontal axis enables the local domain to retain the full dimensional information (ix\_p\*iy\*ivar\*iz, 197 where ix p is the longitudinal dimension of the local domain). Secondly, the localization in local domain permits the analysis 198 only executes within a small domain (ix p\*iy) when the length of horizontal localization radius (Rs) is smaller than iy, which 199 effectively reduces the influence of spurious correlations between different state variables. In this study, we set the horizontal 200 and vertical domain localization radius to 200 km (40 grids) and 1 layer. Besides, we further implemented the observation 201 localization to consider the influence of distance between analysis grid and observational grid (see Sect. 2.3.4). To minimize 202 computational complexity, the observation errors were assumed to be spatially isotropic, with 0.40 µg/m<sup>3</sup>, 1.00 µg/m<sup>3</sup>, 0.50 203  $\mu$ g/m<sup>3</sup>, 3.00  $\mu$ g/m<sup>3</sup>, and 0.50  $\mu$ g/m<sup>3</sup> for NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, OC and EC, respectively.



204

205 Figure 2: The structure of state variables in NAQPMS-PDAF v2.0.

# 206 2.3.3 Generation of ensemble members

207 In ensemble DA, ensemble members interpret the uncertainty of the model or system, characterized by BEC, which 208 significantly impacts the DA performance (Dai et al., 2014). For CTMs, emission input directly influences the chemical 209 calculation and substantially contributes to the uncertainty. Perturbing emission input can effectively represent the uncertainty 210 in aerosol emissions and enhance the consistency of ensemble error spread, thereby improving aerosol DA (Huang et al., 2023). 211 CTMs are nonlinear, and model state errors are non-Gaussian distributions. To obtain non-Gaussian error distributions, we 212 followed Kong et al. (2021)'s method to assume that the emission errors are spatially correlated by an isotropic correlation 213 model with the decorrelation length of 150 km and generate perturbation coefficient matrixes with the same Gaussian 214 distribution as the emission species, which are subsequently transformed into non-Gaussian distribution matrixes through non-215 Gaussian process generation v1.2 (https://github.com/ECheynet/Gaussian to nonGaussian/).





216

217 The target PM<sub>2.5</sub> chemical components are NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, OC, and EC, and the perturbed emission species correspondingly 218 include SO<sub>2</sub>, NOx, VOCs, NH<sub>3</sub>, CO, PM<sub>10</sub>, PM<sub>2.5</sub>, EC, and OC, with the corresponding uncertainties ( $\delta$ ) listed in Table 1. As 219 shown in Eq. (1), the original emission input matrix  $(E_p)$  is multiplied by the corresponding perturbation coefficient matrix 220  $(\theta_i)$  to generate the perturbed emission input matrix  $(E_i)$  for each emission species. The calculation of the perturbation 221 coefficient matrix ( $\theta_i$ ) is followed by Eq. (2)-(3). Firstly, N two-dimensional pseudorandom perturbation fields (P<sub>i</sub>) are created 222 using Evensen's method (Evensen, 1994). The uncertainties ( $\delta$ ) of the emission species are incorporated into the two-223 dimensional pseudorandom perturbation fields (P<sub>i</sub>) to obtain the final perturbation coefficient matrixes ( $\theta_i$ ). Finally, the 224 Gaussian-distribution perturbation coefficient matrixes ( $\theta_i$ ) were transformed into non-Gaussian distribution coefficient 225 matrixes with a given target skewness (set to 1) and kurtosis (set to 6) by non-Gaussian process generation v1.2, which employs 226 the Moment Based Hermite Transformation Model and a cubic transformation. 227 Table 1: The uncertainties of emission species in NAQPMS-PDAF v2.0 Species  $SO_2$ NOx VOCs NH<sub>3</sub> CO  $PM_{10}$ PM<sub>2.5</sub> EC OC Uncertainty  $\delta$ 2.00 0.31 0.53 0.70 2.58 0.68 1.32 1.30 2.08

228 
$$E_i = E_p \times \theta_i, i = 1, 2, ..., N$$
, (1)

229 
$$\ln \theta_{o_{i}} = \left(\frac{(P_{i} - \frac{1}{N} \times \sum_{i=1}^{N} P_{i})}{\sqrt{\frac{1}{N} \times \sum_{i=1}^{N} (P_{i} - \frac{1}{N} \times \sum_{i=1}^{N} P_{i})^{2}} - \frac{1}{2} \times \ln(1 + \delta^{2})\right) \times \sqrt{\ln(1 + \delta^{2})} , \qquad (2)$$

$$230 \qquad \theta_{i} = \frac{\left(\theta_{o_{i}} - \frac{1}{N} \times \sum_{i=1}^{N} \theta_{o_{i}}\right)}{\sqrt{\frac{1}{N} \times \sum_{i=1}^{N} \theta_{o_{i}}}^{1} \times \sum_{i=1}^{N} \theta_{o_{i}}\right)} \times \left(\frac{1}{N} \times \sum_{i=1}^{N} \theta_{o_{i}}\right) \times \delta + \frac{1}{N} \times \sum_{i=1}^{N} \theta_{o_{i}} , \qquad (3)$$

Notably, all matrix operations involved are Schur Product. Where  $E_i$  denotes the i<sup>th</sup> ensemble perturbed emission input matrix,  $E_p$  denotes the original unperturbed emission input matrix and  $\theta_i$  represents the i<sup>th</sup> ensemble perturbation coefficient matrix.  $\theta_{o_i}$  is the i<sup>th</sup> ensemble original perturbation coefficient matrix, which is obtained by mathematical transformation of the i<sup>th</sup> ensemble pseudorandom perturbation matrix  $P_i$ , including standardization, scaling by uncertainty ( $\delta$ ), and logarithm.

#### 235 2.3.4 Hybrid nonlinear DA algorithm with adaptive forgetting factor

236 To thoroughly integrate the stability of EnKFs with the nonlinearity of nonlinear filters and be ideal for the nonlinear and non-

237 Gaussian-distribution situations, the hybrid LKNETF is used in this study. This section reviews the algorithms of LETKF,

- 238 LNETF, and their combination (LKNETF).
- 239
- 240 ETKF, a deterministic filter in EnKFs, efficiently obtains analysis samples using a transformation matrix and the square root

of the forecast error covariance (Bishop et al., 2001). In contrast to stochastic filters in EnKFs, ETKF prevents underestimation

- 242 of the analysis error covariance resulting from the random observation perturbations. And it is particularly applicable in
- 243 situations with small ensemble sizes (Lawson and Hansen, 2004). The realization of ETKF can be divided into the forecast





and analysis steps.

245 246 In the forecast step, the forecast state vector  $(\mathbf{x}_t^f)$  at t is generated by numerical model (**M**) integration of the analysis state 247 vector  $(\mathbf{x}_{t-1}^a)$  at t-1. The forecast error covariance matrix ( $\mathbf{P}_t^f$ ) can be calculated by the perturbation of the forecast ensemble 248  $(\mathbf{X}_t^{f'})$ .

249 
$$\mathbf{x}_{t}^{f} = \mathbf{M}(\mathbf{x}_{t-1}^{a}), \mathbf{X}_{t}^{f} = [\mathbf{x}_{1t}^{f}, \mathbf{x}_{2t}^{f}, ..., \mathbf{x}_{Kt}^{f}],$$
 (4)

$$\mathbf{250} \qquad \mathbf{P}_{t}^{f} = \mathbf{X}_{t}^{f'} \mathbf{X}_{t}^{f'^{\mathrm{T}}} , \qquad (5)$$

251 Where  $\mathbf{X}_{t}^{f}$  is the forecast ensemble at t, and K is the number of ensemble members.  $\mathbf{X}_{t}^{f'}$  is the perturbation of the forecast 252 ensemble at t, calculated by  $\mathbf{X}_{t}^{f}$  and the forecast ensemble mean  $\overline{\mathbf{X}_{t}^{f}}$  at t.

253

In the analysis step, the forecast error covariance matrix ( $\mathbf{P}_{t}^{f}$ ) at t is transformed to the analysis error covariance matrix ( $\mathbf{P}_{t}^{a}$ ) at t by a transform matrix (**T**).

$$256 \qquad \mathbf{P}_{t}^{a} = \mathbf{X}_{t}^{f'} \mathbf{T} \mathbf{X}_{t}^{f'^{T}} , \qquad (6)$$

The transform matrix (**T**) is defined as follows and can be decomposed to a left singular vector matrix (**U**), a singular value matrix (**S**), and a right singular vector matrix (**V**) through the singular value decomposition.

259 
$$\mathbf{T}^{-1} = \rho_{adaptive} (\mathbf{K} - 1)\mathbf{I} + (\mathbf{H}\mathbf{X}_{t}^{f'})^{\mathrm{T}} (\mathbf{L} \cdot \mathbf{R}^{-1}) \mathbf{H}\mathbf{X}_{t}^{f'} = \mathbf{U}\mathbf{S}\mathbf{V} , \qquad (7)$$

$$260 \qquad \rho_{adaptive} = \frac{\sigma_{ens}^2}{\sigma_{resid}^2 - \sigma_{obs}^2} , \tag{8}$$

Where  $\rho_{adaptive}$  is an adaptive forgetting factor, used for the inflation of error covariance estimation (the initial  $\rho_{adaptive}$  is set to 0.9 in this study).  $\sigma_{ens}^2$  is the mean ensemble variance,  $\sigma_{resid}^2$  is mean of observation-minus-forecast residual,  $\sigma_{obs}^2$  is mean observation variance. I is the identity matrix. H is the observation operator. L is the localization matrix, a weight matrix calculated by the 5<sup>th</sup>-order polynomial (Nerger, 2015), implemented in LETKF for observation localization analysis to avoid observational spurious correlation and filter divergence effectively (Hunt et al., 2007). R is the observation error covariance matrix.

267

The analysis state vector 
$$(\mathbf{x}_t^a)$$
 at t is calculated by the forecast state vector  $(\mathbf{x}_t^f)$  at t, the perturbation of the forecast ensemble  
( $\mathbf{x}_t^{f'}$ ) at t and a weight vector ( $\mathbf{w}$ ).

$$270 \qquad \mathbf{x}_{t}^{a} = \mathbf{x}_{t}^{f} + \mathbf{X}_{t}^{f'} \mathbf{w} , \qquad (9)$$

271 The weight vector (**w**) is given by the following equation.





272
$$\mathbf{w} = \mathbf{T} (\mathbf{H} \mathbf{X}_{1}^{\ell})^{\mathsf{T}} (\mathbf{L} \cdot \mathbf{R}^{-1}) (\mathbf{y} - \mathbf{H} \mathbf{x}_{1}^{\ell})$$
, (10)273Where  $\mathbf{y}$  is observations.274275The analysis ensemble  $(\mathbf{X}_{1}^{*})$  at t can be obtained by forecast ensemble mean  $(\overline{\mathbf{X}_{1}^{\ell}})$  at t, the perturbation of the forecast ensemble276 $(\mathbf{X}_{1}^{\ell})$  at t and a transform matrix ( $\mathbf{C}$ ) represented by the symmetric square root of  $\mathbf{T}$ .277 $\mathbf{x}_{1}^{*} = \overline{\mathbf{x}_{1}^{*}} + \sqrt{\mathbf{K} - 1} \mathbf{X}_{1}^{\ell} \mathbf{C}$ , (11)278The transform matrix ( $\mathbf{C}$ ) is calculated as follows.279 $\mathbf{C} = \mathbf{U}\mathbf{S}^{-1/2}\mathbf{U}^{\mathsf{T}}$ , (12)280NETF is a 2<sup>nd</sup>-order exact ensemble square root filter effectively applied to the nonlinear and non-Gaussian DA (Tödter and281Ahrens, 2015). Like PF, NETF indirectly updates the model state by using observations to affect the weights of the prior282ensemble. However, PF and NETF differ in the sampling method. PF utilizes the Monte Carlo and Bayesian methods to283calculate particle weights based on observations, which are then used to generate the analysis ensemble by weighting the284resampling forecast ensemble. In high-dimensional systems, as the DA progresses, the weight differences of particles increase,285with most particles having weights close to 0, leading to filter degeneration. In contrast, NETF generates the analysis ensemble289 $\mathbf{x}^{*} = \frac{1}{\mathbf{x}} \sum_{i=1}^{K} \mathbf{u}_{i} \mathbf{x}_{i}^{\ell}$ ,  $(13)$ 290Where  $\mathbf{x}^{*}$  is the analysis state vector mean, K is the number of ensemble members,  $\mathbf{x}_{i}^{*}$  is the i<sup>h</sup> analysis state vector,  $\mathbf{w}_{i}$  is291the i<sup>h</sup> particle weight vector in PF, which is calcu

293 
$$\mathbf{P}^{a} = \frac{1}{K-1} \sum_{i=1}^{K} (\mathbf{x}_{i}^{a} - \bar{\mathbf{x}}^{a}) (\mathbf{x}_{i}^{a} - \bar{\mathbf{x}}^{a})^{\mathrm{T}} = \sum_{i=K}^{K} w_{i} (\mathbf{x}_{i}^{f} - \bar{\mathbf{x}}^{f}) (\mathbf{x}_{i}^{f} - \bar{\mathbf{x}}^{f})^{\mathrm{T}} , \qquad (14)$$

Where P<sup>a</sup> is the error covariance matrix of the analysis ensemble, calculated by the perturbation of the analysis ensemble.
In NETF, A performs as a transform matrix like the transform matrix (T) in ETKF, which can be obtained from the weight matrix (w).

$$P^{a} = \mathbf{X}^{f'} \mathbf{A} \mathbf{X}^{f'^{T}} , \qquad (15)$$

298 
$$\mathbf{A}^{1/2} = (\mathbf{W} - \mathbf{w}\mathbf{w}^{\mathrm{T}})^{1/2} = \mathbf{V}\mathbf{D}^{1/2}\mathbf{V}^{\mathrm{T}}$$
, (16)





- Where the matrix  $\mathbf{W} \equiv \text{diag}(\mathbf{w})$  is defined as a diagonal matrix created from the weight matrix ( $\mathbf{w}$ ). A can be decomposed ( $\mathbf{A} = \mathbf{V}\mathbf{D}\mathbf{V}^{T}$ ) by a singular value decomposition as it is a real, symmetric, positive semidefinite matrix. V is the orthogonal matrix, and **D** is a diagonal matrix.
- 302

Then, the perturbation of the analysis ensemble  $(X^{a'})$  and the analysis ensemble  $(X^{a})$  can be obtained by applying the square root of **A** as a transform matrix.

305 
$$\mathbf{X}^{a'} = \sqrt{\mathbf{K}} \mathbf{X}^{t'} \mathbf{A}^{1/2}$$
, (17)

$$306 \qquad \mathbf{X}^{\mathrm{a}} = \overline{\mathbf{X}}^{\mathrm{f}} + \mathbf{X}^{\mathrm{f}'} \left( \overline{\mathbf{W}} + \sqrt{\mathrm{K}} \mathbf{A}^{1/2} \right) , \tag{18}$$

307 LKNETF combines the LETKF and LNETF through a hybrid weight  $\gamma$  to perform better in systems with different non-308 linearity degrees and implement in situations with smaller ensemble sizes (Nerger, 2022). When  $\gamma$  approaches 1, the analysis 309 increment ( $\Delta X_{\text{LETKF}}$ ) computed by LETKF becomes more significant and appropriate for linear systems with Gaussian 310 distributions. Conversely, when  $\gamma$  approaches 0, the analysis increment ( $\Delta X_{\text{LNETF}}$ ) computed by LNETF becomes more 311 significant and appropriate for non-linear systems with non-Gaussian distributions. The one-step update scheme is used in this 312 study.

313 
$$\mathbf{X}_{\text{HSync}}^{a} = \overline{\mathbf{X}}^{f} + (1 - \gamma)\Delta \mathbf{X}_{\text{LNETF}} + \gamma\Delta \mathbf{X}_{\text{LETKF}} , \qquad (19)$$

314 2.4 Data

#### 315 2.4.1 Observation

316 The one-month (February 2022) hourly mass concentration observations of five PM<sub>2.5</sub> chemical components (NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, 317 NO3-, OC, and EC) from 33 ground-based sites in Northern China and surrounding areas were collected for this work (Fig. 3). 318 Out of the 33 sites, 24 (DA sites) were utilized for DA and internal validation, and the remaining 9 (VE sites) were used for 319 independent verification to assess the influence of DA sites on neighboring areas. These sites were divided using the K-means 320 clustering algorithm (Lloyd, 1982; Arthur and Vassilvitskii, 2007). The supplement provides a detailed description (Text S1). 321 PM2.5 hourly observations from the China National Environmental Monitoring Centre (CNEMC, http://www.cnemc.cn/) were 322 employed to assess the overall mass concentration of PM2.5 chemical components in NAQPMS-PDAF v2.0. Due to incomplete 323 spatial overlap between the PM2.5 sites and the chemical component sites, the PM2.5 sites were selected based on the closest 324 coordinate Euclidean distance between PM2.5 sites and chemical component sites.









Figure 3: The model domains in WRF simulation (a) and the location of observations (b). The domain 3 in (a) is the target area in this study. Twenty-four red sites in (b) represent the sites for data assimilation, and nine green sites in (b) represent the sites for spatial independent validation. The topographic dataset is from the ETOPO1 1 arc-minute Global Relief Model, taken from the National Geophysical Data Center (Amante and Eakins, 2009).

# 330 2.4.2 Global reanalysis dataset

- 331 The global reanalysis datasets of PM2.5 chemical components in February 2022 were obtained from the Copernicus Atmosphere 332 Monitoring Service ReAnalysis (CAMSRA, 0.75°×0.75°) (Inness et al., 2019) and the Modern-Era Retrospective analysis for 333 Research and Applications, Version 2 (MERRA-2, 0.5°×0.625°) (Randles et al., 2017) to compare with reanalysis dataset 334 generated by NAQPMS-PDAF v2.0. For the consistency of data comparison, the global reanalysis surface grid data located in 335 the observation sites of PM2.5 chemical component were extracted through the k-nearest neighbor search method (Friedman et 336 al., 1977), which can efficiently match grid points and observation sites based on longitude and latitude data and Euclidean 337 distances. Our 3-hourly NAQPMS-PDAF v2.0 output of NO3<sup>-</sup> and NH4<sup>+</sup> were extracted to compare with the CAMSRA dataset, 338 and hourly NAQPMS-PDAF v2.0 output of SO42-, OC, and EC were extracted to compare with MERRA-2 M2T1NXAER 339 dataset.
- 340 2.5 Experimental setting and evaluation method
- In our study, four tests were conducted to evaluate the performance of NAQPMS-PDAF v2.0 with hourly observations of five PM<sub>2.5</sub> chemical components, including (1) the dependence on ensemble size and assimilation frequency, (2) the interpretation ability on mass concentration and spatiotemporal characteristics, (3) the superiority compared to other reanalysis dataset, and (4) the uncertainty in ensemble assimilation. In practice, the ratio of ensemble size to the number of processes with 1:50 in high-performance computers was the best parallel scheme to balance computing efficiency and computing resources (Wang et al., 2022).
- All the tests were run in NAQPMS-PDAF v2.0 after a spin-up experiment with 24 timesteps from 00:00 to 23:00 (LST) on February 1<sup>st</sup>, 2022. (1) For the first test, we assimilated the hourly observations of five PM<sub>2.5</sub> chemical components from all sites with 48 timesteps from 00:00 (LST) on February 2<sup>nd</sup> to 23:00 (LST) on February 3<sup>rd</sup>, 2022. In the first situation, we





- 351 controlled a fixed assimilation frequency of 1 hour and changed the ensemble size to 2, 5, 10, 15, 20, 30, 40, and 50. In the 352 second situation, we controlled a fixed ensemble size of 20 and changed the assimilation frequency to 1 hour, 2 hours, 3 hours, 353 4 hours, 5 hours, 6 hours, 8 hours, and 12 hours. (2) For the second test, we set an ensemble size of 20 and an assimilation 354 frequency of 1 h and assimilated the hourly observations of five PM2.5 chemical components from DA sites with 648 timesteps 355 from 00:00 (LST) on February 2nd to 23:00 (LST) on February 28th, 2022. We also conducted a free running (FR) experiment 356 without assimilation in the same period for comparison. (3) For the third test, we followed the settings in the second test but 357 assimilated the observation from all sites to generate a high-quality reanalysis dataset of five PM2.5 chemical components. (4) 358 The last test was like the first but with a different situation to investigate the impact of ensemble perturbation on ensemble 359 assimilation. From Table 2, we fixed species uncertainty (M4 setting) with five distribution types in the first situation and fixed
- $360 \qquad distribution type (T2 \ setting) \ with \ five \ SO_2 \ uncertainties \ in \ the \ second.$

Experiment	Distribution (Fixed species uncertainty)		
T1	Gaussian		
T2	Non-Gaussian (m3=1, m4=6)		
Т3	Non-Gaussian (m3=-1, m4=6)		
T4	Non-Gaussian (m3=1, m4=12)		
Т5	Non-Gaussian (m3=-1, m4=12)		
	SO <sub>2</sub> uncertainty (Fixed distribution)		
M1	12%		
M2	50%		
M3	100%		
M4	200%		
M5	300%		

361 Table 2: The experiment settings for emission perturbation

362

363 We used the Continuous Ranked Probability Score (CRPS) to evaluate ensemble size dependency, which measures the 364 consistency between ensemble forecast distribution and corresponding observations (Jolliffe and Stephenson, 2012). The 365 calculation rules are referred to in Hersbach's study (Hersbach, 2000). Besides, four common statistical indicators, the Pearson 366 correlation coefficient (CORR), root mean square error (RMSE), mean absolute error (MAE), and coefficient of determination 367  $(R^2)$ , were used to assess the DA system performance in interpreting PM<sub>2.5</sub> chemical components  $(SO_4^{2-}, NO_5^{-}, NH_4^{+}, OC, and$ 368 EC). The CORR measures the correlation between the system outputs and corresponding observations, the RMSE and MAE 369 indicates the overall system accuracy, and the R<sup>2</sup> reflects the proportion of variability in the observations explained by the 370 assimilation system.





#### 371 3 Results and discussion

#### 372 3.1 The Dependence on Ensemble Size and Assimilation Frequency for Five Components

373 Ensemble size is a crucial parameter in ensemble assimilation, determining the model state's uncertainty range. A larger 374 ensemble size more accurately represents the error distribution of state variables but requires considerable computing resources 375 and time, especially for high-dimension systems. A smaller ensemble size can easily lead to underestimating the error 376 covariance matrix, especially for the fine-resolution model (Kong et al., 2021). Thus, identifying an appropriate ensemble size 377 to balance computational efficiency and accuracy is the primary step in ensemble DA. A prior study (NAQPMS-PDAF v1.0) 378 only evaluated the correlation between ensemble size and parallel efficiency and concluded that the ratio of ensemble size to 379 high-performance computing processors was 1:50 (Wang et al., 2022), while the impact of ensemble size on the accuracy and 380 computational efficiency was neglected. In this study, we assessed the NAQPMS-PDAF v2.0 dependency on ensemble size 381 through three statistical indicators (CRPS, RMSE, and CORR).

382

383 From Fig. 4a, when the ensemble size is at its minimum level of 2, the mean CRPS values of the five PM<sub>2.5</sub> chemical 384 components are more significant, with NO3<sup>-</sup> exhibiting the most considerable difference between the simulation distribution 385 and observations (more than 4). With each increase in ensemble size, the mean CRPS values of the five chemical components 386 progressively reduce and eventually reach convergence when the ensemble size is 10, implying that a hybrid nonlinear filter 387 can maintain high accuracy and reliability in ensemble assimilation with an ensemble size that is smaller than the traditional 388 minimum of 20 ensemble members, as observed in prior ensemble assimilation studies (Constantinescu et al., 2007; Miyazaki 389 et al., 2012; Schwartz et al., 2014; Rubin et al., 2017; Kong et al., 2021; Tsikerdekis et al., 2021; Wang et al., 2022), including 390 NAQPMS-PDAF v1.0. The mean CRPS value of EC is the lowest among the five chemical components, indicating the highest 391 accuracy and reliability of EC ensemble DA. The performance of other components is similar. Like CRPS values, the values 392 of RMSE and CORR decrease and increase, respectively, as the ensemble size increases, and convergence begins to occur 393 when the ensemble size is 10 (Fig. 4b and c). Compared with other chemical components, the CORR value of SO42- is 394 significantly lower, less than 0.8, possibly due to its estimated background field error covariance driven by the inadequate 395 ensemble perturbations. Therefore, in the Discussion section, we deeply discuss the uncertainties of ensemble perturbations.







396

Figure 4: Assessment of ensemble size dependency based on mean continuous ranked probability score (CRPS) (a), root mean square
 error (RMSE) (b), correlation coefficient (CORR) (c), and time (d).

399 Figure 4d shows the time required for the four processes of ensemble assimilation under different ensemble sizes, including 400 initialization, model integration, assimilation, and post-processing. The model integration process in NAQPMS-PDAF v2.0 401 takes the longest, followed by post-processing, initialization, and assimilation. The required time for initialization and post-402 processing increases with increasing ensemble size, while for model integration and assimilation, except for ensemble size 30, 403 the required time is the same under different ensemble sizes. Generally, the time needed for ensemble sizes of 30-50 is 404 considerably higher than that for smaller ones. Although convergence occurs with an ensemble size of 10, our work illustrates 405 a similar time required between ensemble sizes 10 and 20. Consequently, we selected an ensemble size of 20 to ensure optimal 406 performance of NAQPMS-PDAF v2.0, considering both assimilation efficiency and accuracy.

407

The assimilation frequency is the interval at which observational data is introduced into the DA system, directly affecting the practical assimilation data volume and computation cost. High-frequency DA with high-quality observations is crucial for improving numerical simulations and forecasts (Liu et al., 2021). Figure 5 demonstrates that the MAE values of the five chemical components analysis fields range from 0.02 to 0.12  $\mu$ g/m<sup>3</sup>, RMSE values range from 0.23 to 2.61  $\mu$ g/m<sup>3</sup>, and CORR values range from 0.71 to 0.98 at a 1-hour assimilation time interval, which is significantly better than the statistical indicators at lower assimilation frequencies. Even at a 2-hour assimilation frequency, the assimilation effect drops sharply compared to the 1-hour interval, especially for NO<sub>3</sub><sup>-</sup>, OC, and EC. The values of MAE and RMSE increase by 2.6-5.82  $\mu$ g/m<sup>3</sup> and 4.72-





415 9.57 µg/m³, respectively, and the CORR values decrease by 0.27-0.81. Gradual increasing trends in MAE and RMSE values 416 and a slight decreasing trend in CORR values are observed as assimilation frequency decreases from the 2-hour interval. 417 Therefore, the fast-updating assimilation with a 1-hour interval significantly improves the NAQPMS simulation. For the 418 forecasting field (Fig. S2), the low sensitivity of state variables to assimilation frequency suggests that NAQPMS-PDAF v2.0 419 can appropriately reduce assimilation frequency during the actual forecasting phase, lowering the demand for high temporal 420 resolution observations and computational resources.



421

Figure 5: Assessment of assimilation interval dependency based on mean absolute error (MAE) (a), root mean square error (RMSE)
(b), and correlation coefficient (CORR) (c) at the analysis step.

# 424 **3.2 Evaluation of NAQPMS-PDAF v2.0 performance**

#### 425 **3.2.1** Overall validation of DA results

426 We conducted a control experiment (free-running field, FR) without any DA and a DA experiment. This section verified the 427 forecast filed (FOR) and analysis field (ANA) at 24 DA sites and 9 VE sites, respectively. Figure 6 shows the scatter distribution 428 of observations and simulations at DA sites. For FR (Fig. 6a1-a5), five chemical components have CORR values ranging from 429 0.32 to 0.56, and R<sup>2</sup> values do not exceed 0.3, indicating poor consistency between observations and simulations. In detail, the 430 simulated mass concentrations of SO42-, OC, and EC are significantly overestimated, while the simulated concentrations of 431  $NH_4^+$  and  $NO_3^-$  are underestimated. OC has the most significant error, with an RMSE value of 25.84  $\mu$ g/m<sup>3</sup> and an MAE value 432 of 19.41  $\mu$ g/m<sup>3</sup>. Besides, the error distributions of SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> are close to a symmetric distribution with a mean 433 value of 0, while the error distributions of OC and EC are skewed to the left from the mean value of 0 (Fig. 7a1-a5), showing 434 the relatively better simulations in SO42-, NO3- and NH4+ than in OC and EC. Overall, NAQPMS cannot interpret the mass 435 concentrations of the five chemical components with significant errors, mainly due to the uncertainties in chemical mechanisms 436 (Miao et al., 2020).







437

438 Figure 6: Scatterplots of the DA-site simulations versus the DA-site observations with probability density for the free-running field

439 (FR, a1-a5), forecast field (FOR, b1-b5), and analysis field (ANA, c1-c5). The dotted gray lines represent the 2:1, 1:1, and 1:2 lines, 440 and the solid red line represents the fitting regression line.







Figure 7: Probability distributions of bias between DA-site observations and DA-site simulations for the free-running field (FR, a1-443 a5), forecast field (FOR, b1-b5), and analysis field (ANA, c1-c5).

444 After DA, FOR shows a slight improvement with a slight increase in CORR and R<sup>2</sup> and a decrease in RMSE and MAE, 445 especially for NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub> (Fig. 6b1-b5). Although SO<sub>4</sub><sup>2-</sup>, OC, and EC are significantly overestimated with a slight decrease 446 in CORR and R<sup>2</sup>, the RMSE and MAE values decrease. Besides, the error distributions of the five chemical components are 447 concentrated at 0, and the overestimation of OC and EC has been improved compared to FR (Fig. 7b1-b5). These results 448 indicate that DA reduces the overall FOR errors in NAQPMS due to improved forecasting ability by obtaining optimal initial





449 fields. However, further improvements are necessary to address the NAQPMS uncertainties in emission sources, 450 meteorological input, and imperfect physiochemical mechanisms. For ANA (Fig. 6c1-c5), DA significantly improves the 451 simulations of the five chemical components, making the ANA consistent with the observations. The CORR values are not 452 less than 0.86, the RMSE and MAE values do not exceed 3.23 µg/m<sup>3</sup> and 1.49 µg/m<sup>3</sup>, respectively, and the R<sup>2</sup> values are not 453 less than 0.74. Specifically, the CORR values for NO3, OC, and EC are not less than 0.96, and the R<sup>2</sup> values are not less than 454 0.93. The error distributions of the five chemical components concentrate to 0 with the mean bias ranging from  $0\pm0.08 \ \mu g/m^3$ 455 to 1.02±3.07 µg/m<sup>3</sup> (Fig. 7c1-c5). The results of VE sites show similar characteristics to the DA sites (Fig. S3 and S4). 456 Compared to FR, the overall errors of the FOR and ANA for the five chemical components decrease with a significant 457 improvement in ANA, showing that the CORR values of NH4<sup>+</sup> and NO3<sup>-</sup> increase by 0.15 and 0.45, respectively, the R<sup>2</sup> values 458 of NH4<sup>+</sup> and NO3<sup>-</sup> increase by 0.22 and 0.81, respectively, the RMSE values of OC and EC decrease by 21.77 µg/m<sup>3</sup> and 17.79 459 µg/m3, respectively. Overall, the FOR and ANA errors decreased significantly. The ANA of the five chemical components at 460 DA sites is almost entirely consistent with the observations, indicating excellent DA performance.

# 461 **3.2.2** Assessment of temporal variation in chemical components

462 The ensemble DA employs a cyclic updating process wherein the forecast and analysis steps are continuously completed at 463 each iteration (Evensen, 2003; Houtekamer and Zhang, 2016). In the forecast step, the ANA at the current time step serves as 464 the optimal initial field to advance the model integration and obtain the FOR at the next step. In the analysis step, the FOR at 465 the next time step provides background field information for the subsequent DA analysis to generate the ANA at the next time 466 step. The FOR and ANA interact with each other in the temporal dimension. Therefore, in this section, we assess the ability of 467 NAQPMS-PDAF v2.0 to interpret the temporal variations of the five chemical components. Figure 8 illustrates the time series 468 of the five chemical components at two representative sites, including a DA site in Tianjin City and a VE site in Heze City. For 469 the DA site (Fig. 8a), the temporal variations of NH4+ and NO3- in FR and FOR exhibit better agreement with the observed 470 temporal variations (OBS) than those of SO<sub>4</sub><sup>2-</sup>, OC, and EC. However, NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> mass concentrations are significantly 471 lower than the high-value mass concentrations observed on February  $25^{\text{th}}$ . The mass concentration of  $SO_4^{2-}$  in FR is greatly 472 overestimated during the periods of Feb. 8th-11th, Feb. 18th-19th, and Feb. 24th-25th. The mass concentrations of OC and EC in 473 FR are overestimated throughout February with substantial temporal fluctuations. Although the time series of SO42-, OC, and 474 EC in FOR show some improvement, noticeable differences from the OBS are still apparent. After DA, the ANA time series 475 for the five chemical components align well with the OBS, indicating good consistency and accurate representation of temporal 476 characteristics, such as the NH<sub>4</sub>NO<sub>3</sub> pollution captured on February 25th. Notably, the mass concentrations of SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, and 477 NH4<sup>+</sup> peaked on Feb. 8<sup>th</sup>-11<sup>th</sup> and February 25<sup>th</sup>, indicating intensified atmospheric secondary chemical reactions primarily due 478 to neutralization reactions of acidic pollutants capturing NH<sub>3</sub>. The temporal variations of NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> are more similar 479 because atmospheric NO3<sup>-</sup> primarily exists as NH4NO3 rather than other metal nitrates, and NH4NO3 can form before the





- 480 complete neutralization of H<sub>2</sub>SO<sub>4</sub> (Ge et al., 2017). The improvements at the VE site (Fig. 8b) are like those at the DA site,
- 481 with the ANA time series of the five chemical components showing closer agreement with the OBS, which suggests that the
- 482 localization analysis in DA effectively facilitates the propagation of observations within a specific spatial range and mitigates
- 483 the assimilation anomalies caused by spurious correlations from the distant sites (Hunt et al., 2007).



484

485 Figure 8: Hourly variation of five PM2.5 chemical components in a representative DA site (a) and a representative VE site (b).

486 NH4<sup>+</sup>, SO4<sup>2-</sup>, NO3<sup>-</sup>, OC, and EC are critical chemical components of PM2.5, and the sum of their mass concentrations can be 487 approximated as the PM2.5 mass concentration. We further assessed the simulation enhancement of PM2.5 time series based on 488 ground-level PM2.5 observations. Six representative sites were selected, including 3 DA sites (Fig. 9a1-a3) and 3 VE sites (Fig. 489 9b1-b3). The FR and FOR in DA and VE sites show significant overestimation and poor consistency with the OBS, mainly 490 due to the overestimation of OC and EC mass concentrations. Conversely, the PM2.5 time series in ANA closely matches that 491 of the OBS, accurately capturing the actual variation of PM2.5. In some specific instances, such as on February 26th at 00:00 in 492 Tianjin City and Langfang City, the peak value of ANA was lower than that of OBS, which could be attributed to the negligence 493 of other PM<sub>2.5</sub> components (such as mineral dust and sea salt) and the inconsistency in location between ground-level PM<sub>2.5</sub> 494 observational sites and chemical components observational sites. Overall, the DA of chemical component observations 495 significantly enhanced the simulation of PM2.5 time series in NAQPMS. Compared to the CORR values of FR and FOR, the 496 CORR values of ANA at the six representative sites increased by 13.64%-89.58% and 17.19%-75.00%, respectively, while the 497 RMSE values decreased by 56.03%-83.13% and 40.74%-72.20% (Table S3).











# 500 3.2.3 Assessment of spatial distribution in chemical components

501 DA can improve the interpretation of model states in the analysis domain by using a limited number of observations. The

502 ability to represent spatial distribution accurately is a crucial performance for aerosol DA. Figure 10 displays the spatial

503 distribution of the monthly average mass concentrations for the five chemical components, including OBS, FR, FOR, ANA,

504 and analysis increment (INC). The spatial distributions of bias and statistical indicators for FR, FOR, and ANA are shown in

505 Fig. 11 and Fig. 12, respectively.





Figure 10: Spatial concentration distribution of site observation (OBS, a1-e1), free-run field (FR, a2-e2), forecast field (FOR, a3-e3),
 analysis field (ANA, a4-e4), and increment (INC) between ANA and FR (a5-e5) for five PM<sub>2.5</sub> chemical components.







509

510 Figure 11: Spatial distribution of DA-site bias for five PM2.5 chemical components from observation (OBS) for the free-running field



511 (FR, a1-e1), forecast field (FOR, a2-e2) and analysis field (ANA, a3-e3).

512

Figure 12: Spatial distribution of DA-site statistical indictors for five PM<sub>2.5</sub> chemical components. (a1-e1) represents the values of RMSE and CORR for the forecast field (FOR), (a2-e2) same as (a1-e1) but for analysis field (ANA), (a3-e3) represents the improvement of RMSE and CORR for the forecast field (FOR), (a4-e4) same as (a3-e3) but for analysis field (ANA). The size represents the value of RMSE in (a1-e2) and the improvement percentage compared to non-assimilation in (a3-e4), respectively.

The spatial characteristics of  $NH_4^+$  and  $NO_3^-$  are similar. Compared to the OBS (Fig. 10a1 and c1), the FR (Fig. 10a2 and c2) and FOR (Fig. 10a3 and c3) have failed to capture the high-value mass concentrations in the border area between Hebei province, Shanxi province, Henan province, and Shandong province, especially in the northern region of Henan province. The primary reason is the uncertainties in emission inventories in winter heating periods, which results in insufficient emission statistics of gaseous precursors NOx and NH<sub>3</sub> (Aleksankina et al., 2018). After DA, this situation is significantly improved





522 with the ANA (Fig. 10a4 and c4). The INCs in the Beijing-Tianjin-Hebei region, Shanxi province, Henan province, and 523 Shandong province are positive (Fig. 10a5 and c5), indicating varying degrees of improvement in correcting the 524 underestimation of mass concentrations. Specifically, for NH4+ and NO3- at DA sites, the biases between the OBS and ANA 525 are significantly reduced compared to the biases between the OBS and FR (Fig. 11), with the mean absolute bias decreasing 526 by 0.93 µg/m<sup>3</sup> and 4.27 µg/m<sup>3</sup>, respectively. Moreover, the overall biases at VE sites also decrease (Fig. S5). As for the spatial 527 statistical indicators of NH<sub>4</sub><sup>+</sup> (Fig. 12a1 and a2), the CORR values in FOR and ANA range from 0.39 to 0.79 and 0.70 to 0.97, 528 respectively, and the RMSE values range from 3.16 µg/m3 to 7.65 µg/m3 and 1.20 µg/m3 to 3.49 µg/m3, respectively. As for 529 the spatial statistical indicators of NO3<sup>-</sup> (Fig. 12c1 and c2), the CORR values in FOR and ANA range from 0.09 to 0.76 and 530 0.89 to 0.99, respectively, and the RMSE values range from 4.88  $\mu$ g/m<sup>3</sup> to 15.69  $\mu$ g/m<sup>3</sup> and 1.34  $\mu$ g/m<sup>3</sup> to 5.39  $\mu$ g/m<sup>3</sup>, 531 respectively. For the FOR, the improvement in accuracy for NO3 is more significant than that for NH4+, with the CORR values 532 of most DA sites increasing by more than 10% and the RMSE of most DA sites decreasing by not less than 10% (Fig. 12a3 533 and c3). For the ANA, NH4<sup>+</sup>, and NO3<sup>-</sup> exhibit significant improvements in CORR and RMSE, as most DA sites show over 534 150% in CORR and over 50% in RMSE (Fig. 12a4 and c4). The improvements can also be found for NH4<sup>+</sup> and NO3<sup>-</sup> at VE sites (Fig. S6). The spatial consistency of NH4+ and NO3- indicates that NH4NO3 is the primary aerosol chemical component, 535 536 highlighting the necessity of coordinated control of precursor NOx and NH<sub>3</sub>.

537

538 Unlike NH4<sup>+</sup> and NO3<sup>-</sup>, compared to the OBS (Fig. 10b1), the mass concentrations of SO4<sup>2-</sup> in the FR and FOR (Fig. 10b2 and 539 b3) are significantly overestimated, especially in Shandong province. In contrast, the ANA has dramatically improved (Fig. 540 10b4), with most areas showing negative INCs (Fig. 10b5). The mean absolute biases in DA and VE sites have decreased by 541 1.80 µg/m3 and 2.68 µg/m3, respectively (Fig. 11 and Fig. S5). Specifically, after DA, the CORR values of the FOR and ANA range from 0.22 to 0.71 and 0.58-0.97, and the RMSE values range from 3.42 µg/m<sup>3</sup> to 11.07 µg/m<sup>3</sup> and 1.20 µg/m<sup>3</sup> to 4.30 542 543 µg/m3, respectively (Fig. 12b1 and b2). The CORR and RMSE values in FOR have significantly improved (Fig. 12b3) at DA 544 sites around Beijing. While the CORR values in ANA have increased by more than 13%, with most DA sites showing an 545 increase of over 50%, and RMSE values have decreased by no less than 30%, with most DA sites showing a decrease of over 546 70% (Fig. 12b4). Besides, half of the VE sites show significant improvement in the CORR and RMSE in the FOR and ANA, 547 mainly due to their proximity to more DA sites (Fig. S6). The OBS and ANA indicate a considerable control in  $SO_4^{2-}$  pollution 548 during the winter heating period due to the emission reduction of gaseous precursors (Zhai et al., 2019; Yan et al., 2021).

549

The spatial distributions of OC and EC exhibit similarities (Fig. 10d1 and e1), consistent with the finding of a strong correlation between OC and EC in winter (Cao et al., 2007). Since the low temperature and weakened photochemical reactions in winter reduced secondary OC (SOC) generation, and primary OC (POC) and EC mainly originate from direct anthropogenic emissions, such as combustion (Guo, 2016). Compared to the OBS, the mass concentrations in FR (Fig. 10d2-d3) and FOR





554 (Fig. 10e2-e3) are significantly overestimated over a wide range. Similar overestimations have also been reported in the global 555 reanalysis datasets of CAMS and MERRA-2, likely attributed to the hygroscopic growth scheme of carbonaceous aerosols in 556 the models, poorly constrained semi-volatile species escaping from primary organic aerosols (Soni et al., 2021), and aging 557 mechanisms in the models (Huang et al., 2013). After DA, the spatial distribution of the ANA aligns entirely with that of the 558 OBS (Fig. 10d4 and e4), with the improvements in all overestimations (Fig. 10d5 and e5) and the average biases of OC and 559 EC at DA sites both significantly decreasing to 0.14 µg/m<sup>3</sup> (Fig. 11d3 and e3). The VE sites show similar results to the DA 560 sites, with the average biases of less than 2 µg/m3 (Fig. S5d3 and e3). Specifically, for OC (Fig. 12d1 and d2), the CORR 561 values in FOR and ANA are 0.18-0.71 and 0.92-1.00, respectively, with RMSE values of 7.91 µg/m3-26.27 µg/m3 and 0.16 562 μg/m<sup>3</sup>-1.45 μg/m<sup>3</sup>, respectively. For EC (Fig. 12e1 and e2), the CORR values in FOR and ANA are 0.01-0.66 and 0.97-1.00, 563 respectively, with RMSE values of 5.33 µg/m3-16.91 µg/m3 and 0.01 µg/m3-0.26 µg/m3, respectively. Although significant 564 improvements are not observed in FOR at some specific DA sites, the RMSE values at all DA sites decrease by 10%-50% (Fig. 565 12d3 and e3). The CORR values of OC and EC in ANA increase by more than 30%, with most DA sites exceeding 200%, and 566 the RMSE values decrease by more than 90% (Fig. 12d4 and e4). At VE sites (Fig. S6), significant improvements in the CORR 567 are not observed, but the RMSE values in the FOR and ANA decrease, which indicates that DA has limited benefits for whole 568 areas but can effectively reduce biases of whole areas.

#### 569 3.3 Compared to NAQPMS-PDAF v1.0 and global reanalysis dataset

570 To comprehensively evaluate the competitiveness and superiority of NAQPMS-PDAF v2.0 in generating the reanalysis 571 datasets of the PM2.5 chemical compositions, we assimilated the mass concentrations of the five PM2.5 chemical components 572 from all sites (sum of DA sites and VE sites) in February 2022 to generate a reanalysis dataset. We compared our reanalysis 573 dataset with the global reanalysis (RA) datasets (CAMSRA and MERRA-2) and NAQPMS-PDAF v1.0 output. Figure 13 574 illustrates the spatial distribution of the monthly average mass concentrations for the five chemical components. Compared to 575 the OBS (Fig. 13a1 and c1), CAMSRA underestimates the NH4<sup>+</sup> and NO3<sup>-</sup> concentrations and fails to capture the high-value 576 concentration in northern Henan Province (Fig. 13a2 and c2). Meanwhile, MERRA-2 overestimates the concentrations of 577 SO<sub>4</sub><sup>2</sup>, OC, and EC (Fig. 13b2, d2 and e2), particularly SO<sub>4</sub><sup>2</sup>, exhibiting a large region with inaccurately high concentrations. 578 Besides, CAMSRA (approximately 80\*80 km<sup>2</sup>) and MERRA-2 (55\*70 km<sup>2</sup>) have significantly lower spatial resolutions 579 compared to NAQPMS-PDAF v2.0 (5\*5 km<sup>2</sup>). Therefore, NAQPMS-PDAF v2.0 provides a more detailed description of the 580 pollution characteristics of chemical components in Northern China and surrounding areas compared to RA.









Figure 13: Spatial distribution of the monthly averaged concentration of five PM<sub>2.5</sub> chemical components for observations (OBS, a1global reanalysis data (RA, a2-e2), NAQPMS-PDAF v1.0 analysis data (a3-e3) and NAQPMS-PDAF v2.0 analysis data (a4-e4).

584 Although NAQPMS-PDAF v1.0 demonstrates a superior spatial representation of the five chemical components when 585 compared to RA, it fails to capture the high-value concentrations of NH4<sup>+</sup> in the northwest of Henan Province and correct the 586 high-value concentrations of NH4+ in the central and western areas of Hebei Province (Fig. 13a3). Moreover, the scattered 587 high-value concentrations of  $SO_4^{2^2}$  in the North China Plain do not align with the spatial characteristics of the OBS (Fig. 13b3). 588 Notably, NAQPMS-PDAF v1.0 exhibits poor performance in interpreting OC and EC with significant overestimations in a 589 wide range (Fig. 13d3 and e3), which indicates that NAQPMS-PDAF v1.0 is weaker than NAQPMS-PDAF v2.0 in terms of 590 DA performance on chemical components, primarily due to insufficient propagation of observations. In NAQPMS-PDAF v2.0, 591 the LKNETF algorithm with an adaptive forgetting factor is more suitable for the nonlinear and non-Gaussian situations 592 compared to EnKFs in NAQPMS-PDAF v1.0, and the ensemble perturbation with non-Gaussian distribution can better 593 represent the reasonable error distribution of model states.

594

595 Table 3 presents a quantitative comparison of three reanalysis datasets. Compared to the CORR of NAQPMS-PDAF v2.0 596 (0.86-0.99), the CORR of RA for the five chemical components is significantly lower (0.42-0.55). Moreover, NAQPMS-PDAF 597 v1.0 exhibits significantly poorer consistency in SO42-, OC, and EC, with CORR values ranging from 0.35 to 0.57. NAQPMS-598 PDAF v2.0 has lower overall RMSE values (0.14 µg/m<sup>3</sup>-3.18 µg/m<sup>3</sup>) compared to RA and NAQPMS-PDAF v1.0, with RMSE 599 values ranging from 4.51 µg/m<sup>3</sup> to 12.27 µg/m<sup>3</sup> and 2.46 µg/m<sup>3</sup> to 15.50 µg/m<sup>3</sup>, respectively. The characteristics of the R<sup>2</sup> are 600 like those of the CORR and RMSE. For  $NH_4^+$  and  $NO_3^-$ , NAQPMS-PDAF v2.0 (0.85 and 0.93) and v1.0 (0.80 and 0.96) are 601 much higher than RA (0.09 and 0.13). Notably, for SO4<sup>2-</sup>, OC, and EC, NAQPMS-PDAF v2.0 (0.74-0.98) is significantly 602 higher than v1.0 (-0.16-0.25) and RA (-0.15-0.25). Overall, NAQPMS-PDAF v2.0 more accurately and consistently interprets





the five chemical components, particularly for NH4<sup>+</sup>, SO4<sup>2-</sup>, OC, and EC. The reasons are summarized as follows. (1) The DA 603 604 frequency of CAMSRA is 12 hours, which is lower than the hourly DA frequency in NAQPMS-PDAF v2.0. (2) CAMSRA 605 only assimilates satellite retrievals (Inness et al., 2019), and MERRA-2 only assimilates aerosol optical depth (AOD) from 606 both ground-based and space-based remote sensing platforms (Randles et al., 2017). The aerosol optical information analysis 607 increment cannot be allocated to each chemical component accurately and reasonably due to the lack of a deterministic 608 relationship between aerosol optical information and PM2.5 chemical components. (3) NAQPMS-PDAF v1.0 has evident DA 609 shortcomings for chemical components due to the limited DA algorithm under the assumption of linear model or system, 610 inappropriate ensemble perturbation under the assumption of Gaussian distribution, and inadequate observational modules. (4) 611 The state variable structure in NAQPMS-PDAF v1.0 lacks the capacity to effectively mitigate the impact of spurious 612 correlations between chemical component variables, even when using analytical localization.

613 Table 3: Statistical indicators (CORR, RMSE, R<sup>2</sup>) of five PM<sub>2.5</sub> chemical components for global reanalysis data (RA), NAQPMS-

614 PDAF v1.0 analysis data and NAQPMS-PDAF v2.0 analysis data.

Components	CORR		RMSE (µg/m <sup>3</sup> )			$\mathbb{R}^2$			
	RA	v1.0	v2.0	RA	v1.0	v2.0	RA	v1.0	v2.0
$\mathrm{NH_4}^+$	0.49	0.90	0.92	5.59	2.53	2.22	0.09	0.80	0.85
$SO_4^{2-}$	0.55	0.57	0.86	12.27	5.45	2.61	0.25	0.25	0.74
NO <sub>3</sub> -	0.54	0.98	0.96	10.27	2.46	3.18	0.13	0.96	0.93
OC	0.50	0.42	0.97	4.51	12.92	0.93	0.15	-0.09	0.93
EC	0.42	0.35	0.99	7.59	15.50	0.14	-0.15	-0.16	0.98

### 615 3.4 The uncertainty in NAQPMS-PDAF v2.0

616 In ensemble DA, the ensemble members represent possible values of the model states, and the ensemble sampling can 617 determine the uncertainties of the model states. Therefore, the ensemble generation directly affects the propagation of 618 observations and subsequently impacts the final DA performance. Previous studies generated ensemble members based on the 619 uncertainties of emission species and the Gaussian-distribution assumption to satisfy the requirements of EnKFs algorithms 620 (Kong et al., 2021; Wang et al., 2022). However, the true error probability distribution of emission species is not an ideal 621 Gaussian distribution, and the assumption will introduce errors. In this study, we coupled the hybrid nonlinear DA algorithm 622 (LKNETF) with NAQPMS to handle the nonlinear and non-Gaussian situations, which combines the stability of LETKF with 623 the nonlinearity of LNETF. Therefore, we evaluate the performance of ensemble members with different uncertainties and 624 error probability distributions in NAOPMS-PDAF v2.0 through two groups of sensitivity experiments. 625

The first group of experiments (T1-T5) involves controlling the SO<sub>2</sub> uncertainty as a fixed value of 200% and transforming

- 627 the distribution of the perturbation coefficient matrix. The second group of experiments (M1-M5) focuses on assessing the
- 628 influence of SO<sub>2</sub> uncertainty on NH<sub>4</sub><sup>+</sup> and SO<sub>4</sub><sup>2-</sup> DA based on a fixed non-Gaussian distribution (m3=1, m4=6). Figure 14





shows the statistical indicators of the five chemical components under different error probability distributions, including a Gaussian distribution (T1) and four non-Gaussian distributions (T2-T5). The mean CRPS and RMSE in T2 and T4 are lower than those in T1, T3, and T5, and the CORR values in T2 and T4 are higher than those in T1, T3, and T5, indicating that the DA performance of non-Gaussian-distribution assumption is superior to that of Gaussian-distribution. Except for SO<sub>4</sub><sup>2-</sup>, the positively skewed non-Gaussian distribution performs better than negatively skewed distribution. Except for SO<sub>4</sub><sup>2-</sup>, the performance in T2 outweighs that in T4 for other chemical components, implying that higher kurtosis harms the chemical components DA.



<sup>636</sup> 

639 SO<sub>2</sub> is a crucial precursor of NH<sub>4</sub><sup>+</sup> and SO<sub>4</sub><sup>2-</sup>, and perturbing SO<sub>2</sub> affects the forecast and simulation of NH<sub>4</sub><sup>+</sup> and SO<sub>4</sub><sup>2-</sup>. Table 640 4 presents statistical indicators of  $NH_4^+$  and  $SO_4^{2-}$  analysis fields based on ensemble perturbations with different  $SO_2$ 641 uncertainties (12%-300%). Increasing the uncertainty of SO<sub>2</sub> from 12% to 200% leads to a decrease in the mean CRPS in the 642 SO42- analysis field from 2.67 to 1.40, an increase in the CORR from 0.51 to 0.74, and a reduction in the RMSE from 4.10 643 µg/m3 to 2.37 µg/m3. Similarly, the mean CRPS in the NH4+ analysis field decreases from 0.98 to 0.77, the CORR increases 644 from 0.88 to 0.91, and the RMSE decreases from  $1.55 \,\mu g/m^3$  to  $1.33 \,\mu g/m^3$ . It indicates that increasing the uncertainty of SO<sub>2</sub> 645 improves the DA performance on NH4+ and SO42- because the higher SO2 uncertainty makes SO2 perturbed sufficiently, and 646 the estimated error probability distribution is closer to the real distribution, resulting in a sufficient spread of observations. 647 However, when the uncertainty of SO<sub>2</sub> reaches 300%, the statistical indicators do not significantly improve and even worsen 648 because excessively high SO2 uncertainty causes the estimated error probability distribution to deviate from the true 649 distribution. Thus, selecting appropriate uncertainties for emission species is crucial in aerosol chemical component DA.

Figure 14: Statistical indicators (mean CRPS (a), RMSE (b), and CORR (c)) of five PM<sub>2.5</sub> chemical components for five perturb experiments based on distribution.





650

- 651 To summarize, the non-Gaussian-distribution assumption outperforms the Gaussian-distribution assumption in NAQPMS-
- 652 PDAF v2.0. Positive skewness performs better than negative skewness, and excessively high kurtosis should be avoided.
- 653 Additionally, appropriately increasing the uncertainty of SO<sub>2</sub> enhances the DA performance of NH<sub>4</sub><sup>+</sup> and SO<sub>4</sub><sup>2-</sup>. Future studies
- 654 should conduct more sensitivity experiments on emission species perturbation to determine the suitable schemes for different
- 655 aerosol chemical components.

### Table 4: Statistical indicators (mean CRPS (a), RMSE (b), and CORR (c)) of five PM<sub>2.5</sub> chemical components for five perturb experiments based on SO<sub>2</sub> emission uncertainty.

Experiment	SO4 <sup>2-</sup>			$\mathrm{NH_4^+}$			
	CRPS	CORR	RMSE	CRPS	CORR	RMSE	
M1	2.67	0.51	4.10	0.98	0.88	1.55	
M2	2.07	0.59	3.24	0.92	0.89	1.48	
M3	1.61	0.69	2.63	0.83	0.91	1.39	
M4	1.40	0.74	2.37	0.77	0.91	1.33	
M5	1.41	0.74	2.39	0.78	0.91	1.33	

## 658 4 Conclusions

In this paper, we online coupled NAQPMS with PDAF-OMI to develop a novel hybrid nonlinear DA system (NAQPMS-PDAF v2.0) with level-2 parallelization based on a hybrid Kalman-Nonlinear Ensemble Transform Filter (LKNETF) for the first time. Compared to NAQPMS-PDAF v1.0, NAQPMS-PDAF v2.0 with OMI can be applied with multiple component types and nonlinear/non-Gaussian situations in chemical analysis to effectively interpret five PM<sub>2.5</sub> chemical components (NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, OC and EC), which is not achieved in previous studies. The background error covariance was calculated by ensemble perturbation based on adaptive uncertainties and non-Gaussian-distribution assumption of emission species. The DA experiments were conducted based on 33 observational sites in Northern China and surrounding areas.

667 NAQPMS-PDAF v2.0 with LKNETF can maintain high accuracy and reliability in ensemble DA with an ensemble size of 10, 668 smaller than the traditional minimum of 20 ensemble members, as observed in prior ensemble assimilation studies. The FR 669 (free-run fields without DA) have a poor consistency with the observations, with the CORR values ranging from 0.32-0.56 670 and the R<sup>2</sup> values less than 0.3, showing that SO4<sup>2-</sup>, OC and EC are significantly overestimated, while NH4<sup>+</sup> and NO3<sup>-</sup> are 671 underestimated. A significant improvement was observed in the ANA (analysis fields) of the DA sites. The CORR values are 672 not less than 0.86, the RMSE and MAE values do not exceed 3.23 µg/m<sup>3</sup> and 1.49 µg/m<sup>3</sup>, respectively, and R<sup>2</sup> is not less than 673 0.74. Specifically, the CORR values for NO<sub>3</sub>, OC, and EC are not less than 0.96, and  $R^2$  is not less than 0.93. The error 674 distributions of the five chemical components concentrate to 0 with the mean bias ranging from  $0\pm0.08 \ \mu g/m^3$  to  $1.02\pm3.07$ 675 µg/m3. These improvements are also found in the ANA at VE sites, indicating an excellent DA performance of NAQPMS-





676 PDAF v2.0.

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678 The ability of NAQPMS-PDAF v2.0 to interpret the spatiotemporal characteristics of the five chemical components was 679 examined. For temporal variations, compared to the FR and FOR (forecast fields), the ANA closely aligned with the OBS 680 (observations) and accurately captured the peak concentrations of SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, and NH<sub>4</sub><sup>+</sup> on specific periods (such as February 681 25<sup>th</sup>), indicating good consistency and accurate characterization. Specifically, the CORR of the ANA at the six representative 682 sites increased by 13.64%-89.58% and 17.19%-75.00%, respectively, while the RMSE decreased by 56.03%-83.13% and 683 40.74%-72.20%. For spatial distributions, after DA, both NH4+ and NO3 with positive analysis increments exhibit significant 684 improvements in CORR and RMSE, as most DA sites show improvements of over 150% in CORR and over 50% in RMSE. 685 SO42, OC, and EC with negative analysis increments were also improved. Especially for OC and EC, the improvements of 686 CORR and RMSE at most DA sites were over 200% and over 90%, respectively. The improvements at VE sites were also 687 identified. Consequently, DA successfully aligned the spatiotemporal characteristics of the ANA with OBS and significantly 688 reduced the biases of five chemical components.

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 $690 \qquad \text{Compared to the global reanalysis datasets} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.42\text{-}0.55, \text{RMSE: } 4.51\text{-}12.27 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.25 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.25 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.25 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.25 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.55 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.55 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{CORR: } 0.51\text{-}12.55 \ \mu\text{g/m}^3) \text{ and } \text{NAQPMS-PDAF v1.0} (\text{NAQPMS-PDAF v1.0} (\text{NAQPMS-PDAF v1.0} (\text{NAQPMS-PDAF v1.0} (\text{N$ 

691 0.35-0.98, RMSE: 2.46-15.50  $\mu$ g/m<sup>3</sup>), the NAQPMS-PDAF v2.0 (CORR: 0.86-0.99, RMSE: 0.14-3.18  $\mu$ g/m<sup>3</sup>) has significant 692 superiority in generating the reanalysis datasets of the PM<sub>2.5</sub> chemical compositions with high spatiotemporal resolution. 693 Besides, NAQPMS-PDAF v1.0 cannot capture the high-value concentrations and exhibits poor performance when interpreting 694 SO<sub>4</sub><sup>2-</sup>, OC, and EC with CORR values ranging from 0.35 to 0.57. In contrast, NAQPMS-PDAF v2.0 interprets the five chemical 695 components more accurately and consistently.

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Finally, the uncertainties of NAQPMS-PDAF v2.0 are examined by identifying the influence of ensemble generation on ensemble DA performance. The non-Gaussian-distribution assumption outperforms the Gaussian-distribution assumption in NAQPMS-PDAF v2.0. Positive skewness performs better than negative skewness, and excessively high kurtosis should be avoided. Additionally, appropriately increasing the uncertainty of SO<sub>2</sub> enhances the DA performance of NH<sub>4</sub><sup>+</sup> and SO<sub>4</sub><sup>2-</sup>. Future studies should conduct more sensitivity experiments on emission species perturbation to determine the suitable schemes for different aerosol chemical components.

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The novel hybrid nonlinear DA system (NAQPMS-PDAF v2.0) can be effectively applied in the interpretation of chemical components and outperform in generating the reanalysis dataset of the five  $PM_{2.5}$  chemical components with high accuracy and high consistency, thus providing the sufficient channel to investigate the spatiotemporal characteristics, identify the regional transport and prevent and control aerosol composition pollution. In future work, we plan to research the vertical DA





- 708 of chemical components, introduce more vertical information from more observational platforms, and verify the simultaneous
- 709 DA performance of surface and vertical mass concentrations.
- 710

# 711 Code and data availability

712 The source codes in our work are available online via Zenodo (https://doi.org/10.5281/zenodo.10886914).

#### 713 Author contributions

- 714 HL developed the data assimilation system, performed numerical experiments, carried out the analysis and wrote the original
- 715 manuscript. TY provided scientific guidance, designed the paper struttre and wrote this paper. LN developed PDAF and
- 716 provided help for the model code. DWZ, DZ, and GT provided PM<sub>2.5</sub> chemical component data. HW provided help for the
- 717 model code. YS, PF, HS, ZW did overall supervision. All authors reviewed and revised this paper.

# 718 Competing interests

719 The contact author has declared that neither they nor their co-authors have any competing interests.

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