# 1 A modeling study of the nonlinear response of fine

# 2 particles to air pollutant emissions in the Beijing-Tianjin-

# 3 Hebei region

4

- 5 Bin Zhao<sup>1,2,3</sup>, Wenjing Wu<sup>1,2</sup>, Shuxiao Wang<sup>1,2</sup>, Jia Xing<sup>1,2</sup>, Xing Chang<sup>1,2</sup>, Kuo-
- 6 Nan Liou<sup>3</sup>, Jonathan H. Jiang<sup>4</sup>, Yu Gu<sup>3</sup>, Carey Jang<sup>5</sup>, Joshua S. Fu<sup>6</sup>, Yun Zhu<sup>7</sup>,
- 7 Jiandong Wang<sup>1,2</sup>, Jiming Hao<sup>1,2</sup>
- 8 [1] School of Environment, and State Key Joint Laboratory of Environment Simulation and
- 9 Pollution Control, Tsinghua University, Beijing 100084, China
- 10 [2] State Environmental Protection Key Laboratory of Sources and Control of Air Pollution
- 11 Complex, Beijing 100084, China
- 12 [3] Joint Institute for Regional Earth System Science and Engineering and Department of
- 13 Atmospheric and Oceanic Sciences, University of California, Los Angeles, CA 90095, USA
- 14 [4] Jet propulsion Laboratory, California Institute of Technology, Pasadena, CA 91109, USA
- 15 [5] U.S. Environmental Protection Agency, Research Triangle Park, NC 27711, USA
- 16 [6] Department of Civil and Environmental Engineering, University of Tennessee, Knoxville,
- 17 TN 37996, United States
- 18 [7] School of Environmental Science and Engineering, South China University of
- 19 Technology, Guangzhou 510006, China

20

21

22 Correspondence to: Shuxiao Wang (shxwang@tsinghua.edu.cn)

23

#### 24 Abstract.

- 25 The Beijing-Tianjin-Hebei (BTH) region has been suffering from the most severe fine particle
- 26 (PM<sub>2.5</sub>) pollution in China, which causes serious health damage and economic loss.
- 27 Quantifying the source contributions to PM<sub>2.5</sub> concentrations has been a challenging task
- 28 because of the complicated non-linear relationships between PM<sub>2.5</sub> concentrations and
- 29 emissions of multiple pollutants from multiple spatial regions and economic sectors. In this
- 30 study, we use the Extended Response Surface Modeling (ERSM) technique to investigate the

nonlinear response of PM<sub>2.5</sub> concentrations to emissions of multiple pollutants from different regions and sectors over the BTH region, based on over 1000 simulations by a chemical transport model (CTM). The ERSM-predicted PM<sub>2.5</sub> concentrations agree well with independent CTM simulations, with correlation coefficients larger than 0.99 and mean normalized errors less than 1%. Using the ERSM technique, we find that, among all air pollutants, primary inorganic PM<sub>2.5</sub> makes the largest contribution (24-36%) to PM<sub>2.5</sub> concentrations. The contribution of primary inorganic PM<sub>2.5</sub> emissions is especially high in heavily polluted winter, and is dominated by the industry as well as residential and commercial sectors, which should be prioritized in PM<sub>2.5</sub> control strategies. The total contributions of all precursors (nitrogen oxides, NO<sub>X</sub>; sulfur dioxides, SO<sub>2</sub>; ammonia, NH<sub>3</sub>; non-methane volatile organic compounds, NMVOC; intermediate-volatility organic compounds, IVOC; primary organic aerosol, POA) to PM<sub>2.5</sub> concentrations range between 31% and 48%. Among these precursors, PM<sub>2.5</sub> concentrations are primarily sensitive to the emissions of NH<sub>3</sub>, NMVOC+IVOC, and POA. The sensitivities increase substantially for NH<sub>3</sub> and NO<sub>X</sub>, and decrease slightly for POA and NMVOC+IVOC with the increase in the emission reduction ratio, which illustrates the nonlinear relationships between precursor emissions and PM<sub>2.5</sub> concentrations. The contributions of primary inorganic PM<sub>2.5</sub> emissions to PM<sub>2.5</sub> concentrations are dominated by local emission sources, which account for over 75% of the total primary inorganic PM<sub>2.5</sub> contributions. For precursors, however, emissions from other regions could play similar roles as local emission sources in the summer and over the northern part of BTH. The source contribution features for various types of heavy-pollution episodes are distinctly different from each other, and from the monthly mean results, illustrating that control strategies should be differentiated based on the major contributing sources during different types of episodes.

2425

26

27

28

29

30

31

32

1

2

3

5

6

7

8

9

10

11

12

13

14

15

16

17

18 19

20

21

22

23

#### 1 Introduction

China is one of the regions with highest concentration of PM<sub>2.5</sub> (particulate matter with aerodynamic diameter equal to or less than 2.5 μm) in the world (van Donkelaar et al., 2015). The problem is especially serious over the Beijing-Tianjin-Hebei (BTH) region, one of the most populous and developed regions in China. Annual average PM<sub>2.5</sub> concentrations in this region reached 85-110 μg/m<sup>3</sup> during 2013-2015, which approximately triple the standard threshold (35 μg/m<sup>3</sup>) and far exceed those in other metropolitan regions (Wang et al., 2017). It

has been estimated that the severe PM<sub>2.5</sub> pollution leads to about 1.05-1.23 million premature

deaths per year in China (Lim et al., 2012; Burnett et al., 2014; Wang et al., 2016b), and the

3 monetized loss over the BTH region is as high as 134 billion Chinese Yuan, representing 2.2%

4 of regional gross domestic product (GDP) (Lv and Li, 2016). Additionally, PM<sub>2.5</sub> substantially

5 affects global and regional climate by absorbing and scattering solar radiation and by altering

cloud properties (IPCC, 2013; Seinfeld et al., 2016; Zhao et al., 2017a), which in turn exert

7 impact on regional air quality (Wang et al., 2014a; Zhao et al., 2017b).

6

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

2728

29

30

31

32

To tackle the heavy PM<sub>2.5</sub> pollution problem, Chinese government issued the "Action Plan on Prevention and Control of Air Pollution" in September 2013, which aimed at a 25% reduction in PM<sub>2.5</sub> concentrations over the BTH region by 2017 from the 2012 levels (The State Council of the People's Republic of China, 2013). The attainment of ambient PM<sub>2.5</sub> standard would further require substantial reductions in air pollutant emissions (Wang et al., 2017; Wang et al., 2015). To establish emission control strategies, many studies have apportioned the sources of PM<sub>2.5</sub> over the BTH region, either by mining monitoring data using the Positive Matrix Factorization and Chemical Mass Balance methods (e.g., Zhang et al., 2007; Yu et al., 2013) or by embedding chemical tracers in chemical transport models (CTMs) (e.g., Wang et al., 2016c; Li et al., 2015b; Ying et al., 2014). While these studies can capture the current contributions of various sources to PM<sub>2.5</sub> concentrations, these contributions could differ significantly from the PM<sub>2.5</sub> reductions induced by reducing emissions from the corresponding sources, due to highly nonlinear chemical mechanisms (Han et al., 2016; Wang et al., 2011). Therefore, it is imperative to assess the nonlinear response of PM<sub>2.5</sub> to pollutant emissions from multiple sources, which could provide direct support for the development of effective control policies.

The most widely used technique to evaluate the responses of PM<sub>2.5</sub> concentrations to emission changes is the "Brute force" method, which involves perturbing emissions from a certain source and repeated solution of a CTM (Russell et al., 1995). A number of studies have utilized the "Brute force" method to quantify the sensitivities of PM<sub>2.5</sub> concentrations over the BTH region to emissions from different spatial regions (Streets et al., 2007; Wang et al., 2008; Li and Han, 2016; Wang et al., 2014b) or different economic sectors (Wang et al., 2008; Han et al., 2016; Wang et al., 2014b; Liu et al., 2016), either on a seasonal basis (Streets et al., 2007; Wang et al., 2008; Han et al., 2016; Liu et al., 2016) or during a specific heavy-pollution episode (Li and Han, 2016; Wang et al., 2014b). To improve the

1 computational efficiency, several mathematic techniques embedded in CTMs have been 2 developed to simultaneously calculate the sensitivities of the modeled concentrations to 3 multiple emission sources, including the Decoupled Direct Method (Yang et al., 1997) and 4 Adjoint Analysis (Sandu et al., 2005; Hakami et al., 2006). Zhang et al. (2016) used the 5 Adjoint Analysis method to examine sensitivities of PM<sub>2.5</sub> concentrations in the BTH region 6 to pollutant emissions during several pollution periods. However, all the preceding studies 7 only quantified first-order sensitivities and therefore could not capture the nonlinearity in the 8 responses of PM<sub>2.5</sub> concentrations to pollutant emissions, which can be extremely strong in 9 metropolitan regions like BTH due to complex chemical mechanisms (Wang et al., 2011). 10 Moreover, no studies have simultaneously evaluated the response of PM<sub>2.5</sub> concentrations in 11 BTH to emissions of multiple pollutants from different sectors and regions, which we need to 12 consider and balance to develop cost-effective control strategies.

13

14

15

16

17

18

1920

21

22

23

24

25

26

27

28

29

30

31

32

In light of the drawbacks of the preceding methods, the Response Surface Modeling (RSM) technique (denoted by "conventional RSM" technique hereafter to distinguish from the ERSM technique) has been developed by using advanced statistical techniques to characterize the complex nonlinear relationship between model outputs and inputs (U.S. Environmental Protection Agency, 2006; Xing et al., 2011; Wang et al., 2011). This technique has been applied to the United States (U.S. Environmental Protection Agency, 2006) and the Eastern China (Wang et al., 2011) to evaluate the response of PM<sub>2.5</sub> and its chemical components to pollutant emissions. However, the number of emission scenarios required to build conventional RSM depends on the variable number via an equation of fourth or higher order (Zhao et al., 2015b). Therefore, the required scenario number would be tens of thousands for over 15 variables and even hundreds of thousands for over 25 variables, which is computationally impossible for most three-dimensional CTMs. To overcome this major limitation, we recently developed the Extended Response Surface Modeling (ERSM) technique (Zhao et al., 2015b), which substantially reduced the scenario number needed to build the response surface and hence extended its applicability to an increased number of regions, pollutants, and sectors with an acceptable computational burden.

Given the advantage of the ERSM technique, here we apply it to over 1000 simulations by the Community Multi-scale Air Quality model with Two-Dimensional Volatility Basis Set (CMAQ/2D-VBS) to systematically evaluate the nonlinear response of PM<sub>2.5</sub> to emission changes of multiple pollutants from different sectors and regions over the BTH region. The

- 1 major sources contributing to PM<sub>2.5</sub> and its major components are identified and the
- 2 nonlinearity in the response of PM<sub>2.5</sub> to emission changes is characterized. Based on results of
- 3 this study, suggestions for PM<sub>2.5</sub> control policies over the BTH region are proposed.

#### 4 2 Methodology

### 5 2.1 CMAQ/2D-VBS configuration and evaluation

- 6 The CMAQ/2D-VBS model was developed in our previous study (Zhao et al., 2016) by
- 7 incorporating the 2D-VBS model framework into CMAQv5.0.1. Compared with the default
- 8 CMAQ, the CMAQ/2D-VBS model explicitly simulates aging of secondary organic aerosol
- 9 (SOA) formed from non-methane volatile organic compounds (NMVOC), aging of primary
- 10 organic aerosol (POA), and photo-oxidation of intermediate-volatility organic compounds
- 11 (IVOC), thereby significantly improving the simulation results of organic aerosol (OA),
- 12 particularly SOA. The model parameters within the 2D-VBS framework have been optimized
- in our previous studies (Zhao et al., 2015a; Zhao et al., 2016) based on a series of smog-
- 14 chamber experiments. Here we use the same model parameters as those of the "High-Yield
- VBS" configuration reported in Zhao et al. (2016), which agrees best with surface OA and
- 16 SOA observations among three model configurations. An application in the Eastern China
- 17 reveals that CMAO/2D-VBS reduces the underestimation in OA concentrations from 45%
- 18 (default CMAQv5.0.1) to 19%. More importantly, while the default CMAQv5.0.1
- 19 substantially underestimates the fraction of SOA in OA by 5-10 times and can not track
- 20 oxygen-to-carbon ratio (O:C), the SOA fraction and O:C simulated by CMAQ/2D-VBS agree
- 21 fairly well with observations.
- We apply the CMAQ/2D-VBS model over the BTH region. One-way, double nesting
- 23 simulation domains are used, as shown in Fig. 1. Domain 1 covers East Asia with a grid
- 24 resolution of 36 km×36 km; domain 2 covers the BTH and its surrounding regions with a grid
- 25 resolution of 12 km×12 km. We use the SAPRC99 gas-phase chemistry module and the
- 26 AERO6 aerosol module, in which the treatment of OA is replaced with the 2D-VBS
- 27 framework. The aerosol thermaldynamics is based on ISORROPIA-II. The initial and
- boundary conditions for Domain 1 are kept constant as the model default profile, and those
- 29 for Domain 2 are extracted from the output of Domain 1. A 5-day spin-up period is used to
- reduce the influence of initial conditions on modeling results.
- The Weather Research and Forecasting Model (WRF, version 3.7) is used to generate the
- 32 meteorological fields. The National Center for Environmental Prediction (NCEP)'s FNL

(Final) Operational Global Analysis data (ds083.2) at 1.0° × 1.0° and 6-h resolution are used 1 2 to generate the first guess field. The NCEP's Automated Data Processing (ADP) data (ds351.0 and ds461.0) are used in objective analysis (i.e., grid nudging). The major physics 3 4 options for WRF include the Kain-Fritsch cumulus scheme, the Pleim-Xiu land-surface 5 module, the Asymmetric Convective Model with non-local upward mixing and local 6 downward mixing (ACM2) for planetary boundary layer (PBL) parameterization, the 7 Morrison double-moment scheme for cloud microphysics, and the Rapid Radiative Transfer 8 Model for GCMs (RRTMG) radiation scheme. The land cover type data are obtained from the 9 Moderate resolution Imaging Spectroradiometer (MODIS). The simulation periods are 10 January, March, July, and October in 2014, representing winter, spring, summer, and fall. We select these four months because the occurrence frequencies of various meteorological types 11 12 in these months are statistically most similar to the average conditions in winter, spring, 13 summer, and fall during 2004-2013 (Wu, 2016).

14

15

16

17

18

19

2021

22

23

24

25

26

27

28

29

30

31

32

A high-resolution anthropogenic emission inventory in 2014 has been developed using an "emission factor method" (Fu et al., 2013; Zhao et al., 2013b) for the BTH region by Tsinghua University. The emissions from area and mobile sources are first calculated for each prefecture-level city based on statistical data, and subsequently distributed into the model grids according to spatial distribution of population, GDP, and road networks. A unit-based method (Zhao et al., 2008) is applied to estimate and locate the emissions from large point sources (LPS) including power plants, iron and steel plants, and cement plants. The anthropogenic emission inventory in other provinces of China was originally developed for 2010 and 2012 in our previous studies (Zhao et al., 2013b; Zhao et al., 2013a; Wang et al., 2014c; Cai et al., 2016), which has been updated to 2014 in this study following the same methodology. In both the BTH and national emission inventories, the emissions from open burning of agricultural residue are calculated using crop yields, straw to grain ratio, fraction of biomass burned in the open field, and emission factors (Fu et al., 2013; Zhao et al., 2013b; Wang and Zhang, 2008). We do not include the emissions from forest and grassland fires, which typically account for less than 5% of the total biomass burning emissions over the BTH region (Qin and Xie, 2011) and are not the focus of the present study. Table S1 summarizes emissions of major air pollutants in each prefecture-level city over the BTH region in 2014; Table S2 gives the provincial emissions in the whole China in 2014. The emissions for other countries are obtained from the MIX emission inventory (Li et al., 2015a) for 2010, which is

- 1 the latest year available. Following our previous study (Zhao et al., 2016), we assume IVOC
- 2 emissions to be 30 times, 4.5 times, 1.5 times, and 3.0 times the POA emissions from gasoline
- 3 vehicles, diesel vehicles, biomass burning, and other emission sources, respectively, which is
- 4 based on a series of laboratory measurements (Gordon et al., 2014b; Gordon et al., 2014a;
- 5 Hennigan et al., 2011; Jathar et al., 2014). The biogenic emissions were calculated by the
- 6 Model of Emissions of Gases and Aerosols from Nature (MEGAN; Guenther et al., 2006).
- 7 We compared the simulation results of WRFv3.7 and CMAQ/2D-VBS with
- 8 meteorological observations obtained from the National Climatic Data Center (NCDC), PM<sub>2.5</sub>
- 9 observations at 138 state-controlled observational sites, and observations of major PM<sub>2.5</sub>
- 10 chemical components at 7 sites within the modeling domain. We show that the meteorological
- and chemical simulations generally agree well with observations, with performance statistics
- mostly within the benchmark values proposed by previous studies. Details of the model
- evaluation methods and results are given in the Supplementary Information (Section 1, Table
- 14 S3-S5, Fig. S1-S5).

#### 2.2 Development of ERSM prediction system

- 16 The detailed methodologies of the conventional RSM and ERSM techniques have been
- described in our previous papers (Zhao et al., 2015b; Xing et al., 2011). Here we only
- 18 summarize some key components. The conventional RSM technique characterizes the
- 19 relationships between a response variable (e.g., PM<sub>2.5</sub> concentration) and a set of control
- variables (i.e., emissions of particular pollutants from particular sources) based on a number
- of randomly generated emission control scenarios (Xing et al., 2011; Wang et al., 2011). The
- 22 PM<sub>2.5</sub> concentration for each emission scenario is calculated with a CTM (CMAQ/2D-VBS in
- 23 this study), and the conventional RSM is subsequently established using the Maximum
- 24 Likelihood Estimation Empirical Best Linear Unbiased Predictors (MLE-EBLUPs)
- developed by Santner et al. (2003). Due to the limitation of the conventional RSM technique
- with respect to variable number, we have developed the ERSM technique (Zhao et al., 2015b)
- 27 to extend the applicability to an increased number of variables and geographical regions. The
- 28 ERSM technique first quantifies the relationship between PM<sub>2.5</sub> concentrations and precursor
- 29 emissions for each single region using the conventional RSM technique as described above,
- and then assesses the effects of inter-regional transport of PM<sub>2.5</sub> and its precursors on PM<sub>2.5</sub>
- 31 concentration in the target region. In order to quantify the interaction among regions, we
- 32 introduce a key assumption that the emissions of precursors in the source region affect PM<sub>2.5</sub>

concentrations in the target region through two major processes: (1) the inter-regional transport of precursors enhancing the chemical formation of secondary PM<sub>2.5</sub> in the target region; (2) the formation of secondary PM<sub>2.5</sub> in the source region followed by transport to the target region. We quantify the individual contributions of these two processes as well as the contribution of local emissions in the target region, which are subsequently integrated to derive the total PM<sub>2.5</sub> concentrations in the target region. The development of the ERSM prediction system requires several hundred to over 1000 emission scenarios, but once built, it enables real-time prediction of PM<sub>2.5</sub> concentrations for any given control strategy and proves to be an efficient and user-friendly decision making tool. Moreover, ERSM can be applied to design least-cost control strategy once it is coupled with control cost models/functions that links the emission reductions with economic costs.

1

2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

2122

23

24

25

26

27

28

29

30

31

32

For application of the RSM/ERSM techniques to the BTH region, we define 5 target regions in the inner modeling domain (Domain 2), i.e., Beijing, Tianjin, Northern Hebei (N Hebei), Eastern Hebei (E Hebei), and Southern Hebei (S Hebei), as shown in Fig. 1. The decomposition of the Hebei province is based on a preliminary analysis of the pollutant transport patterns over the BTH region (Section 2 in the Supplementary Information). The simulation using back trajectory method indicates that four major types of heavy-pollution episodes in Beijing are primarily contributed by air mass from the south, the local area, the northwest, and the southeast. We develop two RSM/ERSM prediction systems (Table 1). The response variables for the first prediction system, which is built using the conventional RSM technique, are concentrations of PM<sub>2.5</sub>, SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, and OA over the urban areas of prefecture-level cities in the five target regions. For the second prediction system that is established using the ERSM technique, the response variables are only PM<sub>2.5</sub> concentrations. The first prediction system use 101 emission control scenarios generated by the Latin Hypercube Sample (LHS) method (Iman et al., 1980) to map atmospheric concentrations versus emissions of five PM<sub>2.5</sub> precursors, i.e., NO<sub>X</sub>, SO<sub>2</sub>, NH<sub>3</sub>, NMVOC+IVOC, and POA, in all five target regions (Table 1). It is on one hand intended for the validation of the second system (Section 3.1), and on the other hand used to study the source contributions of major PM<sub>2.5</sub> components. For the second system, the emissions of the preceding PM<sub>2.5</sub> precursors as well as primary inorganic PM<sub>2.5</sub> (i.e., the chemical components of primary PM<sub>2.5</sub> other than POA) in each of the 5 regions are categorized into 7 and 4 control variables, respectively, resulting in 55 control variables in total (Table 1). Note that we distinguish POA and primary

1 inorganic PM<sub>2.5</sub> because the former undergoes chemical reactions and produces SOA, while 2 the latter is mostly chemically inert in the CMAQ/2D-VBS model. We generate 1121 3 scenarios (see Table 1) to build the response surface, following the method detailed in Zhao et 4 al. (2015b). Specifically, the scenarios include (1) 1 CMAQ/2D-VBS base case; (2) 200 5 scenarios generated by applying LHS method for the control variables of precursors in 6 Beijing, 200×4 scenarios generated in the same way for Tianjin, Northern Hebei, Eastern 7 Hebei, and Southern Hebei; (3) 100 scenarios generated by applying LHS method for the total 8 emissions of NO<sub>X</sub>, SO<sub>2</sub>, NH<sub>3</sub>, NMVOC+IVOC, and POA in all 5 regions; and (4) 20 9 scenarios where one of the control variables of primary inorganic PM<sub>2.5</sub> emissions is set to 0.25 for each scenario. Here the scenario numbers (200 in group 2 and 100 in group 3) are 10 11 determined based on numerical experiments conducted in our previous studies (Xing et al., 12 2011; Wang et al., 2011), which showed that the response surface for 7 and 5 variables could 13 be built with good prediction performance (mean normalized error < 1%; correlation 14 coefficient > 0.99) using 200 and 100 scenarios, respectively. Finally, we generate 54 15 independent scenarios for out-of-sample validation, which will be detailed in Section. 3.1.

For application of the ERSM prediction system to quantitatively characterize the sensitivity of PM<sub>2.5</sub> concentrations to emission changes, we define "PM<sub>2.5</sub> sensitivity" as the change ratio of PM<sub>2.5</sub> concentration divided by the reduction ratio of a emission source, following previous studies (Zhao et al., 2015b; Wang et al., 2011).

$$S_a^X = \left[ \left( C^* - C_a \right) / C^* \right] / (1-a) \tag{4}$$

where  $S_a^X$  is the PM<sub>2.5</sub> sensitivity to emission source X at its emission ratio a;  $C^*$  and  $C_a$  are  $PM_{2.5}$  concentrations in the base case (when the emission ratio of X is 1) and in the control scenario where the emission ratio of X is a, respectively. Similar indices can be defined for chemical components of  $PM_{2.5}$ , such as  $NO_3^-$ ,  $SO_4^{2-}$ , and OA.

24 25

26

27

16

17 18

19

21 22

23

#### Results and discussion 3

#### Validation of ERSM performance 3.1

28 The conventional RSM technique has been extensively demonstrated to have high accuracy 29 and stability in previous papers (Xing et al., 2011; Wang et al., 2011), so we only describe the 30

validation of the ERSM technique. Following Zhao et al. (2015b), we assess the performance

of the ERSM prediction system using the "out-of-sample" and 2D-isopleths validation methods, which focus on the accuracy and stability of the prediction system, respectively.

For out-of-sample validation, we use the ERSM prediction system to calculate the PM<sub>2.5</sub> concentrations for 54 "out-of-sample" control scenarios, i.e., scenarios independent from those used to build the prediciton system, and compare with the corresponding CMAQ/2D-VBS simulation results. These 54 out-of-sample scenarios (summarized in Table S6) include 40 cases (case 1-40) where the control variables of precursors change but those of primary inorganic PM<sub>2.5</sub> stay the same as the base case, 4 cases (case 41-44) the other way around, and 10 cases (case 45-54) where control variables of precursors and primary inorganic PM<sub>2.5</sub> change simultaneously. Most cases are generated randomly with the LHS method (case 4-6, 10-12, 16-18, 22-24, 28-54), and some cases are designed where all control variables are subject to large emission changes (case 1-3, 7-9, 13-15, 19-21, 25-27).

Figure 2 compares the ERSM-predicted and CMAQ/2D-VBS-simulated PM<sub>2.5</sub> concentrations and PM<sub>2.5</sub> responses (defined as the difference between PM<sub>2.5</sub> concentration in an emission control scenario and that in the base case) for the out-of-sample scenarios using scatter plots. Table 2 summarizes the statistics of the model performance. The definitions of normalized error (NE), mean normalized error (MNE), and normalized mean error (NME) are given as follows:

$$NE = |P_i - S_i| / S_i \tag{1}$$

20 MNE=
$$\frac{1}{N_S} \sum_{i=1}^{N_S} [|P_i - S_i| / S_i]$$
 (2)

21 
$$NME = \sum_{i=1}^{N_S} |P_i - S_i| / \sum_{i=1}^{N_S} S_i$$
 (3)

where  $P_i$  and  $S_i$  are the ERSM-predicted and CMAQ/2D-VBS-simulated value of the i<sup>th</sup> out-of-sample scenario; Ns is the number of out-of-sample scenarios. Figure 2 shows that the ERSM predictions and CMAQ/2D-VBS simulations agree well with each other. For PM<sub>2.5</sub> concentrations, the correlation coefficients are larger than 0.99, and the MNEs and NMEs are less than 1% for all four months. The maximum NEs could be as large as 11% for particular month and region, but the 95% percentiles of NEs are all within 4.4%. NEs exceeding 4.4% happen only for the scenarios where most control variables are reduced substantially, indicating relatively large errors at low emission rates, which is consistent with our previous study (Zhao et al., 2015b). Note that all sensitivity scenarios used in Sections 3.2-3.4 have  $\leq$  80% emission reductions, which helps to avoid relatively large errors. We also examine the errors in predicted PM<sub>2.5</sub> response. Since the CMAQ/2D-VBS-simulated PM<sub>2.5</sub> responses are

very close to zero in several scenarios, their normalized errors (NEs) and mean normalized 2 errors (MNEs) could be extremely large even if the absolute errors are small, which cannot 3 properly characterize the accuracy of the ERSM technique. For this reason, we only calculate 4 the correlation coefficients and NMEs (Table 2). The correlation coefficients of PM<sub>2.5</sub> 5 response are larger than 0.99, and the NMEs are within 5.6% for all months. In summary, the 6 out-of-sample validation indicates an overall good agreement between ERSM predictions and 7 CMAQ/2D-VBS simulations.

1

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

We further examine whether the ERSM technique can capture the trends in PM<sub>2.5</sub> concentrations in response to continuous changes in precursor emissions, i.e., the stability of the ERSM technique. To this end, we compare the 2D-isopleths of PM<sub>2.5</sub> concentrations as a function of simultaneous changes in two precursors' emissions in all five regions derived from the ERSM and conventional RSM techniques. It should be noted that, although the ERSM technique is applicable to a much larger number of control variables than conventional RSM, the assumptions in the treatment of inter-regional transport (Section 2.2) in ERSM might affect its accuracy. Nevertheless, the predictions by conventional RSM can be regarded as proxies for real CMAQ/2D-VBS simulations since it has been extensively demonstrated to have high accuracy and stability in previous studies (Xing et al., 2011; Wang et al., 2011). For this reason, the comparison between the ERSM and conventional RSM techniques helps to evaluate the stability of the ERSM technique. Figure 3 illustrates the PM<sub>2.5</sub> isopleths in Beijing as a function of three combinations of precursors, i.e., NO<sub>X</sub> vs NH<sub>3</sub>, SO<sub>2</sub> vs NH<sub>3</sub>, and VOC+IVOC vs POA; the isopleths for other regions are very similar and thus not shown. The X- and Y-axis of the figures represent the "emission ratio", defined as the ratios of the changed emissions to the emissions in the base case. For example, an emission ratio of 0.7 means the emission of a particular control variable accounts for 70% that of the base case. The colour isopleths represent PM<sub>2.5</sub> concentrations. The comparison shows that the shapes of isopleths derived from both prediction systems generally agree with each other. The agreement is very good for the case of VOC+IVOC vs POA, and for the cases of NO<sub>X</sub> vs NH<sub>3</sub> and SO<sub>2</sub> vs NH<sub>3</sub> when the emission ratios for NO<sub>X</sub> and NH<sub>3</sub> are larger than 0.2. Relatively large errors occur at very low NO<sub>X</sub>/NH<sub>3</sub> emission ratios (< 0.2) due primarily to an extremely strong nonlinearity. Within these low emission ranges, the ERSM technique can capture the general trends in PM<sub>2.5</sub> concentrations in response to emission changes, but the concentration gradients predicted by ERSM are smaller than those given by conventional RSM. More

- studies are needed to further improve the performance of ERSM at very low NO<sub>X</sub>/NH<sub>3</sub>
- 2 emission ratios. Despite the existing errors, the general consistency between RSM and
- 3 ERSM-predicted isopleths demonstrates the stability of the ERSM prediction system. In other
- 4 words, the discrepancies between ERSM and CMAQ/2D-VBS cannot challenge the major
- 5 conclusions on the effectiveness of emission reductions. Finally, as stated in the last
- 6 paragraph, all sensitivity scenarios used in the following discussions have emission ratios  $\geq$
- 7 0.2, since < 0.2 emission reductions are quite rare as limited by the technologically feasible
- 8 reduction potentials (Wang et al., 2014c).

#### 3.2 Response of PM<sub>2.5</sub> concentrations to emissions of air pollutants

- 10 Having demonstrated the reliability of the ERSM prediction system, we employ it to
- 11 investigate the responses of PM<sub>2.5</sub> concentrations to emissions of various pollutants from
- 12 different sectors and regions. We use "PM2.5 sensitivity" defined in Section 2.2 to
- 13 quantitatively characterize the sensitivity of PM<sub>2.5</sub> concentrations to emission changes. Figure
- 4 illustrates the sensitivity of 4-month (January, March, July, and October) mean PM<sub>2.5</sub>
- 15 concentrations to stepped control of individual air pollutants (left panel) and individual
- 16 pollutant-sector combinations (right panel) in the BTH region, which are derived from the
- 17 ERSM technique. The left panel of Fig. 4 can be obtained from both the RSM and ERSM
- prediction systems and their results are consistent, whereas the right panel of Fig. 4, as well as
- 19 the results shown in Fig. 5 and 6 can only be derived from ERSM. Among all pollutants, the
- 4-month mean PM<sub>2.5</sub> concentrations are most sensitive to the emissions of primary inorganic
- 21 PM<sub>2.5</sub> in all five regions, and the PM<sub>2.5</sub> sensitivities vary from 24% to 36% according to
- 22 region. When primary inorganic PM<sub>2.5</sub> emissions from various sectors are differentiated, the
- 23 industry sector is found to make the largest contribution to PM<sub>2.5</sub> concentrations, followed by
- 24 the residential and commercial sectors; the contribution of power plants is negligibly small
- because of smaller emissions and higher stacks. The PM<sub>2.5</sub> sensitivities to primarily inorganic
- 26 PM<sub>2.5</sub> emissions remain constant at various reduction ratios.
- While primary inorganic PM<sub>2.5</sub> makes the largest contribution to PM<sub>2.5</sub> concentrations
- among all air pollutants, the total contributions of all precursors (NO<sub>X</sub>, SO<sub>2</sub>, NH<sub>3</sub>, NMVOC,
- 29 IVOC, and POA), which range between 31% and 48%, exceed that of primary inorganic
- 30 PM<sub>2.5</sub> (24-36%). Among the precursors, PM<sub>2.5</sub> concentrations are primarily sensitive to the
- 31 emissions of NH<sub>3</sub>, NMVOC+IVOC, and POA, and their relative importance differ according
- 32 to reduction ratio. The PM<sub>2.5</sub> sensitivity to NH<sub>3</sub> increases substantially with the increase of

reduction ratio, primarily attributable to the transition from NH<sub>3</sub>-rich to NH<sub>3</sub>-poor regimes when more controls are enforced. The PM<sub>2.5</sub> sensitivies to POA and NMVOC+IVOC, however, decrease slightly with the increase of reduction ratio. This is because that, based on the gas-particle absorptive partitioning theory, organics have a higher tendency to partition into the particle phase at larger OA concentrations. As a result of the nonlinearity, the PM<sub>2.5</sub> sensitivities to POA and NMVOC+IVOC emissions are larger than those to NH<sub>3</sub> emissions at small reduction ratios (e.g., 20%), while it is the other way around at large reduction ratios (e.g., 80%).

1

2

3

4

5

6

7

8

9

10

11

1213

14

15

16

17

18 19

20

21

22

23

24

25

26

27

28

29

30

31

32

The PM<sub>2.5</sub> sensitivity to SO<sub>2</sub> emissions is considerably smaller compared with the three precursors above, and does not change significantly as a function of reduction ratio. From 2007 to 2014 (the base year of this study), both  $SO_2$  emissions and  $SO_4^{2-}$  concentrations in PM<sub>2.5</sub> have been continuously decreasing due to effective control policies (Wang et al., 2017), which partly explains the small sensitivity of PM<sub>2.5</sub> to SO<sub>2</sub> emissions. The response of PM<sub>2.5</sub> concentrations to NO<sub>X</sub> emissions could change from negative to positive with the increase of reduction ratio, which has been reported in several previous studies (Dong et al., 2014; Zhao et al., 2013c; Cai et al., 2016). Small NO<sub>X</sub> emission reductions could lead to increase in O<sub>3</sub> and HO<sub>X</sub> concentrations in several seasons owing to a NMVOC-limited photochemical regime, which on one hand enhances  $SO_4^{2-}$  and SOA formation, and on the other hand, could also increase NO<sub>3</sub> concentrations by accelerating the nocturnal formation of N<sub>2</sub>O<sub>5</sub> and HNO<sub>3</sub> through the  $NO_2 + O_3$  reaction at low temperatures. A substantial reduction in  $NO_X$  emissions, however, transforms the NMVOC-limited regime to a NO<sub>X</sub>-limited regime, resulting in a successive decline in concentrations of O<sub>3</sub>, HO<sub>X</sub>, and most PM<sub>2.5</sub> chemical components. Judging from the our simulation results (Fig. 4), if only the NO<sub>X</sub> emissions within the BTH region are controlled, a very large reduction ratio of about 80% is required to realize a reduction in annual PM<sub>2.5</sub> concentrations in most areas. However, the effects could be distinctly different if NO<sub>X</sub> emissions outside the BTH region are jointly reduced. Our previous studies using the CMAQ model (Zhao et al., 2013c; Wang et al., 2010; Wang et al., 2011) have shown that uniform reductions in NO<sub>X</sub> emissions in the whole China by 23-50% result in considerable annual PM<sub>2.5</sub> reduction over the BTH region. This is because NO<sub>X</sub> emission reductions in upwind regions are more likely to result in a net PM<sub>2.5</sub> decrease compared with local emission reductions, since the photochemistry typically changes from a NMVOC-limited regime in local urban areas at surface to a NO<sub>X</sub>-limited regime in downwind

areas or at upper levels (Xing et al., 2011). The results shown in Fig. 4 also support the above-mentioned pattern and mechanism to some extent: even a 20% NO<sub>X</sub> emission reduction in BTH can lead to PM<sub>2.5</sub> decrease in Northern Hebei, because, as the northernmost region in BTH, it is significantly affected by emissions in other regions within BTH. Note that some recently discovered chemical pathways are missing in the model, such as the oxidation of SO<sub>2</sub> by NO<sub>2</sub> in aerosol water and the SO<sub>2</sub> heterogeneous reactions on the dust surface (Fu et al., 2016; Cheng et al., 2016; Wang et al., 2016a). Incorporation of these processes in the model may affect the simulated responses of PM<sub>2.5</sub> to NO<sub>X</sub> and SO<sub>2</sub> emissions. Regarding emission sectors, the contributions of SO<sub>2</sub> and NO<sub>X</sub> emissions are domiated by "other sources" (sources other than LPS) because they emit larger amount of pollutants at lower height compared with LPS.

The black dotted lines in Fig. 4 show the PM<sub>2.5</sub> sensitivity when all pollutants from all sectors are controlled simultaneously. The sum of PM<sub>2.5</sub> sensitivities to individual pollutants sector combinations (stacked columns) is mostly larger than the sensitivity to all pollutants and sectors (black dotted lines), especially under large reduction ratios. This is mainly attributed to the overlapping effect of two precursors (e.g., SO<sub>2</sub> and NH<sub>3</sub>) involved in the formation of ammonium sulfate and ammonium nitrate. Nevertheless, at small reduction ratios, the sum of individual sensitivities is sometimes smaller, because the negative effects of reducing NO<sub>X</sub> are mitigated when we simultaneously reduce NO<sub>X</sub> emissions from multiple sectors as well as emissions of other air pollutants such as NMVOC. When all pollutants and sectors are controlled together, the PM<sub>2.5</sub> sensitivity generally increases with reduction ratio, indicating that additional air quality benefit could be achieved, larger than the expectation from linear extropolation, if more control measures are implemented.

Figure 5 illustrates the PM<sub>2.5</sub> sensitivities to individual pollutant-sector combinations in each month. The source contribution features are significantly discrepant in different months. The contributions of primary inorganic PM<sub>2.5</sub> emissions to PM<sub>2.5</sub> concentrations are notably higher in January than in other months, which is probably attributed to weaker dilution and slower chemical reactions in January. Regarding different emission sectors of primary inorganic PM<sub>2.5</sub>, the industrial sector plays a dominant role in all months except January, when the residential and commercial sectors make a similar or even larger contribution as compared to the industrial sector. The higher contribution of the residential and commercial sectors in January is on one hand because of the higher emissions due to heating, and on the

other hand explained by weaker vertical mixing in winter, which results in a larger relative contribution of low-level sources. This result highlights the importance of residential and commercial sources for PM<sub>2.5</sub> pollution controls in the winter. The contributions of precursors are dominated by POA and NMVOC+IVOC in January, while in July, NO<sub>X</sub>, SO<sub>2</sub>, and NH<sub>3</sub>, which are known to be precursors of secondary inorganic aerosols, make larger contributions than POA and NMVOC+IVOC. The responses of PM<sub>2.5</sub> concentrations to NO<sub>X</sub> emissions can be opposite in different seasons. Specifically, in July, NO<sub>X</sub> emission reductions always induce decrease in PM<sub>2.5</sub> concentrations due to a NO<sub>X</sub>-limited photochemical regime. In January, however, even a 80% reducion in NO<sub>X</sub> emissions (roughly the maximum technically feasible reduction ratio) could result in a net PM<sub>2.5</sub> increase, as a result of a strong NMVOC-limited regime. To achieve a net PM<sub>2.5</sub> reduction in January, it would be necessary to simultaneously reduce NO<sub>X</sub> emissions outside the BTH region.

We further evaluate the contributions of primary inorganic PM<sub>2.5</sub> and precursor emissions from various regions to PM<sub>2.5</sub> concentrations (Fig. 6, Fig. S6). Here the contributions are quantified by comparing the base case with sensitivity scenarios in which emissions from a specific source are reduced by 80%, which reaches the maximum technologically feasible reduction ratios of major pollutants in most areas (Wang et al., 2014c). Obviously, the contributions of total primary inorganic PM<sub>2.5</sub> emissions in the BTH region are dominated by local sources, which account for over 75% of the total primary inorganic PM<sub>2.5</sub> contributions. When precursor emissions are decomposed into different regions, local sources usually also represent the largest contributors, but precursor emissions from other regions (denoted by "regional precursor emissions" hereafter) could also make significant contributions, depending on regions and seasons. The precursor emissions from the northern part of BTH (e.g., Northern Hebei, Beijing) mainly contribute to local PM<sub>2.5</sub> concentrations, whereas those from the southern part of BTH (e.g., Southern Hebei) significantly affect the PM<sub>2.5</sub> concentrations in both the local region and other regions. Over the BTH, heavy pollution is frequently associated with southerly wind while strong northerly wind often blows away PM<sub>2.5</sub> pollution (Jia et al., 2008; Zheng et al., 2015), which explains the higher contribution of emissions from southern BTH to other regions. Moreover, the importance of regional precursor emissions relative to local ones is remarkably higher in July than in January, which can be explained by the sourtherly monsoon and stronger vertical mixing in summer that favors inter-regional transport of air pollutants. We also examine the contributions of

- 1 emissions outside the BTH region to PM<sub>2.5</sub> concentrations in the five target regions. The
- 2 results reveal that these emissions contribute 24-33% of the 4-month mean PM<sub>2.5</sub>
- 3 concentrations, among which more than 80% could be attributed to precursor emissions.
- 4 Among the four months, the contribution of emissions outside BTH is considerably smaller in
- 5 January (12-21%) as compared to other months (29-38%).

## 3.3 Response of PM<sub>2.5</sub> chemical components to emissions of air pollutants

- 7 Ambient PM<sub>2.5</sub> is comprised of complicated chemical components with distinctly different
- 8 formation pathways. To gain deeper insight into the formation mechanisms and source
- 9 attribution of PM<sub>2.5</sub>, we examine the sensitivities of major PM<sub>2.5</sub> components, including NO<sub>3</sub>,
- 10 SO<sub>4</sub><sup>2</sup>, and OA, to stepped control of individual air pollutants, as shown in Fig. 7 (January and
- July) and Fig. S7 (March and October). NO<sub>3</sub> concentrations are most sensitive to NH<sub>3</sub>
- emissions in all months except July, when the sensitivities of NO<sub>3</sub> concentrations to NH<sub>3</sub> and
- 13 NO<sub>X</sub> emissions are similar. The NO<sub>3</sub> sensitivities to NO<sub>X</sub> emissions differ significantly
- according to season. In most months, NO<sub>3</sub> concentrations are positively correlated with NO<sub>X</sub>
- emissions. In January, however, the sensitivities of NO<sub>3</sub> concentrations to NO<sub>X</sub> emissions are
- mostly negative and could be positive at large reduction ratios, which can be explained by a
- 17 very strong NMVOC-limited photochemical regime, and abundant ice water for
- heterogeneous formation of HNO<sub>3</sub> from N<sub>2</sub>O<sub>5</sub> at cold temperatures. The sensitivites of NO<sub>3</sub><sup>-</sup> to
- 19 both NH<sub>3</sub> and NO<sub>X</sub> emissions show pronounced increasing trends with the increase of
- 20 reduction ratio, in agreement with the strong nonlinearity in these two pollutants described in
- 21 Section 3.2. NMVOC emissions make moderate positive contributions to NO<sub>3</sub>, with the
- 22 largest and smallest contributions occuring in January and July in conjunction with NMVOC-
- 23 limited and NO<sub>X</sub>-limited photochemical regimes, respectively. Finally, SO<sub>2</sub> emissions have
- very small influences on NO<sub>3</sub> concentrations.
- For SO<sub>4</sub><sup>2-</sup>, SO<sub>2</sub> emissions represent the dominant contributor in all months. The sensitivity
- of  $SO_4^{2-}$  concentrations to  $SO_2$  emissions does not change significantly with respect to
- 27 reduction ratio, consistent with the results shown in Section 3.2. The contributions of NH<sub>3</sub>
- 28 emissions to  $SO_4^{2-}$  concentrations are quite small except in October, when NH<sub>3</sub> accounts for
- 29 approximately one fourth the contribution of SO<sub>2</sub>. NO<sub>X</sub> emissions affect SO<sub>4</sub><sup>2-</sup> concentrations
- 30 mainly by altering O<sub>3</sub> and HO<sub>X</sub> concentrations, the effects of which are positive in July at
- 31 large reduction ratios, and mostly negative in other months. NMVOC emissions can impose
- 32 small impact on  $SO_4^{2-}$  concentrations primarily through changing  $O_3$  and  $HO_X$  concentrations.

The emissions of POA and NMVOC+IVOC are obviously two major contributors to OA concentrations. The relative importance of the two is strongly dependent on season. In July, POA and NMVOC+IVOC make similar contributions to OA concentrations, while POA usually contributes more in other months. In January, the contribution of POA could account for about four times those of NMVOC+IVOC. The higher relative contribution of POA emissions in January can be explained by several reasons. First, the POA emissions are relatively higher in January due to residential heating, while the NMVOC emissions from solvent use and biogenic sources are higher in July. Second, lower temperature in winter favors the partitioning of the semi-volatile components comprising POA to the particle phase, whereas higher temperature and stronger radiation in July accelerate the formation of SOA from NMVOC+IVOC. Similar to  $SO_4^{2-}$ , the impact of  $NO_X$  emissions on OA concentrations also works through two pathways. Besides the abovementioned photochemical pathway, NO<sub>X</sub> emission reductions could lead to OA increases due to the fact that SOA yield, defined as the ratio of SOA formation to the consumption of a precursor, is generally higher at a low-NO<sub>X</sub> condition than at a high-NO<sub>X</sub> condition. As an integrated effect, the responses of OA concentrations to NO<sub>X</sub> emissions are negative in most situations.

1

2

3

5

6

7

8

9

10

11

12

13

14

15

16

17

## 3.4 PM<sub>2.5</sub> responses to emission reductions during heavy-pollution episodes

Having shown the responses of monthly-mean PM<sub>2.5</sub> concentrations to pollutant emissions, 18 19 we are also interested in heavy-pollution episodes, in which the source contributions could be 20 quite different from the monthly-mean results, largely due to variations in meteorological 21 conditions. To provide more insight into the control strategies for heavy pollution, we use the 22 ERSM technique to investigate the source contribution features during three typical heavy-23 pollution episodes. We first select 47 heavy-pollution episodes over the BTH region during 24 2013-2015 (Table S7). Subsequently, we employ the Hybrid Single Particle Lagrangian 25 Integrated Trajectory (HYSPLIT) model (Stein et al., 2015) and Concentration Weighted 26 Trajectory (CWT) method (Cheng et al., 2013) to identify the potential source regions for 27 PM<sub>2.5</sub> during each episode, and categorize these episodes according to their source regions. 28 We then select a representative episode from each of three most important pollution types in 29 which the air mass primarily originates from local areas ("Local" type), from the south 30 ("South" type), and from the southeast ("Southeast" type). We give preference to episodes 31 within the four-month simulation period of this study to facilitate a comparison with the 32 monthly-mean source contribution features. For this reason, we select (1) January 5-7, 2014,

1 (2) October 7-11, 2014, and (3) October 29-31, 2014 as representatives of the "Local",

2 "South", and "Southeast" types. The selection of heavy-pollution episodes is detailed in

3 Section 2 of the Supplementary Information.

Figure 8 shows the contribution of precursor and primary inorganic PM<sub>2.5</sub> emissions from individual regions to PM<sub>2.5</sub> concentrations during the three heavy-pollution episodes, and Fig. 9 illustrates the sensitivity of PM<sub>2.5</sub> concentrations to stepped control of individual pollutant-sector combinations. During January 5-7, 2014 ("Local" type), the contributions of local emission sources to PM<sub>2.5</sub> concentrations far exceed those from other regions within BTH as well as from outside of BTH (Fig. 8). In contrast to the monthly mean results (Section 3.2), the contributions of primary inorganic PM<sub>2.5</sub> emissions are comparable to, and even larger than those of precursor emissions in the BTH region. The total contributions of primary PM<sub>2.5</sub> (including POA) account for as high as 70-80% of the contributions of all pollutants within the BTH region, which highlights the crucial importance of primary PM<sub>2.5</sub> controls during this episode. Moreover, the controls of NMVOC, NH<sub>3</sub>, and SO<sub>2</sub> emissions could contribute moderately to reducing PM<sub>2.5</sub> concentrations. However, NO<sub>X</sub> emission reduction induces an increase in PM<sub>2.5</sub> concentrations, even at an 80% reduction ratio. Therefore, effective temporary control measures for this episode should focus on the controls of local emissions, with emphasis laid on primary PM<sub>2.5</sub>.

During October 7-11, 2014 ("South" type), the contributions of emissions outside BTH to PM<sub>2.5</sub> concentrations are as large as 33% in Beijing, and 40-50% in other regions. Within the BTH region, the emissions from Southern Hebei can have similar effects to local emissions on PM<sub>2.5</sub> concentrations in Beijing, indicating a strong long-range transport from the south. In addition, the total contributions of precursor emissions about double those of primary inorganic PM<sub>2.5</sub> emissions. Among all precursors, PM<sub>2.5</sub> concentrations are mainly sensitive to emissions of NH<sub>3</sub>, NMVOC+IVOC, and POA. The sensitivity of PM<sub>2.5</sub> concentrations to NO<sub>X</sub> emissions increases dramatically with reduction ratio. Although small NO<sub>X</sub> reductions may slightly elevate PM<sub>2.5</sub> concentrations, large NO<sub>X</sub> emission reduction (> 50%) can result in significant PM<sub>2.5</sub> reduction. To effectively mitigate PM<sub>2.5</sub> pollution during this episode, we should implement control measures for precursor emissions in both the BTH region (especially the southern part) and regions south of BTH. The NO<sub>X</sub> emissions, if controlled, should be reduced by at least 50% to avoid adverse side effect.

For October 29-31, 2014 ("Southeast" type), PM<sub>2.5</sub> concentrations are also significantly affected by emissions outside the BTH region. Within the BTH region, the PM<sub>2.5</sub> concentrations in Beijing and Northern Hebei are about equally affected by local emissions and emissions from Eastern Hebei and Southern Hebei, while local emissions play dominant roles in other regions. The emissions of both precursor and primary inorganic PM<sub>2.5</sub> within the BTH region make important contributions to PM<sub>2.5</sub> concentrations, and the relative significance of the two is dependent on region. All precursors except NO<sub>X</sub> can contribute considerably to PM<sub>2.5</sub> reductions, and the sensitivity of PM<sub>2.5</sub> to NH<sub>3</sub> increase rapidly with emission ratio. NO<sub>X</sub> emissions are negatively correlated with PM<sub>2.5</sub> concentrations in most cases. Regarding the temporary control strategy for this episode, it is preferable to implement joint controls of primary PM<sub>2.5</sub> and precursors both within and outside the BTH region, with stringent measures over the Eastern and Southern Hebei.

From the analysis above, we conclude that the source contributions are tremendously different in these three episodes, which have been demonstrated to represent some key features of the corresponding pollution types ("Local", "South", and "Southeast" types). Therefore, episode-specific control strategies need to be formulated based on the source contribution features of individual pollution types. Nevertheless, the results of this study are not yet sufficient to guide the development of temporary control strategies for all heavy-pollution episodes, because the conclusions drawn from the three episodes may not be generalized to pollution types. In future studies, we need to simulate more episodes to improve their classification and to comprehensively understand the source contribution features of each pollution type. For a coming heavy-pollution episode, we can predict its pollution type using an air quality forecasting model, and subsequently formulate the temporary control strategies based on the source contribution features of this specific pollution type.

#### 4 Conclusion and implications

- In the present study, we investigated the nonlinear response of PM<sub>2.5</sub> concentrations to emission changes of multiple pollutants from different sectors and regions over the BTH region, using the ERSM technique coupled with the CMAQ/2D-VBS model.
- Among all pollutants, primary inorganic PM<sub>2.5</sub> makes the largest contribution (24-36%) to the 4-month mean PM<sub>2.5</sub> concentrations. The contribution from primary inorganic PM<sub>2.5</sub> is especially high in heavily polluted winter, and is dominated by the industry as well as

residential and commercial sectors. The total contributions of all precursors to PM<sub>2.5</sub> concentrations range between 31% and 48%. Among the precursors, PM<sub>2.5</sub> concentrations are primarily sensitive to the emissions of NH<sub>3</sub>, NMVOC+IVOC, and POA. With the increase of reduction ratio, the sensitivities of PM<sub>2.5</sub> concentrations to pollutant emissions remain roughly constant for primary inorganic PM<sub>2.5</sub> and SO<sub>2</sub>, increase substantially for NH<sub>3</sub> and NO<sub>X</sub>, and decrease slightly for POA and NMVOC+IVOC. The contributions of primary inorganic PM<sub>2.5</sub> emissions to PM<sub>2.5</sub> concentrations are dominated by local emission sources, which account for over 75% of the total primary inorganic PM<sub>2.5</sub> contributions. For precursors, however, emissions from other regions could play similar roles to local emission sources in the summer and over the northern part of BTH. Different PM<sub>2.5</sub> chemical components are associated with distinct source contribution features. The NO<sub>3</sub><sup>-</sup> and SO<sub>4</sub><sup>2-</sup> concentrations are most sensitive to emissions of NH<sub>3</sub> and SO<sub>2</sub>, respectively. The emissions of the POA and NMVOC+IVOC are two major contributors to OA concentrations, with their relative importance depending on season.

The source contribution features are significantly different for three typical heavy-pollution episodes, which belong to three distinct pollution types. The PM<sub>2.5</sub> concentrations in the first episode ("Local" type) are dominated by local sources and primary PM<sub>2.5</sub> emissions, while the second episode ("South" type) is primarily affected by precursor emissions from local and southern regions. The third episode ("Southeast" type) is significantly influenced by emissions of both primary inorganic PM<sub>2.5</sub> and precursors from multiple regions. Future investigations are needed to acquire generalized patterns for the source contributions of various heavy-pollution types.

The results of the present study have important implications for PM<sub>2.5</sub> control policies over the BTH region. First, the controls of primary PM<sub>2.5</sub> emissions should be a priority in PM<sub>2.5</sub> control strategies. Primary PM<sub>2.5</sub>, including primary inorganic PM<sub>2.5</sub> and POA, contribute over half of the 4-month mean PM<sub>2.5</sub> concentrations, which is even higher in the winter when heavy pollution frequently occurs. The industry sector and the residential and commercial sectors represent 85% of the total primariy PM<sub>2.5</sub> emissions, and therefore should be the focus of primary PM<sub>2.5</sub> controls. In particular, we should pay special attention to the residential and commercial sectors, which account for half of the total contribution of primary PM<sub>2.5</sub> emissions to PM<sub>2.5</sub> concentrations in the winter but have been frequently neglected in China's previous control policies. Second, the control policies for NMVOC and IVOC

emissions should be strengthened. The sensitivity of PM<sub>2.5</sub> concentrations to NMVOC+IVOC is one of the largest among all precursors. In particular, the controls of NMVOC and IVOC emissions are very effective for PM<sub>2.5</sub> reduction even at the initial control stage, as indicated by the large sensitivity at small reduction ratios. Moreover, NMVOC reduction is also crucial for the mitigation of O<sub>3</sub> pollution considering a NMVOC-limited regime over the urban and its surrounding areas (Xing et al., 2011). Third, NO<sub>X</sub> emissions should be substantially reduced in both the BTH and other parts of China; in the long run, the reduction ratio should preferably approach their maximum feasible reduction levels. Fourth, more stringent control policies should be enforced in Southern Hebei, which on one hand suffers from the most severe PM<sub>2.5</sub> pollution (Wang et al., 2014b), and on the other hand, significantly affects both local and regional PM<sub>2.5</sub> concentrations. Last but not least, considering the distinct source contributions in different heavy pollution episodes, episode-specific temporary control strategies should be formulated according to the source contribution feature of the specific pollution type.

The present study has a few limitations. First, the establishment of ERSM requires several hundred or over 1000 emission scenarios, although the scenario number needed for a specific number of control variables has already been dramatically reduced as compared to the conventional RSM technique. Studies are needed to further reduce the scenario number but retain the accuracy of the ERSM technique. Second, the current ERSM is developed based on the meteorological conditions simulated for the base year, and has not considered the impact of inter-annual variations in meteorological conditions on the relationships between emissions and PM<sub>2.5</sub> concentrations. Third, although the ERSM-predicted responses of PM<sub>2.5</sub> concentrations to precursor emissions have been demonstrated to agree well with chemical transport model simulations, evaluating the predicted responses against the actual situation in the real atmosphere still represents a major challenge, because it is extremely difficult to artificially perturb emissions in the atmosphere. Last but not the least, the NMVOC and IVOC emissions have been lumped together in this study to reduce the number of control variables. Considering their differences in sources and SOA formation potentials (Jathar et al., 2014; Wu et al., 2017), a detailed quantification of the individual contributions of NMVOC and IVOC emissions from various sources to PM<sub>2.5</sub> concentrations is required in the future to better inform NMVOC/IVOC control policies.

#### Acknowledgements

- 3 This research has been supported by National Science Foundation of China (21625701 &
- 4 21521064), MOST National Key R & D program (2016YFC0207601), Strategic Pilot Project
- 5 of Chinese Academy of Sciences (XDB05030401), the UCLA Sustainable Los Angeles
- 6 Grand Challenge 2016 YZ-50958, and the Jet Propulsion Laboratory, California Institute of
- 7 Technology, under contract with NASA. The simulations were completed on the "Explorer
- 8 100" cluster system of Tsinghua National Laboratory for Information Science and
- 9 Technology.

# 1011

#### References

- Burnett, R. T., Pope, C. A., Ezzati, M., Olives, C., Lim, S. S., Mehta, S., Shin, H. H., Singh,
  G., Hubbell, B., Brauer, M., Anderson, H. R., Smith, K. R., Balmes, J. R., Bruce, N. G.,
  Kan, H. D., Laden, F., Pruss-Ustun, A., Michelle, C. T., Gapstur, S. M., Diver, W. R.,
  and Cohen, A.: An Integrated Risk Function for Estimating the Global Burden of
  Disease Attributable to Ambient Fine Particulate Matter Exposure, Environ Health
- Persp, 122, 397-403, Doi 10.1289/Ehp.1307049, 2014.

  Cai, S. Y., Wang, Y. J., Zhao, B., Wang, S. X., Chang, X., and Hao, J. M.: The impact of the
  "Air Pollution Prevention and Control Action Plan" on PM2.5 concentrations in JingJin-Ji region during 2012-2020, Sci. Total. Environ., in press, DOI

21 10.1016/j.scitotenv.2016.11.188, 2016.

- Cheng, I., Zhang, L., Blanchard, P., Dalziel, J., and Tordon, R.: Concentration-weighted trajectory approach to identifying potential sources of speciated atmospheric mercury at an urban coastal site in Nova Scotia, Canada, Atmos Chem Phys, 13, 6031-6048, 10.5194/acp-13-6031-2013, 2013.
- Cheng, Y. F., Zheng, G. J., Wei, C., Mu, Q., Zheng, B., Wang, Z. B., Gao, M., Zhang, Q., He,
  K. B., Carmichael, G., Pöschl, U., and Su, H.: Reactive nitrogen chemistry in aerosol
  water as a source of sulfate during haze events in China, Sci Adv, 2, e1601530, DOI:
  10.1126/sciadv.1601530, 2016.
- Dong, X. Y., Li, J., Fu, J. S., Gao, Y., Huang, K., and Zhuang, G. S.: Inorganic aerosols responses to emission changes in Yangtze River Delta, China, Sci Total Environ, 481, 522-532, DOI 10.1016/j.scitotenv.2014.02.076, 2014.
- Fu, X., Wang, S. X., Zhao, B., Xing, J., Cheng, Z., Liu, H., and Hao, J. M.: Emission inventory of primary pollutants and chemical speciation in 2010 for the Yangtze River Delta region, China, Atmos Environ, 70, 39-50, DOI 10.1016/j.atmosenv.2012.12.034, 2013.
- Fu, X., Wang, S. X., Chang, X., Cai, S. Y., Xing, J., and Hao, J. M.: Modeling analysis of secondary inorganic aerosols over China: pollution characteristics, and meteorological and dust impacts, Sci Rep-Uk, 6, 10.1038/srep35992, 2016.
- Gordon, T. D., Presto, A. A., May, A. A., Nguyen, N. T., Lipsky, E. M., Donahue, N. M., Gutierrez, A., Zhang, M., Maddox, C., Rieger, P., Chattopadhyay, S., Maldonado, H.,
- 42 Maricq, M. M., and Robinson, A. L.: Secondary organic aerosol formation exceeds

primary particulate matter emissions for light-duty gasoline vehicles, Atmos Chem Phys, 14, 4661-4678, DOI 10.5194/acp-14-4661-2014, 2014a.

3

4

5

6

7

23

24

25

26

27

28

29

- Gordon, T. D., Presto, A. A., Nguyen, N. T., Robertson, W. H., Na, K., Sahay, K. N., Zhang, M., Maddox, C., Rieger, P., Chattopadhyay, S., Maldonado, H., Maricq, M. M., and Robinson, A. L.: Secondary organic aerosol production from diesel vehicle exhaust: impact of aftertreatment, fuel chemistry and driving cycle, Atmos Chem Phys, 14, 4643-4659, DOI 10.5194/acp-14-4643-2014, 2014b.
- Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P. I., and Geron, C.: Estimates of
   global terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and
   Aerosols from Nature), Atmos Chem Phys, 6, 3181-3210, 2006.
- Hakami, A., Seinfeld, J. H., Chai, T. F., Tang, Y. H., Carmichael, G. R., and Sandu, A.:
  Adjoint sensitivity analysis of ozone nonattainment over the continental United States,
  Environ Sci Technol, 40, 3855-3864, DOI 10.1021/Es052135g, 2006.
- Han, X., Zhang, M. G., Zhu, L. Y., and Skorokhod, A.: Assessment of the impact of emissions reductions on air quality over North China Plain, Atmos Pollut Res, 7, 249-259, 10.1016/j.apr.2015.09.009, 2016.
- Hennigan, C. J., Miracolo, M. A., Engelhart, G. J., May, A. A., Presto, A. A., Lee, T.,
  Sullivan, A. P., McMeeking, G. R., Coe, H., Wold, C. E., Hao, W. M., Gilman, J. B.,
  Kuster, W. C., de Gouw, J., Schichtel, B. A., Collett, J. L., Kreidenweis, S. M., and
  Robinson, A. L.: Chemical and physical transformations of organic aerosol from the
  photo-oxidation of open biomass burning emissions in an environmental chamber,
  Atmos Chem Phys, 11, 7669-7686, DOI 10.5194/acp-11-7669-2011, 2011.
  - Iman, R. L., Davenport, J. M., and Zeigler, D. K.: Latin Hypercube Sampling (Program User's Guide), Sandia National Laboratories, Albuquerque, NM, U.S., 78 pp., 1980.
  - IPCC: Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, edited by: Stocker, T. F., Qin, D., Plattner, G.-K., Tignor, M., Allen, S. K., Boschung, J., Nauels, A., Xia, Y., Bex, V., and Midgley, P. M., Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 1535 pp., 2013.
- Jathar, S. H., Gordon, T. D., Hennigan, C. J., Pye, H. O. T., Pouliot, G., Adams, P. J., Donahue, N. M., and Robinson, A. L.: Unspeciated organic emissions from combustion sources and their influence on the secondary organic aerosol budget in the United States, P Natl Acad Sci USA, 111, 10473-10478, DOI 10.1073/pnas.1323740111, 2014.
- Jia, Y. T., Rahn, K. A., He, K. B., Wen, T. X., and Wang, Y. S.: A novel technique for quantifying the regional component of urban aerosol solely from its sawtooth cycles, J Geophys Res-Atmos, 113, 10.1029/2008jd010389, 2008.
- Li, J. W., and Han, Z. W.: A modeling study of severe winter haze events in Beijing and its neighboring regions, Atmos Res, 170, 87-97, 10.1016/j.atmosres.2015.11.009, 2016.
- Li, M., Zhang, Q., Kurokawa, J., Woo, J. H., He, K. B., Lu, Z., Ohara, T., Song, Y., Streets, D. G., Carmichael, G. R., Cheng, Y. F., Hong, C. P., Huo, H., Jiang, X. J., Kang, S. C., Liu, F., Su, H., and Zheng, B.: MIX: a mosaic Asian anthropogenic emission inventory for the MICS-Asia and the HTAP projects, Atmos Chem Phys Discuss, 15, 34813-34869, doi:10.5194/acpd-15-34813-2015, 2015a.
- Li, X., Zhang, Q., Zhang, Y., Zheng, B., Wang, K., Chen, Y., Wallington, T. J., Han, W. J., Shen, W., Zhang, X. Y., and He, K. B.: Source contributions of urban PM2.5 in the Beijing-Tianjin-Hebei region: Changes between 2006 and 2013 and relative impacts of emissions and meteorology, Atmos Environ, 123, 229-239,

48 10.1016/j.atmosenv.2015.10.048, 2015b.

1 Lim, S. S., Vos, T., Flaxman, A. D., Danaei, G., Shibuya, K., Adair-Rohani, H., AlMazroa, M. 2 A., Amann, M., Anderson, H. R., Andrews, K. G., Aryee, M., Atkinson, C., Bacchus, L. 3 J., Bahalim, A. N., Balakrishnan, K., Balmes, J., Barker-Collo, S., Baxter, A., Bell, M. 4 L., Blore, J. D., Blyth, F., Bonner, C., Borges, G., Bourne, R., Boussinesq, M., Brauer, 5 M., Brooks, P., Bruce, N. G., Brunekreef, B., Bryan-Hancock, C., Bucello, C., 6 Buchbinder, R., Bull, F., Burnett, R. T., Byers, T. E., Calabria, B., Carapetis, J., 7 Carnahan, E., Chafe, Z., Charlson, F., Chen, H., Chen, J. S., Cheng, A. T.-A., Child, J. 8 C., Cohen, A., Colson, K. E., Cowie, B. C., Darby, S., Darling, S., Davis, A., 9 Degenhardt, L., Dentener, F., Des Jarlais, D. C., Devries, K., Dherani, M., Ding, E. L., 10 Dorsey, E. R., Driscoll, T., Edmond, K., Ali, S. E., Engell, R. E., Erwin, P. J., Fahimi, S., 11 Falder, G., Farzadfar, F., Ferrari, A., Finucane, M. M., Flaxman, S., Fowkes, F. G. R., Freedman, G., Freeman, M. K., Gakidou, E., Ghosh, S., Giovannucci, E., Gmel, G., 12 13 Graham, K., Grainger, R., Grant, B., Gunnell, D., Gutierrez, H. R., Hall, W., Hoek, H. 14 W., Hogan, A., Hosgood Iii, H. D., Hoy, D., Hu, H., Hubbell, B. J., Hutchings, S. J., 15 Ibeanusi, S. E., Jacklyn, G. L., Jasrasaria, R., Jonas, J. B., Kan, H., Kanis, J. A., 16 Kassebaum, N., Kawakami, N., Khang, Y.-H., Khatibzadeh, S., Khoo, J.-P., Kok, C., 17 Laden, F., Lalloo, R., Lan, Q., Lathlean, T., Leasher, J. L., Leigh, J., Li, Y., Lin, J. K., 18 Lipshultz, S. E., London, S., Lozano, R., Lu, Y., Mak, J., Malekzadeh, R., Mallinger, L., 19 Marcenes, W., March, L., Marks, R., Martin, R., McGale, P., McGrath, J., Mehta, S., 20 Memish, Z. A., Mensah, G. A., Merriman, T. R., Micha, R., Michaud, C., Mishra, V., Hanafiah, K. M., Mokdad, A. A., Morawska, L., Mozaffarian, D., Murphy, T., Naghavi, 21 22 M., Neal, B., Nelson, P. K., Nolla, J. M., Norman, R., Olives, C., Omer, S. B., Orchard, 23 J., Osborne, R., Ostro, B., Page, A., Pandey, K. D., Parry, C. D. H., Passmore, E., Patra, 24 J., Pearce, N., Pelizzari, P. M., Petzold, M., Phillips, M. R., Pope, D., Pope Iii, C. A., Powles, J., Rao, M., Razavi, H., Rehfuess, E. A., Rehm, J. T., Ritz, B., Rivara, F. P., 25 26 Roberts, T., Robinson, C., Rodriguez-Portales, J. A., Romieu, I., Room, R., Rosenfeld, 27 L. C., Roy, A., Rushton, L., Salomon, J. A., Sampson, U., Sanchez-Riera, L., Sanman, 28 E., Sapkota, A., Seedat, S., Shi, P., Shield, K., Shivakoti, R., Singh, G. M., Sleet, D. A., 29 Smith, E., Smith, K. R., Stapelberg, N. J. C., Steenland, K., Stöckl, H., Stovner, L. J., 30 Straif, K., Straney, L., Thurston, G. D., Tran, J. H., Van Dingenen, R., van Donkelaar, 31 A., Veerman, J. L., Vijayakumar, L., Weintraub, R., Weissman, M. M., White, R. A., 32 Whiteford, H., Wiersma, S. T., Wilkinson, J. D., Williams, H. C., Williams, W., Wilson, 33 N., Woolf, A. D., Yip, P., Zielinski, J. M., Lopez, A. D., Murray, C. J. L., and Ezzati, 34 M.: A comparative risk assessment of burden of disease and injury attributable to 67 risk 35 factors and risk factor clusters in 21 regions, 1990–2010: a systematic analysis for the 36 2010, The Burden of Disease Study Lancet, 380, 2224-2260, 37 http://dx.doi.org/10.1016/S0140-6736(12)61766-8, 2012. 38

Liu, J., Mauzerall, D. L., Chen, Q., Zhang, Q., Song, Y., Peng, W., Klimont, Z., Qiu, X. H.,
 Zhang, S. Q., Hu, M., Lin, W. L., Smith, K. R., and Zhu, T.: Air pollutant emissions from Chinese households: A major and underappreciated ambient pollution source, P
 Natl Acad Sci USA, 113, 7756-7761, 10.1073/pnas.1604537113, 2016.

42 Lv, L. Y., and Li, H. Y.: Economic evaluation of the health effect of PM10 and PM2.5 43 pollution over the Beijing-Tianjin-Hebei region, Acta Scientiarum Naturalium 44 Universitatis Nankaiensis, 69-77, 2016.

Qin, Y., and Xie, S. D.: Historical estimation of carbonaceous aerosol emissions from biomass open burning in China for the period 1990-2005, Environ Pollut, 159, 3316-3323, DOI 10.1016/j.envpol.2011.08.042, 2011.

Russell, A., Milford, J., Bergin, M. S., Mcbride, S., Mcnair, L., Yang, Y., Stockwell, W. R., and Croes, B.: Urban Ozone Control and Atmospheric Reactivity of Organic Gases, Science, 269, 491-495, DOI 10.1126/science.269.5223.491, 1995.

4

5

6

15

16

17

21

22

23

24

25

26

- Sandu, A., Daescu, D. N., Carmichael, G. R., and Chai, T. F.: Adjoint sensitivity analysis of regional air quality models, J Comput Phys, 204, 222-252, DOI 10.1016/j.jcp.2004.10.011, 2005.
- Santner, T. J., Williams, B. J., and Notz, W.: The Design and Analysis of Computer Experiments, Springer Verlag, New York, U.S., 283 pp., 2003.
- Seinfeld, J. H., Bretherton, C., Carslaw, K. S., Coe, H., DeMott, P. J., Dunlea, E. J., Feingold,
   G., Ghan, S., Guenther, A. B., Kahn, R., Kraucunas, I., Kreidenweis, S. M., Molina, M.
   J., Nenes, A., Penner, J. E., Prather, K. A., Ramanathan, V., Ramaswamy, V., Rasch, P.
   J., Ravishankara, A. R., Rosenfeld, D., Stephens, G., and Wood, R.: Improving our
   fundamental understanding of the role of aerosol-cloud interactions in the climate
   system, P Natl Acad Sci USA, 113, 5781-5790, 10.1073/pnas.1514043113, 2016.
  - Stein, A. F., Draxler, R. R., Rolph, G. D., Stunder, B. J. B., Cohen, M. D., and Ngan, F.: NOAA's HYSPLIT atmospheric transport and dispersion modeling system, B Am Meteorol Soc, 96, 2059-2077, 10.1175/bams-d-14-00110.1, 2015.
- Streets, D. G., Fu, J. S., Jang, C. J., Hao, J. M., He, K. B., Tang, X. Y., Zhang, Y. H., Wang,
   Z. F., Li, Z. P., and Zhang, Q.: Air quality during the 2008 Beijing Olympic Games,
   Atmos Environ, 41, 480-492, DOI 10.1016/j.atmosenv.2006.08.046, 2007.
  - The State Council of the People's Republic of China: Notice to issue the "Air Pollution Prevention and Control Action Plan": <a href="http://www.gov.cn/zwgk/2013-09/12/content">http://www.gov.cn/zwgk/2013-09/12/content</a> 2486773.htm, access: September 9, 2016, 2013.
  - U.S. Environmental Protection Agency: Technical support document for the proposed PM NAAQS rule: Response Surface Modeling[R/OL]: <a href="http://www.epa.gov/scram001/reports/pmnaaqs\_tsd\_rsm\_all\_021606.pdf">http://www.epa.gov/scram001/reports/pmnaaqs\_tsd\_rsm\_all\_021606.pdf</a>, access: 2015-02-01, 2006.
- van Donkelaar, A., Martin, R. V., Brauer, M., and Boys, B. L.: Use of satellite observations for long-term exposure assessment of global concentrations of fine particulate matter, Environmental health perspectives, 123, 135, 2015.
- 31 Wang, G. H., Zhang, R. Y., Gomez, M. E., Yang, L. X., Zamora, M. L., Hu, M., Lin, Y., Peng, 32 J. F., Guo, S., Meng, J. J., Li, J. J., Cheng, C. L., Hu, T. F., Ren, Y. Q., Wang, Y. S., 33 Gao, J., Cao, J. J., An, Z. S., Zhou, W. J., Li, G. H., Wang, J. Y., Tian, P. F., Marrero-34 Ortiz, W., Secrest, J., Du, Z. F., Zheng, J., Shang, D. J., Zeng, L. M., Shao, M., Wang, 35 W. G., Huang, Y., Wang, Y., Zhu, Y. J., Li, Y. X., Hu, J. X., Pan, B., Cai, L., Cheng, Y. 36 T., Ji, Y. M., Zhang, F., Rosenfeld, D., Liss, P. S., Duce, R. A., Kolb, C. E., and Molina, M. J.: Persistent sulfate formation from London Fog to Chinese haze, P Natl Acad Sci 37 38 USA, 113, 13630-13635, 10.1073/pnas.1616540113, 2016a.
- Wang, J. D., Wang, S. X., Jiang, J. K., Ding, A. J., Zheng, M., Zhao, B., Wong, D. C., Zhou,
  W., Zheng, G. J., Wang, L., Pleim, J. E., and Hao, J. M.: Impact of aerosol-meteorology
  interactions on fine particle pollution during China's severe haze episode in January
  2013, Environ Res Lett, 9, 094002, DOI 10.1088/1748-9326/9/9/094002, 2014a.
- Wang, J. D., Xing, J., Mathur, R., Pleim, J. E., Wang, S. X., Hogrefe, C., Gan, C.-M., Wong,
   D. C., and Hao, J. M.: Historical Trends in PM2.5-Related Premature Mortality during
   1990-2010 across the Northern Hemisphere, Environ Health Persp, in press, DOI
   10.1289/EHP298, 2016b.
- Wang, J. D., Zhao, B., Wang, S. X., Yang, F. M., Xing, J., Morawska, L., Ding, A. J., Kulmala, M., Kerminen, V. M., Kujansuu, J., Wang, Z. F., Ding, D. A., Zhang, X. Y.,

Wang, H. B., Tian, M., Petaja, T., Jiang, J. K., and Hao, J. M.: Particulate matter pollution over China and the effects of control policies, Sci Total Environ, 584, 426-447, 10.1016/j.scitotenv.2017.01.027, 2017.

4

5

6

7

8

15

16

17

18 19

20

21

22

- Wang, L. T., Hao, J. M., He, K. B., Wang, S. X., Li, J. H., Zhang, Q., Streets, D. G., Fu, J. S., Jang, C. J., Takekawa, H., and Chatani, S.: A modeling study of coarse particulate matter pollution in Beijing: Regional source contributions and control implications for the 2008 summer Olympics, J Air Waste Manage, 58, 1057-1069, Doi 10.3155/1047-3289.58.8.1057, 2008.
- 9 Wang, L. T., Jang, C., Zhang, Y., Wang, K., Zhang, Q. A., Streets, D., Fu, J., Lei, Y., 10 Schreifels, J., He, K. B., Hao, J. M., Lam, Y. F., Lin, J., Meskhidze, N., Voorhees, S., 11 Evarts, D., and Phillips, S.: Assessment of air quality benefits from national air pollution 12 control policies in China. Part II: Evaluation of air quality predictions and air quality 13 assessment, Environ, 44. 3449-3457, benefits Atmos DOI 14 10.1016/j.atmosenv.2010.05.058, 2010.
  - Wang, L. T., Wei, Z., Yang, J., Zhang, Y., Zhang, F. F., Su, J., Meng, C. C., and Zhang, Q.: The 2013 severe haze over southern Hebei, China: model evaluation, source apportionment, and policy implications, Atmos Chem Phys, 14, 3151-3173, 10.5194/acp-14-3151-2014, 2014b.
  - Wang, S. X., and Zhang, C. Y.: Spatial and temporal distribution of air pollutant emissions from open burning of crop residues in China, Sciencepaper Online, 3, 1-6, 2008.
  - Wang, S. X., Xing, J., Jang, C. R., Zhu, Y., Fu, J. S., and Hao, J. M.: Impact assessment of ammonia emissions on inorganic aerosols in east China using response surface modeling technique, Environ Sci Technol, 45, 9293-9300, DOI 10.1021/Es2022347, 2011.
- Wang, S. X., Zhao, B., Cai, S. Y., Klimont, Z., Nielsen, C. P., Morikawa, T., Woo, J. H., Kim,
  Y., Fu, X., Xu, J. Y., Hao, J. M., and He, K. B.: Emission trends and mitigation options
  for air pollutants in East Asia, Atmos Chem Phys, 14, 6571-6603, DOI 10.5194/acp-14-6571-2014, 2014c.
- Wang, S. X., Zhao, B., Wu, Y., and Hao, J. M.: Target and measures to prevent and control ambient fine particle pollution in China, Chinese Journal of Environmental Management, 37-43, 2015.
- Wang, Y. J., Bao, S. W., Wang, S. X., Hu, Y. T., Shi, X., Wang, J. D., Zhao, B., Jiang, J. K.,
  Zheng, M., Wu, M. H., Russell, A. G., Wang, Y. H., and Hao, J. M.: Local and regional
  contributions to fine particulate matter in Beijing during heavy haze episodes, Sci Total
  Environ, in press, DOI: 10.1016/j.scitotenv.2016.12.127, 2016c.
- Wu, W. J.: Health Effect Attributed to Ambient Fine Particle Pollution in the Beijing-Tianjin Hebei Region and its Source Apportionment, Doctor, School of Environment, Tsinghua
   University, Beijing, China, 98 pp., 2016.
- Wu, W. J., Zhao, B., Wang, S. X., and Hao, J. M.: Ozone and secondary organic aerosol formation potential from anthropogenic volatile organic compounds emissions in China, Journal of Environmental Sciences, 53, 224-237, 10.1016/j.jes.2016.03.025, 2017.
- Xing, J., Wang, S. X., Jang, C., Zhu, Y., and Hao, J. M.: Nonlinear response of ozone to precursor emission changes in China: a modeling study using response surface methodology, Atmos Chem Phys, 11, 5027-5044, DOI 10.5194/acp-11-5027-2011, 2011.
- Yang, Y. J., Wilkinson, J. G., and Russell, A. G.: Fast, direct sensitivity analysis of multidimensional photochemical models, Environ Sci Technol, 31, 2859-2868, DOI 10.1021/Es970117w, 1997.

- Ying, Q., Wu, L., and Zhang, H. L.: Local and inter-regional contributions to PM2.5 nitrate and sulfate in China, Atmos Environ, 94, 582-592, 10.1016/j.atmosenv.2014.05.078, 2014.
- Yu, L. D., Wang, G. F., Zhang, R. J., Zhang, L. M., Song, Y., Wu, B. B., Li, X. F., An, K.,
  and Chu, J. H.: Characterization and Source Apportionment of PM2.5 in an Urban Environment in Beijing, Aerosol Air Qual Res, 13, 574-583, 10.4209/aaqr.2012.07.0192,
  2013.
- Zhang, L., Shao, J. Y., Lu, X., Zhao, Y. H., Hu, Y. Y., Henze, D. K., Liao, H., Gong, S. L.,
   and Zhang, Q.: Sources and Processes Affecting Fine Particulate Matter Pollution over
   North China: An Adjoint Analysis of the Beijing APEC Period, Environ Sci Technol, 50,
   8731-8740, 10.1021/acs.est.6b03010, 2016.
- Zhang, W., Guo, J. H., Sun, Y. L., Yuan, H., Zhuang, G. S., Zhuang, Y. H., and Hao, Z. P.:
   Source apportionment for,urban PM10 and PM2.5 in the Beijing area, Chinese Sci Bull,
   52, 608-615, 10.1007/s11434-007-0076-5, 2007.
- Zhao, B., Wang, S. X., Dong, X. Y., Wang, J. D., Duan, L., Fu, X., Hao, J. M., and Fu, J.:
   Environmental effects of the recent emission changes in China: implications for particulate matter pollution and soil acidification, Environ Res Lett, 8, 024031, DOI 10.1088/1748-9326/8/2/024031, 2013a.
- Zhao, B., Wang, S. X., Liu, H., Xu, J. Y., Fu, K., Klimont, Z., Hao, J. M., He, K. B., Cofala,
  J., and Amann, M.: NOx emissions in China: historical trends and future perspectives,
  Atmos Chem Phys, 13, 9869-9897, DOI 10.5194/acp-13-9869-2013, 2013b.
- Zhao, B., Wang, S. X., Wang, J. D., Fu, J. S., Liu, T. H., Xu, J. Y., Fu, X., and Hao, J. M.:
   Impact of national NOx and SO2 control policies on particulate matter pollution in
   China, Atmos Environ, 77, 453-463, DOI 10.1016/j.atmosenv.2013.05.012, 2013c.
- Zhao, B., Wang, S. X., Donahue, N. M., Chuang, W., Hildebrandt Ruiz, L., Ng, N. L., Wang,
   Y. J., and Hao, J. M.: Evaluation of one-dimensional and two-dimensional volatility
   basis sets in simulating the aging of secondary organic aerosols with smog-chamber
   experiments, Environ Sci Technol, 49, 2245-2254, DOI 10.1021/es5048914, 2015a.

30

31

32

- Zhao, B., Wang, S. X., Xing, J., Fu, K., Fu, J. S., Jang, C., Zhu, Y., Dong, X. Y., Gao, Y., Wu, W. J., Wang, J. D., and Hao, J. M.: Assessing the nonlinear response of fine particles to precursor emissions: development and application of an extended response surface modeling technique v1.0, Geosci Model Dev, 8, 115-128, DOI 10.5194/gmd-8-115-2015, 2015b.
- Zhao, B., Wang, S. X., Donahue, N. M., Jathar, S. H., Huang, X. F., Wu, W. J., Hao, J. M.,
   and Robinson, A. L.: Quantifying the effect of organic aerosol aging and intermediate volatility emissions on regional-scale aerosol pollution in China, Sci Rep-Uk, 6,
   10.1038/srep28815, 2016.
- Zhao, B., Liou, K.-N., Gu, Y., Jiang, J. H., Li, Q., Fu, R., Huang, L., Liu, X., Shi, X., Su, H.,
  and He, C.: A water vapor modulated aerosol impact on ice crystal size, Atmos Chem
  Phys Discuss, DOI 10.5194/acp-2017-548, DOI 10.5194/acp-2017-548, 2017a.
- Zhao, B., Liou, K. N., Gu, Y., Li, Q. B., Jiang, J. H., Su, H., He, C. L., Tseng, H. L. R., Wang,
  S. X., Liu, R., Qi, L., Lee, W. L., and Hao, J. M.: Enhanced PM2.5 pollution in China due to aerosol-cloud interactions, Sci Rep-Uk, 7, 10.1038/s41598-017-04096-8, 2017b.
- Zhao, Y., Wang, S. X., Duan, L., Lei, Y., Cao, P. F., and Hao, J. M.: Primary air pollutant emissions of coal-fired power plants in China: Current status and future prediction, Atmos Environ, 42, 8442-8452, DOI 10.1016/j.atmosenv.2008.08.021, 2008.
- Zheng, G. J., Duan, F. K., Su, H., Ma, Y. L., Cheng, Y., Zheng, B., Zhang, Q., Huang, T.,
  Kimoto, T., Chang, D., Poschl, U., Cheng, Y. F., and He, K. B.: Exploring the severe

winter haze in Beijing: the impact of synoptic weather, regional transport and heterogeneous reactions, Atmos Chem Phys, 15, 2969-2983, 10.5194/acp-15-2969-2015, 2015.

# 1 Tables and figures

# 2 Table 1. Description of the RSM/ERSM prediction systems developed in this study.

Method	Control variables	Control scenarios			
Conventional RSM technique	5 control variables: total emissions of $NO_X$ , $SO_2$ , $NH_3$ , $NMVOC+IVOC$ , and $POA$	<ul> <li>101 control scenarios:</li> <li>1) 1 CMAQ/2D-VBS base case;</li> <li>2) 100<sup>a</sup> scenarios generated by applying LHS method for the 5 variables.</li> </ul>			
ERSM technique	55 control variables in total:  11 control variables in each of the 5 regions, including 7 nonlinear control variables, i.e.,  1) NO <sub>X</sub> /large point sources (LPS) <sup>b</sup> 2) NO <sub>X</sub> /other sources  3) SO <sub>2</sub> /LPS  4) SO <sub>2</sub> /other sources  5) NH <sub>3</sub> /all sources  6) NMVOC+IVOC/all sources  7) POA/all sources  and 4 linear control variables, i.e.,  8) Primary inorganic PM <sub>2.5</sub> /power plants  9) Primary inorganic PM <sub>2.5</sub> /Industry  10) Primary inorganic PM <sub>2.5</sub> /residential & commercial  11) Primary inorganic PM <sub>2.5</sub> /transportation	, , , , , , , , , , , , , , , , , , , ,			

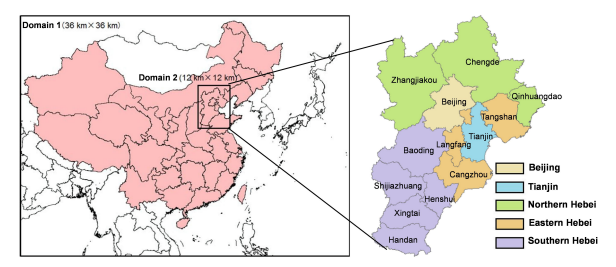
<sup>3</sup> a 100 and 200 scenarios are needed for the response surfaces for 5 and 7 variables, respectively (Xing et al.,

<sup>4 2011;</sup> Wang et al., 2011).

<sup>5</sup> b LPS includes power plants, iron and steel plants, and cement plants

Table 2. Comparison between ERSM-predicted and CMAQ/2D-VBS-simulated PM<sub>2.5</sub> concentrations for
 54 out-of-sample scenarios.

Month	Variable	Statistical index	Beijing	Tianjin	Northern Hebei	Eastern Hebei	Southern Hebei
Jan	PM <sub>2.5</sub> concentration	R	0.998	0.998	0.995	0.997	0.997
		MNE (%)	0.52	0.55	0.64	0.67	0.60
		Maximum NE (%)	7.56	6.98	10.67	8.01	8.03
		95% percentile of NEs (%)	1.61	2.86	2.92	3.46	3.02
		NME (%)	0.44	0.46	0.57	0.53	0.53
	PM <sub>2.5</sub> response	R	0.998	0.998	0.995	0.997	0.997
		NME (%)	3.36	3.48	4.25	4.00	3.88
Mar	PM <sub>2.5</sub> concentration	R	0.999	0.996	0.998	0.995	0.999
		MNE (%)	0.37	0.54	0.39	0.57	0.49
		Maximum NE (%)	3.75	6.58	4.30	5.04	3.22
		95% percentile of NEs (%)	1.53	3.15	2.03	4.35	2.03
		NME (%)	0.31	0.45	0.34	0.49	0.42
	PM <sub>2.5</sub> response	R	0.999	0.996	0.998	0.995	0.999
		NME (%)	2.38	4.32	2.70	4.55	3.59
Jul	PM <sub>2.5</sub> concentration	R	0.997	0.998	0.998	0.999	0.999
		MNE (%)	0.94	0.54	0.46	0.37	0.47
		Maximum NE (%)	5.05	5.02	4.65	1.83	3.62
		95% percentile of NEs (%)	3.47	2.33	2.17	1.49	1.87
		NME (%)	0.80	0.47	0.41	0.33	0.39
	PM <sub>2.5</sub> response	R	0.997	0.998	0.998	0.999	0.999
		NME (%)	4.97	3.71	2.80	2.58	2.78
Oct	PM <sub>2.5</sub> concentration	R	0.996	0.994	0.999	0.999	0.999
		MNE (%)	0.83	0.70	0.36	0.39	0.36
		Maximum NE (%)	8.90	11.19	3.79	3.90	2.46
		95% percentile of NEs (%)	3.04	3.50	1.44	2.10	1.64
		NME (%)	0.67	0.58	0.30	0.35	0.32
	PM <sub>2.5</sub> response	R	0.996	0.994	0.999	0.999	0.999
		NME (%)	4.51	5.64	2.20	3.29	2.79
		( · - )					



2 Figure 1. Double nesting domains used in CMAQ/2D-VBS simulation (left) and the definition

- 3 of five target regions in the innermost domain, denoted by different colours (right). The grey
- 4 lines in the right figure represent the boundaries of prefecture-level cities.

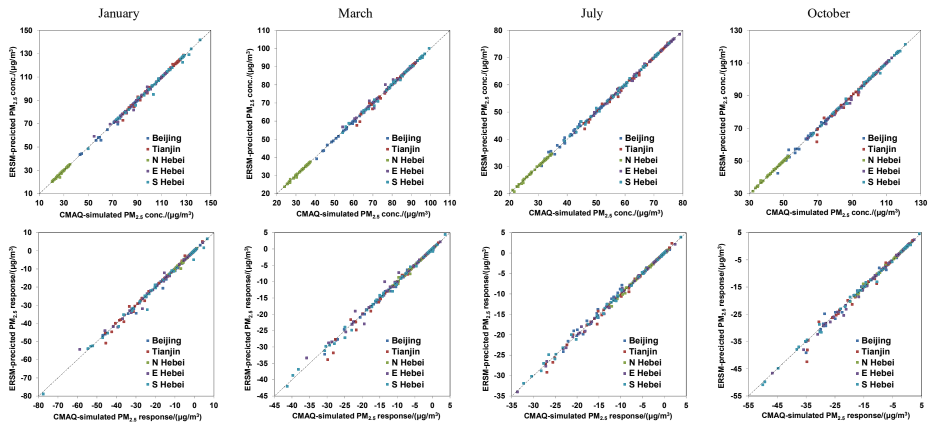


Figure 2. Comparison of PM<sub>2.5</sub> concentrations (top row) and PM<sub>2.5</sub> responses (bottom row) predicted by the ERSM technique with out-of-

2 sample CMAQ/2D-VBS simulations. The dashed line is the one-to-one line indicating perfect agreement.

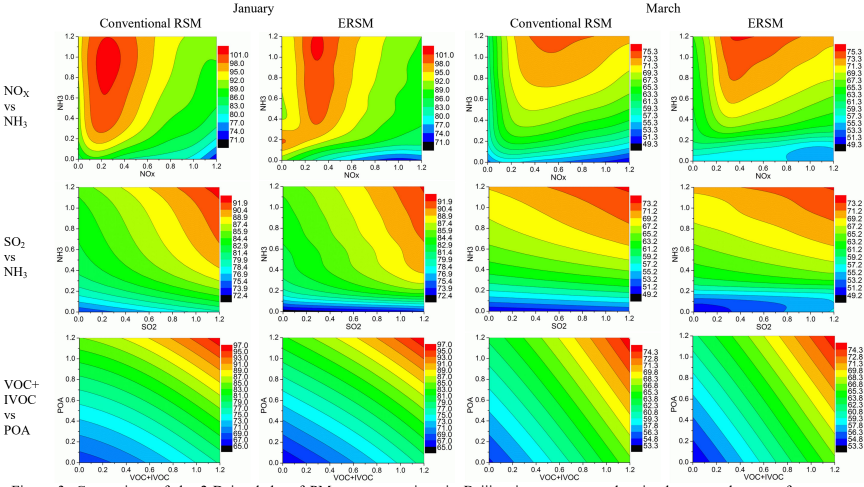


Figure 3. Comparison of the 2-D isopleths of  $PM_{2.5}$  concentrations in Beijing in response to the simultaneous changes of precursor emissions in all five regions derived from the conventional RSM technique and the ERSM technique. The X- and Y-axis represent the emission ratio, defined as the ratios of the changed emissions to the emissions in the base case. The colour contours represent  $PM_{2.5}$  concentrations (unit:  $\mu g m^{-3}$ ).



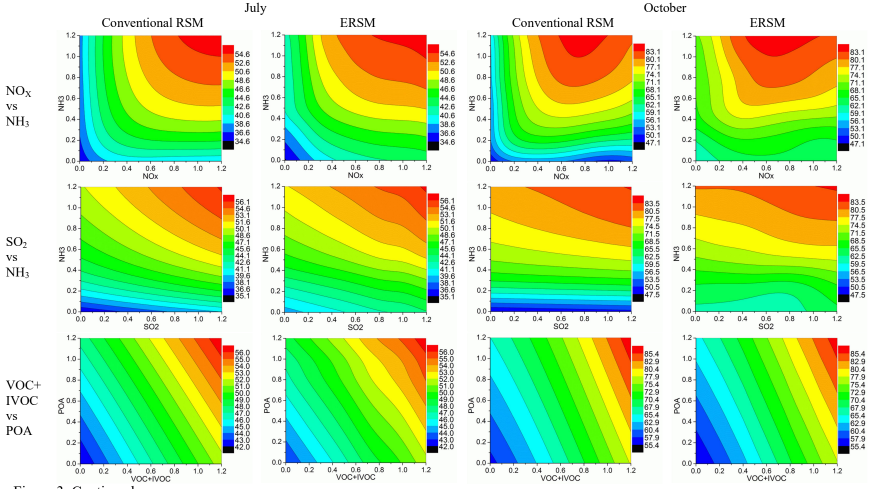


Figure 3. Continued.

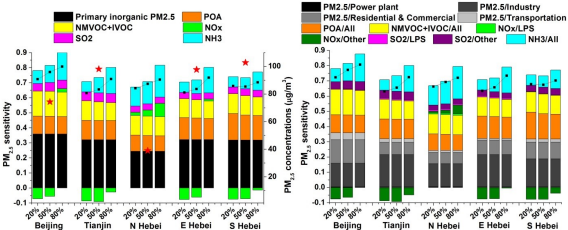


Figure 4. Sensitivity of 4-month mean  $PM_{2.5}$  concentrations to stepped control of individual air pollutants (left) and individual pollutant-sector combinations (right). The X-axis shows the reduction ratio (= 1 – emission ratio). The Y-axis shows  $PM_{2.5}$  sensitivity, which is defined as the change ratio of concentration divided by the reduction ratio of emissions. The coloured bars denote the  $PM_{2.5}$  sensitivities when a particular emission source is controlled while the others stay the same as the base case; the black dotted line denotes the  $PM_{2.5}$  sensitivity when all emission sources are controlled simultaneously. The red stars represent  $PM_{2.5}$  concentrations in the base case.

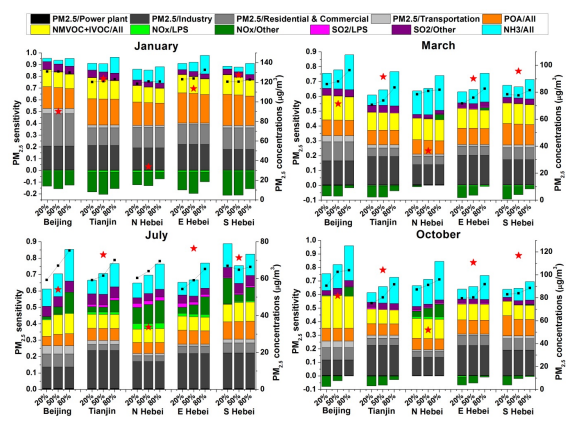


Figure 5. Sensitivity of monthly mean PM<sub>2.5</sub> concentrations to stepped control of individual air pollutants from individual sectors in January, March, July, and October. The meanings of X-axis, Y-axis, coloured bars, black dotted lines, and red stars are the same as Fig. 4.

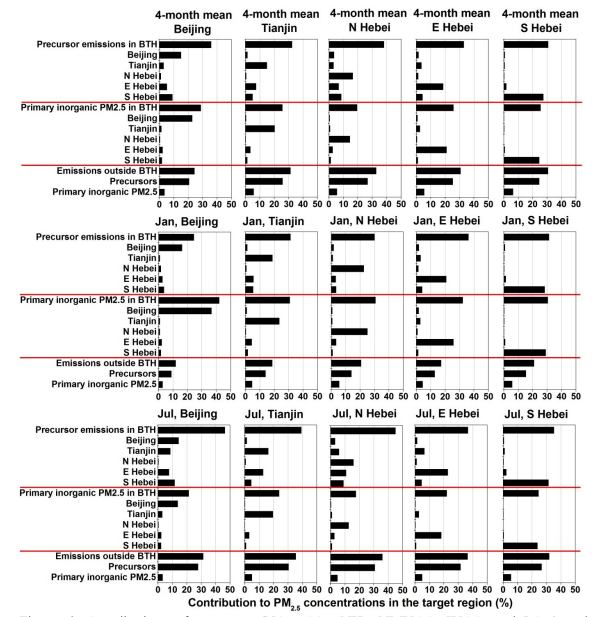


Figure 6. Contributions of precursor (NO<sub>X</sub>, SO<sub>2</sub>, NH<sub>3</sub>, NMVOC, IVOC, and POA) and primary inorganic PM<sub>2.5</sub> emissions from individual regions to PM<sub>2.5</sub> concentrations. The contributions are quantified by comparing the base case with sensitivity scenarios in which emissions from a specific source are reduced by 80%. This figure illustrates contributions to 4-month mean PM<sub>2.5</sub> concentrations and monthly mean PM<sub>2.5</sub> concentrations in January and July. The results for March and October are given in Fig. S6.

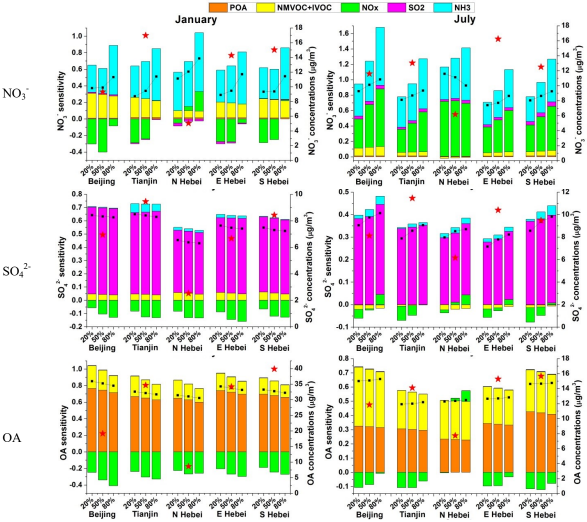


Figure 7. Sensitivity of monthly mean NO<sub>3</sub>-, SO<sub>4</sub><sup>2</sup>-, and OA concentrations to stepped control of individual air pollutants in January and July. The meanings of X-axis, Y-axis, coloured bars, black dotted lines, and red stars are the same as Fig. 4 but for NO<sub>3</sub>-/SO<sub>4</sub><sup>2</sup>-/OA. The results for March and October are given in Fig. S7.

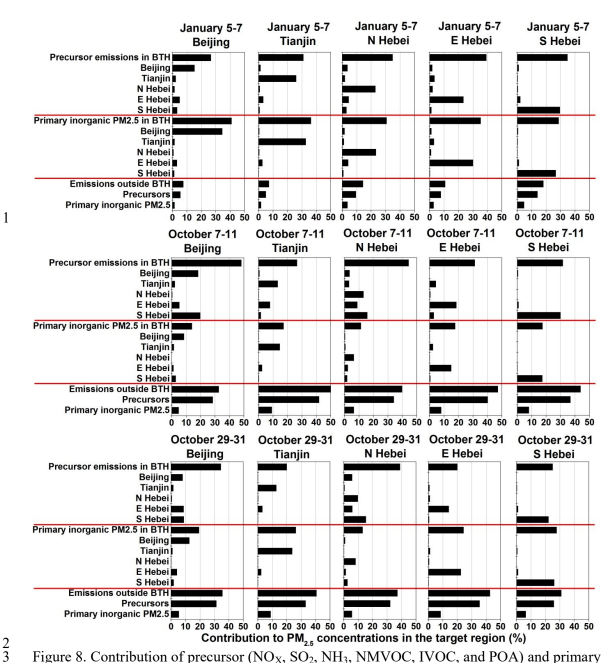


Figure 8. Contribution of precursor (NO<sub>X</sub>, SO<sub>2</sub>, NH<sub>3</sub>, NMVOC, IVOC, and POA) and primary inorganic PM<sub>2.5</sub> emissions from individual regions to PM<sub>2.5</sub> concentrations during three typical heavy-pollution episodes.

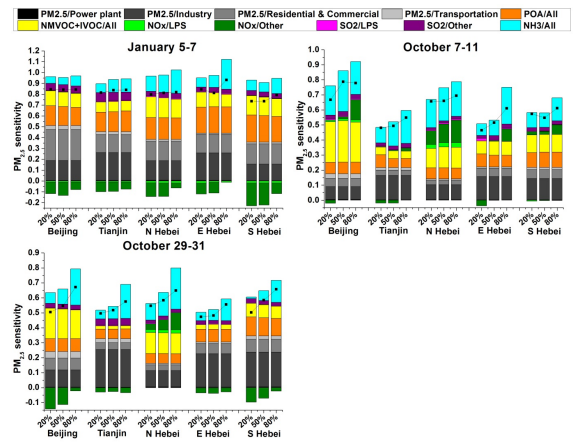


Figure 9. Sensitivity of  $PM_{2.5}$  concentrations to stepped control of individual air pollutants from individual sectors during three heavy-pollution episodes. The meanings of X-axis, Y-axis, coloured bars, and black dotted lines are the same as Fig. 4.