- ¹ Observations and modeling of air quality trends over 1990-2010
- ² across the northern hemisphere: China, the United States and

3 Europe

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9 Abstract

Trends in air quality across the northern hemisphere over a 21-year period (1990-2010) were simulated using the CMAQ multiscale chemical transport model driven by meteorology from WRF simulations and internally consistent historical emission inventories obtained from EDGAR. Thorough comparison with several ground observation networks mostly over Europe and North America was conducted to evaluate the model performance as well as the ability of CMAQ to reproduce the observed trends in air quality over the past two decades in three regions: eastern China, the continental United States and Europe.

The model successfully reproduced the observed decreasing trends in SO₂, NO₂, maxima 8h O₃, SO₄²⁻ and EC in the U.S. and Europe. However, the model fails to reproduce the decreasing trends in NO₃⁻ in the US, potentially pointing to uncertainties of NH₃ emissions. The model failed to capture the 6-year trends of SO₂ and NO₂ in CN-API from 2005-2010, but reproduced the observed pattern of O₃ trends shown in three WDCGG sites over eastern Asia. Due to the coarse spatial resolution employed in these calculations, predicted SO₂ and NO₂ concentrations are underestimated relative to all urban networks, i.e., US-AQS (NMB=-38% and

1 -48%), EU-AIRBASE (NMB=-18% and -54%) and CN-API (NMB=-36% and -68%). Conversely, at the rural network EU-EMEP SO₂ is overestimated (NMB from 4% to 150%) 2 while NO₂ is simulated well (NMB within $\pm 15\%$) in all seasons. Correlations between simulated 3 4 and observed winter time daily maxima 8-hr (DM8) O₃ are poor compared to other seasons for all networks. Better correlation between simulated and observed SO42- was found compared to 5 that for SO₂. Underestimation of summer SO_{4²⁻} in the US may be associated with the uncertainty 6 in precipitation and associated wet scavenging representation in the model. The model exhibits 7 worse performance for NO_3^{-} predictions, particularly in summer, due to high uncertainties in the 8 gas/particle partitioning of NO₃⁻ as well as seasonal variations of NH₃ emissions. There are high 9 correlations (R>0.5) between observed and simulated EC, although the model underestimates the 10 EC concentration by 65% due to the coarse grid resolution as well as uncertainties in the PM 11 speciation profile associated with EC emissions. 12

The almost linear response seen in the trajectory of modeled O₃ changes in the eastern 13 China over the past two decades, suggests that control strategies that focus on combined control 14 of NO_x and VOC emissions with a ratio of 0.46 may provide the most effective means for O_3 15 reductions for the region devoid of non-linear response potentially associated with NO_x or VOC 16 limitation resulting from alternate strategies. The response of O₃ is more sensitive to changes in 17 NO_x emissions in the eastern U.S because the relative abundance of biogenic VOC emissions 18 tends to reduce the effectiveness of VOC controls. Increasing NH₃ levels offset the relative 19 20 effectiveness of NO_x controls in reducing the relative fraction of aerosol NO_3^- formed from declining NO_x emissions in the eastern U.S., while the control effectiveness was assured by the 21 22 simultaneous control of NH₃ emission in Europe.

23 Keywords: Trends, CMAQ, modeling, air quality, sulfate, nitrate, ozone, northern hemisphere

1 1. Introduction

The last two decades have witnessed significant changes in air pollutant emissions across 2 the globe. Developed countries in North America and Europe have implemented emission 3 reduction measures which have led to a continuous improvement in air quality. Conversely, in 4 5 developing regions of the world, in Asia in particular, though control actions have been taken, their effectiveness has been overwhelmed by the sharp increase in emissions resulting from 6 increased energy demand associated with rapidly growing economies and populations. The 7 8 striking contrast in the trends in air quality between developed and developing countries has been well discussed in recent years (e.g., Richter et al, 2005). It is also believed that the observed 9 "dimming" and "brightening" trends over the past two decades is primarily related to the changes 10 of emission patterns over northern hemisphere (e.g., Wild, 2009; Gan et al, 2014). Therefore, an 11 accurate description of the decadal variations in emissions and associated aerosol burden in the 12 atmosphere is the basis of any attempts to explain the causes of decadal changes in surface solar 13 radiations and short-term climate forcing issues arising from human activities. 14

Improving air quality and protecting the health and welfare of their people is an important 15 goal for any country. Studies on historical trends in air quality can provide an indication of 16 progress in the direction as well as an assessment of future steps towards the goal. On the basis 17 of long-term records, the effectiveness of past or current control policy can be evaluated and 18 19 suitable control strategies can be designed for the future. In Europe and North America, several monitoring networks have been in operation for decades and observational records available at 20 some networks are long enough to be used in trends analysis studies (e.g., Sickles and Shadwick 21 22 (2007)). Such records are vital not only because they reflect the changes in air quality over time, 23 but also because they can be used to evaluate long-term trends in air quality arising from

1 estimated changes in historical emissions, simulated by air quality models. Colette et al (2011) analyzed the air quality trends during 1998-2007 over Europe by using observations of European 2 Monitoring and Evaluation Programme (EU-EMEP, http://www.emep.int) and the European Air 3 quality data Base (EU-AIRBASE, http://acm.eionet.europa.eu/databases/airbase/) records as 4 well as model simulations. Hogrefe et al (2009) adjusted six-year model simulations (2000-2005) 5 by using the observed PM_{2.5} species concentrations from the observations of Interagency 6 Monitoring of Protected Visual Environments (US-IMPROVE, 7 http://vista.cira.colostate.edu/improve/) and Chemical Speciation Network (CSN) sites in the 8 northeastern US. Trends in O₃ concentration and SO₄²⁻, NO₃⁻ depositions from 1988-2005 9 simulated by the same model were also compared with long term observations (Civerolo et al, 10 2010; Hogrefe et al, 2011). However, due to the large computational cost, very few studies have 11 examined in decadal trend in air pollution over large regions such as northern hemisphere. 12 Koumoutsaris and Bey (2012) evaluated the global model performance of O₃ trends simulation 13 (1991–2005) through comparison with long-term observed records from EMEP, the World Data 14 Centre for Greenhouse Gases (WDCGG, http://ds.data.jma.go.jp/gmd/wdcgg/) and the Clean Air 15 Status and Trends Network (US- CASTNET, http://epa.gov/castnet/). Long-term records of lower 16 troposphere O₃ concentrations from selected sites which are believed to represent baseline 17 conditions in Europe (Logan et al., 2012) and the U.S. (Parrish et al., 2009; 2012) were used to 18 make quantitative comparisons of simulation results from three chemistry-climate models 19 (NCAR CAM-chem, GFDL-CM3, and GISS-E2-R) (Parrish et al., 2014). To date however 20 limited attempts have been made to systematically assess long-term trends in multiple linked 21 atmospheric pollutants (oxidants, particles and acidifying substances) across regional to 22 23 hemispheric scales.

As a regional chemistry transport model (CTM), the Community Multiscale Air Quality 1 (CMAQ) modeling (version 5.0) system (Binkowski and Roselle, 2003; Byun and Schere, 2006; 2 Foley et al., 2010) has previously been successfully applied for several quality studies over North 3 America (Eder and Yu, 2006; Appel et al, 2007, 2008; Mathur et al., 2008), Europe (Matthias et 4 al., 2012; Kukkonen et al., 2012) and eastern Asia (Yamaji et al., 2006; Wang et al., 2011a; Xing 5 6 et al., 2011a). However, the need for time varying lateral boundary conditions (LBCs) which are usually derived from global CTMs simulations limits its applications in trend analysis over 7 decades. Recently, the applicability of CMAQ model has been successfully extended to 8 9 hemispheric scales (Mathur et al., 2012; 2014), so that the application of hemispheric CMAQ provides a consistent approach to generate LBCs for nested regional domains employing finer 10 resolution. 11

Changing emission patterns across the globe over the past two decades have influenced 12 background air pollution levels for different regions across the northern hemisphere. To examine 13 air quality trends in different regions over northern hemisphere, we used a multiscale chemical 14 transport model (i.e., CMAQ) driven by historical emission inventories and meteorological 15 dataset to simulate air quality from 1990-2010. The ability of the multiscale model to reproduce 16 17 observed trends over the northern hemisphere, including North America, Europe and East Asia, was assessed. A brief description of the model configuration, emission processing and 18 observations is given in section 2. The evaluation of model performance through comparison 19 with long-term observation records is presented in section 3.1. The trends in both observed and 20 simulated air quality are provided in section 3.2 and further discussed in section 4. 21

1 **2. Method**

2 **2.1 Model configuration**

Unlike the traditional regional studies with CMAQ, this study used a simulation domain extended to cover the entire northern hemisphere with a grid of 108 km×108 km resolution and 44 vertical layers of variable thickness between the surface and 50mb (Mathur et al., 2012; 2014). We selected three sub-regions, i.e., eastern China (20N-40N, 100E-125E), eastern US (28N-50N, 100W-70W) and Europe (35N-65N, 10W-30E), for further analysis and comparison with measurements. These three sub-regions are parts of the original northern hemispheric domain and no nested simulations were conducted.

The meteorological inputs for 21-year WRF simulations were derived from the 10 NCEP/NCAR Reanalysis data which has 2.5 degree spatial, and 6-hour temporal resolution. 11 NCEP ADP Operational Global Surface Observations were used for surface reanalysis which is 12 used for indirect soil moisture and temperature nudging (Pleim and Xiu, 2003; Pleim and Gilliam, 13 2009) in the Pleim-Xiu Land Surface Model (PX LSM) (Pleim and Xiu 1995; Xiu and Pleim 14 2001). The WRF configurations also used MODIS land-use types with 20 categories, RRTMg 15 shortwave and longwave radiation scheme (Iacono et al., 2008), and the ACM2 PBL model 16 (Pleim 2007a, b). WRF performance for the simulation of hourly surface temperature (T), 17 relative humidity, wind speed and direction was evaluated through comparison with observations 18 19 from NOAA's National Climatic Data Center (NCDC) Integrated Surface Data (ISD with liteformat) which provides hourly (or with 3-hour interval) meteorological observations over a long 20 historical period across the globe. The mean bias of T, wind-speed and direction over the 21 simulation domain is -0.4 K, 0.4 m s⁻¹ and -3 degree respectively, within the benchmark range 22 suggested by Emery et al. (2001) for retrospective regional-scale model applications which is \leq 23

1 $\pm 0.5 \text{ K}$, $\leq \pm 0.5 \text{ m s}^{-1}$ and $\leq \pm 10$ degree respectively.

2 2.2 Emission inventories from 1990-2010

Fig. 1 presents a flow chart of the approach to emission processing employed in creating 3 model inputs spanning the 21-year period. EDGAR (Emission Database for Global Atmospheric 4 Research, version 4.2) (European Commission, 2011) provides a consistent global emission 5 inventories for 1970-2008 for 17 anthropogenic sectors on a 0.1 °×0.1 ° resolution. In this study, 6 7 we used year specific EDGAR emission for the period 1990-2008. Estimates for 2009 and 2010 were derived from projections based on three most recent references for the United States (Xing 8 et al, 2013), Europe (EEA, 2012) and China (He, 2012). In Europe and North America, pollutant 9 emissions, SO_2 and NO_x in particular, have seen continuous reductions during 1990-2010 (refer 10 to Fig. 2). In contrast, NO_x and VOC emissions in China have continuously increased, while SO_2 11 12 increased during 1990-2006 then decreased from 2007 to 2010 due to more recent strict controls (Zhao et al., 2013; Wang et al., 2014). Emissions in other areas during 2009-2010 were kept the 13 same as the 2008 values. Additionally, since EDGARv4.2 provides only PM₁₀ emissions, PM_{2.5} 14 15 emissions were estimated by deriving the ratio of PM_{2.5} to PM₁₀ from the 2000-2005 EDGAR HTAP (Hemispheric Transport of Air Pollution, version 1) inventory (Janssens-Maenhout et al, 16 2012) which provides both PM_{10} and $PM_{2.5}$ emissions and then applying this ratio to split 17 EDGARv4.2 PM₁₀ emissions into PM_{2.5} and PM_{2.5-10}. Biogenic VOC and lightning NOx 18 emissions were obtained from GEIA (Global Emission Inventory Activity) (Guenther et al., 1995; 19 Price et al, 1997) and were kept the same for all years during 1990-2010. The 0.1° resolution 20 gridded data was spatially allocated to the CMAQ grid ensuring conservation of mass. Vertical 21 profiles for anthropogenic sectors and lightning were based on Simpson et al (2003) and Ott et al 22 23 (2010), respectively. The annual mean emissions in each sector were distributed into each hour

1 for each simulated day using the EDGAR default temporal profiles which are primarily based on

2 some western European data

3 (http://themasites.pbl.nl/tridion/en/themasites/edgar/documentation/content/Temporal-

variation.html). Emissions of PM_{2.5} and NMVOC were further speciated into AERO6 and CB05
species based on default profiles in Sparse Matrix Operator Kernel Emissions modeling system
(SMOKE, http://cmascenter.org/smoke/) which is primarily based on data for the United States.
Uncertainties are expected when region specific temporal and speciation profiles are applied to
all other counties; however this approach is reasonable given the lack of any additional
information. Further improvement and data are needed to develop more representative profiles

11 **2.3 Observed long-term trends**

12 Table 1 summarizes the dataset used in this study, which includes three networks in the United States, i.e., Air Quality System, (US-AQS, http://www.epa.gov/ttn/airs/airsaqs/), US-13 CASTNET and US-IMPROVE; two networks in Europe, i.e., EU-EMEP and EU-AIRBASE; 14 one in China (CN-API, Air Pollution Index) and one global network (WDCGG). Among these, 15 records of US-CASTNET, US-IMPROVE and EU-EMEP are specifically designed for trend 16 17 assessments since most of their sites are located in rural background areas to represent regional atmospheric pollution. Sites in US-AQS and EU-AIRBASE are typically closer to urban areas 18 and may be impacted by local pollution and features sub-grid to the model resolution, thus are 19 representative of much smaller regions. To obtain a more valid analysis, the US-AQS and EU-20 AIRBASE data were averaged over the 108 km grid cells before comparing with the model. CN-21 API is the average of observed air pollutant concentrations from urban monitoring sites in each 22 23 city and represents records in 7 Chinese cities (i.e., Beijing, Shanghai, Guangzhou, Xi'an, Wuhan,

1 Guiyang, Guilin which are located in north China plain, Yangtze-river delta, Pearl-river delta, northwest China, central China and south China respectively) where long-term observations are 2 available starting from 2005. (Jiang et al, 2004; Wang et al, 2011a). In addition, 3 selected 3 4 WDCGG sites were used for O_3 trends analysis in East Asia. Only data at sites that covered the 75% of entire 21-year period (i.e., at least 18 available years with >75% coverage for each year) 5 is considered except in the case of CN-API which was only recently set up in early 2000s and in 6 the case of US-CASTNET (for O_3 only) because most sites have no O_3 records in winter (criteria 7 set as at least 15 available years with >75% coverage from March to November for each year). 8 9 Details about the time-period covered, the number of sites selected for analysis as well as the record frequency for each network can be found in Table 1. Model results at each monitor 10 location were matched in time to the available record; thus model data was not considered during 11 periods of missing observations, in either the statistical evaluation or in the trend analysis. 12

To evaluate the model's performance, model-observed comparisons were conducted by network and pollutant. Five statistical measures: correlation coefficient (R), Mean Bias (MB), Normalized Mean Bias (NMB), Root Mean Squared Error (RMSE) and Normalized Mean Error (NME) are employed for evaluation. In consideration of the limited length of record, this study only focuses on linear trends (Colette et al, 2011). The linear least square fit method was employed and significance of trends was examined with a Student t-test at the 95% confidence level (p=0.05).

20 **3. Result**

21 **3.1 Model performance**

Table 2 summaries the statistics of model performance for gaseous species (Table 2a) and fine particles (Table 2b).

1 3.1.1 SO₂ and NO₂ concentration

Model performance characteristics for SO₂, primarily emitted from point sources, can 2 largely be attributed to artificial dilution effects over the large grid volumes employed here. As 3 expected, a hemispherical simulation with relatively coarse spatial resolution is unable to 4 accurately capture the peak values. As seen in Table 2a, SO₂ is underestimated for all urban 5 networks characterized by higher concentrations than rural network, i.e., US-AQS 6 underestimated by 38%, EU-AIRBASE by 17% and CN-API by 36%. For rural network EU-7 EMEP, SO₂ is overestimated in all seasons (4-150%). A small bias is evident for US-CASTNET 8 9 annual concentrations since the overestimation in fall is compensated by the underestimation in spring and winter. 10

Similar performance is noted for simulated NO₂. The model significantly underestimates 11 NO₂ at urban networks: US-AQS by 48%, EU-AIRBASE by 54% and CN-API by 68%. 12 However, much better performance is noted at sites in the rural network EU-EMEP with bias 13 within $\pm 15\%$ in all seasons. Though the model-observation correlation coefficients (R) are low 14 for EU-AIRBASE (0.4) and CN-API (0.08) on annual basis, the MB in EU-AIRBASE (-13.9 µg 15 m^{-3}) is comparable with previous modeling as reported by Colette et al (2011) (-6.5 to -18.1 µg 16 m^{-3}) and the magnitude of NMB in CN-API (67.5%) is comparable with Wang et al (2009) (-61.2) 17 to -81.3%) but in opposite direction. It is expected that the performance should be better when 18 simulations are conducted with finer horizontal resolution and with more accurate spatially-19 20 resolved emissions.

21 **3.1.2 O**₃ concentration

Model performance for O_3 is examined through comparisons of seasonal or annual maxima of the daily maxima 8-hr (DM8) average or 1-hour values since those are the metrics most 1 relevant to air quality standards and health assessments.

2 Correlation coefficients in EU-AIRBASE (0.4) are lower than Colette et al (2011) (0.6-0.8) because the frequency of the observed record used in this study is annual-, and therefore, the 3 4 correlation coefficients calculated here do not benefit from the fact that the model simulations generally capture the observed seasonal cycle. However, the MB (14.4 µg m⁻³) is comparable 5 with that reported in Colette et al (2011) (-4.3 to $18.5\mu g m^{-3}$). Simulations in winter (R=0.3-0.5) 6 have the worst correlation with observations for all networks compared to those in other seasons 7 (R=0.6-0.8). On the other hand, both NMB (-13.6 to 16.9%) and NME (< 25.9%) are fairly small 8 in all seasons and comparable with that reported by Zhang et al. (2009) (NMB: -10.6 to 15.9%; 9 NME: <25.4%) and Wang et al. (2009) (INMBI<37.9%). 10

11 3.1.3 SO₄²⁻, NO₃⁻ and NH₄⁺ concentration

SO₄²⁻ which is formed from the oxidation of SO₂, is the predominant inorganic aerosol 12 component. In general, SO_4^{2-} concentrations show a strong positive response to the changes in 13 SO_2 emissions (Butler and Lakens, 1991), though the SO_2 effective cloud oxidation rate can be 14 affected by NH₃ (Pandis and Seinfeld, 1989; Tsimpidi et al., 2007). As a secondary species, SO₄²⁻ 15 is widely spread over the region, unlike SO₂ which is usually more localized to source areas. As 16 seen in Table 2b, correlation coefficients for SO_4^{2-} simulation (0.5-0.9) are higher than those for 17 SO₂ (0.4-0.8). The NMBs for US-CASTNET (-8 to -45%) and US-IMPROVE (-29 to 22%) are 18 comparable with the results reported by Zhang et al. (2009), which are -23 to 22% and -8 to 16%, 19 Eder and Yu. (2006), which are -10% and -5% on annual level, and Wang et al. (2009) 20 (NMB|<55%). Significant SO₄²⁻ underestimation is noted during summer at both US-CASTNET 21 (by 45.2%) and US-IMPROVE (by 28.9%). Some studies also found similar under-prediction in 22 23 their simulations and they attributed such low biases to the uncertainty in precipitation and

overestimation of wet-scavenging. However, precipitation simulated in this study is 1 underestimated domain-wide by 4% (in summer) to 65% (in winter). Wang et al (2009) found 2 similar underestimation of precipitation from -31% to -41%, but SO42- was over-predicted 3 because higher SO₂ emissions were used. Future investigation of the low bias in predicted SO_4^{2-} 4 is still necessary. Better performance is shown at EU-EMEP, with NMB within ±30%. The 5 difference in sulfate biases between the U.S. networks and the European network might be 6 associated with the different SO₂ biases, i.e., a moderate bias (NMB=-9.4%) in US-CASTNET 7 but a relatively larger bias (NMB=+67%) in EU-EMEP. The transition rate from SO₂ to SO₄²⁻ is 8 likely underestimated in both regions, leading to the underestimation of SO_4^{2-} in the U.S. and the 9 better estimates of SO_4^{2-} in Europe. 10

Worse performance for NO₃⁻ prediction is expected because of higher uncertainties in 11 representing the gas/particle partitioning of airborne nitrate (Mathur and Dennis, 2003; Eder and 12 Yu, 2006). Especially in summer when SO_4^{2-} concentrations are higher and available NH₃ 13 preferentially react to form ammonium sulfate, leading to low ambient NO_3^{-} level. Simulated and 14 observed NO₃⁻ have the lowest correlations for both US-CASTNET and US-IMPROVE sites 15 (R=0.31 and 0.10 respectively) during summer compared those in other seasons (R=0.7). Similar 16 magnitudes of NMB (-56 to 59%) and NME (89 to 197%) at US-IMPROVE sites were reported 17 by Wang et al. (2009) and Zhang et al. (2009). The underestimation in summer and 18 overestimation in spring / winter are found relative to both CASTNET (NMB: -48% and 93/75%) 19 and IMPROVE (NMB: -41% and 107/95%) and comparable to previous CMAQ analysis of Eder 20 and Yu (2006) (|NMB| > 40%). Uncertainties in NH₃ emission particularly in the seasonal 21 temporal profile may also contribute to such bias characteristics. Slightly better performance is 22 23 noted for NO_3^- at EU-EMEP sites, with higher R (>0.6) and smaller bias (NMB: -67% to 23%)

1 for all seasons.

Performance for NH₄⁺ simulation is better than that of NO₃⁻ but slightly worse than for 2 SO_4^{2-} . The NMB for US-CASTNET is -54 to 23% which is comparable with Wang et al. (2009) 3 (INMB<50%). Similar performance statistics are shown for EU-EMEP (NMB: -15 to 68%). 4

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3.1.4 Elemental Carbon (EC) concentration

EC being a primary pollutant, its spatial distributions exhibit strong correlation to its 6 emissions. The correlation between the observed and simulated EC concentrations is high with 7 R >0.5, though the model significantly underestimates the concentrations. NMB up to -74%8 9 which is worse than previous modeling studies utilizing relatively higher spatial resolution (Zhang et al., 2009; NMB = -15.4 to 8 %; Eder and Yu, 2006; NMB = -6 %), but the magnitude 10 of NMB is comparable with Wang et al. (2009) (NMB= 101.7%) which also utilized coarse 11 spatial resolution. Some previous CMAQ modeling studies (Tesche et al., 2006; Appel et al., 12 2008) with higher spatial resolution also found the similar underestimation of EC, indicating 13 other factors besides model resolution, such as uncertainties of PM speciation profiles used to 14 estimate the EC emissions might also contribute to such low biases. 15

3.2 Trend analysis 16

Simulated trends in SO₂, NO₂, O₃, SO₄²⁻, NO₃⁻, NH₄⁺ and EC concentrations in three 17 regions (Eastern China, Eastern US and Europe) are given in Table 3. To help understand the 18 changes, trends in input emissions used in this study are also provided in Table 3 as well as 19 depicted in Fig. 2. Capability of the CMAQ model to capture the observed trends was examined 20 through comparisons with network measurements, and both simulated and observed trends are 21 quantified in Table 4 and Figures 3-9. 22

23 3.2.1 SO₂ and NO₂ trend

Simulated trends in both SO₂ and NO₂ concentrations over the northern hemisphere reflect 1 trends in SO₂ and NO_x emissions, respectively (see Fig. 2a-b, Fig. 3a and Fig. 4a), with 2 pronounced increasing trend in Asia and decreasing trend in Europe and North America. 3 Particularly, in China annual change rates of SO₂ and NO₂ concentration are about 2.7% and 4 4.1% which are comparable to their corresponding emission rates (SO₂ and NO_x) of 3.2% and 5 4.3% respectively. Annual change rates of SO₂ / NO₂ concentrations in the US (-5.7% / -1.4%) 6 and Europe (-5.1% / -1.2%) are also close to the rates of emission changes in both regions, at -7 5.4% / -1.8% and -5.4% / -1.5% respectively. 8

9 Such decreasing trends in the US and Europe are comparable with those inferred from observations at the different networks. The annual change rates of SO₂ observed from US-10 CASTNET and US-AQS are -5.0% and -5.3%, close to that simulated by the model as -6.6% and 11 -6.5%. Most of the reductions are located in the eastern US as seen in Fig.3e-f. The model was 12 unable to capture the increasing trend at two of the eastern AQS sites and also the large 13 decreasing trend at a few sites in the mid-west. It should be noted that the AQS SO₂ 14 measurements predominantly represent urban conditions, and the ability of a coarse resolution 15 model in capturing SO₂ levels and trends is influenced both by its inability to accurately 16 17 represent sub-grid variability as well as changes in local emissions. For instance, the monitor in Kansas City, MO shows sharp increase in SO₂ levels starting 2003; in contrast the grid averaged 18 SO_2 emissions in the corresponding model cell show systematic decreasing trends over the 21-19 20 year period resulting in the simulated decreasing SO_2 trend at this location. Also, as seen in the scatter plots in these panels, the pathway of such reductions from 1990 to 2010 is in good 21 agreement between observation and simulation. Stronger trends are noted in winter when SO₂ 22 concentrations are higher compared to other seasons in both observed (-0.368 μ g m⁻³ yr⁻¹) and 23

simulated trend (-0.366 µg m⁻³ yr⁻¹) at US-CASTNET (see Table 4). Annual change rates of SO₂
observed from EU-AIRBASE and EU-EMEP are -8.9% and -7.3% which are close to that
simulated by the model at -5.9% and -6.1%, with higher rates in winter when SO₂ concentration
are at their highest level. Significant reductions are found at locations in Southern UK, Benelux,
Germany, Italy, Czech Republic, Poland, Hungary and Romania.

The overall reductions in NO₂ from 1990 to 2010 are also in good agreement between the 6 observations and model simulations. Observed decreasing trends of NO₂ concentrations (and 7 annual change rate) are shown in urban networks, i.e., US-AQS and EU-AIRBASE are -0.63 µg 8 m^{-3} yr⁻¹ (-2.3%) and -0.64µg m⁻³ yr⁻¹ (-1.9%) respectively. Model simulated trends (and annual 9 change rate) at these two urban network, $-0.32 \ \mu g \ m^{-3} \ yr^{-1} (-2.2\%)$ and $-0.14 \ \mu g \ m^{-3} \ yr^{-1} (-0.9\%)$ 10 respectively, are however underestimated. The reason might be associated with the 11 underestimation of NO₂ concentrations. The model slightly overestimated the trends (annual 12 change rates as well) at the rural EU-EMEP network (-0.16 μ g m⁻³ yr⁻¹ (-2.0%) from the model, 13 compared to the observed trends of -0.13 μ g m⁻³ yr⁻¹ (-1.7%)). Such decreasing trends are more 14 pronounced over the eastern US and California as well as Southern UK, Northern France, 15 Benelux and Germany. 16

Large increases in the remotely sensed NO₂ vertical column density (VCD) over eastern China over the past decade has been noted in many studies (Richter et al., 2005; Irie et al, 2005; Akimoto et al., 2006; Zhang et al., 2007) but very limited in-situ data is available. Trends in SO₂ and NO₂ inferred from available CN-API data (for 6 years) were not significant (Table 4 and Fig. 3-4b); the model was unable to capture these trends, yielding trends more similar to those of the emissions. These discrepancies could likely arise from uncertainties in local emissions as well as the coarse spatial resolution which limits the model's ability to represent pollution distribution at finer scale which is likely captured at these monitors. Some industries were moved out from city center to rural area nearby so that the improvement of local air quality observed in city center cannot be captured by large scale simulations. However, the model results agree with the findings from studies analyzing satellite information over Asia. For example, Zhang et al. (2012) analyzed SCIAMACHY-SO₂ VCD during 2004-2009, suggesting a continuous increase in tropospheric SO₂ loading in West China, but transition from increase to decrease in 2007 in East China resulting from controls.

8 **3.2.2 O**₃ trends

9 Ozone concentrations are sensitive to the control of NO_x and VOC emissions and studies have indicated that control in NO_x emission without a simultaneous significant reduction of VOC 10 might lead to an increase of daily O₃ due to the switch from VOC-limited to NO_x-limited regime 11 (e.g., Chameides et al., 1992; Sillman, 1999). However, O₃ chemistry is likely to be at NO_x-12 limited regime during periods of heavy photochemical pollution (Trainer et al, 1993; Xing et al., 13 2011b), suggesting that NO_x controls are more effective in reducing annual maximum (rather 14 than average) of DM8 O₃. Therefore, trends in NO_x emission are more likely to have positive 15 correlation with trends in annual maximum (rather than average) of DM8 O₃. As expected, 16 17 simulated trend of annual maximum of DM8 O₃ concentration (see Fig.5a) looks quite similar to the NO_x and VOC emission trends (Fig. 2b-c). The simulated annual increasing rate of annual 18 maximum of DM8 O_3 in eastern China is 1.49%, which is associated with the increase in NO_x 19 20 and VOC emissions (by 4.3% and 2.3% per year). In contrast, due to reductions of emissions, substantial decreasing trends in annual maximum of DM8 O_3 are apparent in both the eastern US 21 and Europe, with magnitudes of -0.66% and -0.54% per year, respectively (see Table 3). 22 23 Significant increases of O_3 are also shown in northern India, west-Asia and sub-Saharan Africa 1 where both NO_x and VOC emissions have increased during this period (see Fig.2b-c).

Observed decreasing trends in annual maximum of DM8 O₃ concentrations (and annual 2 change rate) in EU- EMEP, EU-AIRBASE and US-CASTNET are -1.07 µg m⁻³ yr⁻¹ (-0.7%), -3 1.35 μ g m⁻³ yr⁻¹ (-0.8%) and -1.86 μ g m⁻³ yr⁻¹ (-1.1%) respectively. Similar trends are estimated 4 by the model simulation for both networks, i.e., $-1.31 \ \mu g \ m^{-3} \ yr^{-1}$ (-0.9%), $-2.13 \ \mu g \ m^{-3} \ yr^{-1}$ (-5 1.1%) and -0.95 μ g m⁻³ yr⁻¹ (-0.6%) (see Table 4). The failure to capture the slightly increasing 6 trends in observations in the urban network (i.e., EU-AIRBASE) might be associated with the 7 limitation by coarse spatial resolution that causes the model to fail to represent the VOC-limited 8 9 regime at these urban locations and a likely switch of O_3 chemistry from VOC- to NO_x- limited regime which usually goes along with the transition from urban to rural area (e.g., Xing et al., 10 2011b). Such decreasing trends are noted in all seasons except during winter when O_3 is at the 11 lowest level. In contrast, the most significant reduction occurred in summer when O_3 12 concentrations are at the highest. The spatial pattern of O₃ trends is quite similar to that of NO₂, 13 with more pronounced decrease in regions downwind of urban areas across the eastern US and 14 California as well as Southern UK, Northern France, Benelux and Germany. The reason for 15 increasing trends shown in both observed and model in mid-west of US might be explained by 16 the changes in local emissions (less or no controls in mid-west) as well as increasing long-range 17 transport of pollutants across the Pacific (Mathur et al., 2014). Analysis of long-term 18 observations at remote sites along the western U.S. (e.g., Jaffe and Ray, 2007; Parrish et al., 2009) 19 20 also show increasing trends in O_3 within the boundary layer attributable to inflow to the western U.S. from the Pacific. 21

Though long-term observation records of O_3 are not available in China, recent studies have suggested increasing trends similar to those found here. For instance, Xu et al (2011) suggested significant increasing trends in tropospheric ozone residual over the North China Plain. Ding et al (2008) suggest that O₃ in the lower troposphere over Beijing had a strong positive trend (2% per year) during the period 1995 to 2005. Ozonesonde measurements analyzed by Wang et al (2012) suggests a clear positive trend in the maximum summer ozone concentration (3.4% per year) over the Beijing area during 2002-2010. In this study, the trend in summer maximum of DM8 ozone concentration in Beijing during 1990 to 2010 is estimated to be 2% per year, which is comparable to that inferred from observations in these two recent studies.

Observation records at three sites in WDCGG network were used to investigate trends in 8 9 O₃ distribution in eastern Asia. One of these sites, Minamitorishima (noted as S1, lat: 24.28N, lon: 153.98E), is located far from land and can be considered to be a representative of clean 10 conditions, while two sites located on Honshu island, i.e., Tsukuba (noted as S2, lat: 36.05, lon: 11 140.13) which is to the northwest of Tokyo and closest to urban regions, and Ryori (noted as S3, 12 lat: 39.03, lon: 141.82) which is in the north and representative of rural conditions. The model 13 generally captured the observed pattern of O_3 trends at each site. For the clean site (S1), no 14 significant trends are inferred either in the observed or the simulated maximum of DM8 O₃. 15 However, for the urban site (S2), significant reduction, particularly during summer, is noted in 16 17 the observed values and is reflective of emission reductions in Japan during past two decades (e.g., Wakamatsu et al., 2013). In contrast, increasing trends are inferred at the rural site (S3) in 18 all seasons expect fall, presumably, representing transport from upwind locations in East Asia. 19 20 The model produces similar magnitude (though smaller significance) of the decreasing/increasing trends at S2/S3. The contrasting trends at sites S2 and S3 likely result from 21 different controls in local emissions as well as transboundary transport. 22

23 **3.2.3** SO₄²⁻, NO₃⁻ and NH₄⁺ trends

Simulated SO4²⁻ shows a pronounced increasing trend in eastern China (2.8% per year) and
decrease in the US (-3.2% per year) and EUROPE (-3.7% per year) which is consistent with,
though slightly smaller in magnitude, with trends in SO₂ emissions in these regions (see Table 3 and Fig. 6).

Simulated SO₄²⁻ trends are in a good agreement with observed trends inferred from all three 5 networks. Simulated trends in SO₄²⁻ concentrations (and annual change rate) at US-CASTNET, 6 US-IMPROVE and EU-EMEP are -0.09 μ g m⁻³ yr⁻¹(-3.5%), -0.03 μ g m⁻³ yr⁻¹(-2.1%) and -0.09 7 μ g m⁻³ yr⁻¹ (-3.6%), which is comparable with the observed trends of -0.10 μ g m⁻³ yr⁻¹ (-2.9%), -8 0.03 μ g m⁻³ yr⁻¹ (-2.4%) and -0.10 μ g m⁻³ yr⁻¹ (-4.1%), respectively. More significant trends are 9 noted in summer compared to other seasons because of relatively higher summer time SO42-10 concentrations. Average trends at US-CASTNET are more significant than those at IMPROVE 11 12 because majority of CASTNET sites are located in the eastern US which witnessed stronger reductions in SO₂ emissions. In Europe, most SO_4^{2-} reductions are found in central to eastern 13 Europe, i.e., Germany, Czech, Poland, Hungary, Benelux, Italy, and Romania. 14

 NH_3 emission plays an important role in NO_3^- formation (Mathur and Dennis, 2003; Wang 15 et al., 2011b). Growth in NH₃ emission or reduction in SO₂ emission (consequently more free 16 NH_3 due to less association with SO_4^{2-}) without simultaneous reduction in NO_x emission can 17 enhance NO₃⁻ concentration especially under NH₃ poor conditions (Pinder et al., 2008a; 18 Blanchard et al., 2007). As illustrated in Fig. 7, growth in both NO_x and NH_3 emissions results in 19 the increasing trend in airborne NO_3^- in China (5.4% per year), while reductions in emissions of 20 both results in the decreasing trend in Europe (-1.8% per year). In contrast, over the past two 21 decades in the US, a reduction in SO₂ and NO_x accompanied with a growth in NH₃ emission 22 23 results in different trends across different seasons. The model fails to reproduce the decreasing

1 trend in NO₃⁻ at both US-CASTNET and US-IMPROVE in spring, summer and fall though the significance of the trend is small. However, both simulated and observed NO_3^- show an 2 increasing trend in winter values when NO_3^{-1} is at the highest level. Similar observed increasing 3 trend is noted during winter at the EU-EMEP monitors, which is not captured by the model. The 4 decreasing trend at the EU-EMEP locations during other seasons is however captured by the 5 6 model. Successful reproduction of NO₃⁻ trends depends on an accurate baseline emission as well as an accurate representation of changes in historical NH₃ emission. Unfortunately, both current 7 NH₃ emission and their historical trends over the globe still suffer from large uncertainties (e.g., 8 9 Heald et al, 2012) and likely contribute to the significant bias in the simulated NO_3^- trend.

10 NH_4^+ is simulated based on the thermodynamic equilibrium between the NO_x - SO_x - NH_x 11 species. It shows a similar increasing trend in China (3.4%) and a decreasing trend in the US (-12 0.7%) and Europe (-2.9%), as illustrated in Fig. 8. NH_4^+ simulation suffers the same 13 uncertainties as NO_3^- which leads to difficulties in reproducing the trend in observations (see 14 Table 4).

15 **3.2.4 Elemental Carbon (EC) trends**

Growth of human activities such as biomass burning and open fires results in the simulated increasing trends in EC levels in China (1.0%; see Table 3), India and sub-Saharan Africa (see Fig. 9). In contrast, continuous controls have led to a decreasing trend in EC concentrations in the US (-3.4%) and Europe (-2.5%). The observed trend in EC at US-IMPROVE, i.e., -0.006 μ g m⁻³ yr⁻¹ (-2.6%) is well reproduced by the model, i.e., -0.003 μ g m⁻³ yr⁻¹ (-3.3%). Both observations and the model suggest higher magnitudes of trends during fall and winter, and are likely associated with higher ambient levels during these seasons.

23 Decreasing trend of EC in Europe has also been observed in other studies (Järvi et al.,

1 2008). The model estimates a consistent decreasing EC trend in the Canadian Arctic (see Fig. 9)
2 which is mainly impacted by emissions from Europe and Russia during winter and spring as
3 demonstrated by Sharma et al (2004) who analyzed in-situ ground-level observations of aerosol
4 black carbon between 1989 and 2002. The increasing trend of EC in southern Asia is
5 corroborated by the evidence found from the Nam Co Lake (located in the central Tibetan
6 Plateau) sediments indicating a recent rise in BC deposition flux (Cong et al., 2013).

7 4. Discussion

8 4.1 O₃ chemistry

9 As discussed in section 3.2.2, the response of O_3 concentration depends on changes in NO_x 10 and VOC emissions, and the non-linear chemistry associated with the subsequent VOC- or NO_{x-1} limited environment. The response of O₃ to changing levels of NO_x and VOC have previously 11 been examined through a variety of methods ranging from isopleths created from chemistry box-12 model calculations to detailed spatially varying response surfaces developed from output of 13 14 hundreds of simulations with detailed air pollution modeling systems (e.g., Xing et al., 2011b). Exploration of the changes in O₃ levels in response to historical (and geographically varying) 15 changes in NO_x and VOC emissions, as captured by the multi-decadal simulations presented here, 16 17 provide a unique opportunity to develop insights into factors controlling changes in O₃ production and distributions. 18

Fig. 10 attempts to summarize the changes in NO_x and VOC emissions as well as the surface O_3 response during the 1990-2010 period for the three regions; the figures in the left panel illustrate the changes in emissions relative to the 1990 values and the figures in the right panel show the corresponding percentage change in both the maximum and the average of the DM8 O_3 for each year. As can be noted, the relative changes in NO_x and VOC emissions vary

significantly over different time-period for different regions. Based on the emission estimates, 1 simultaneous growth of VOC and NO_x emissions is noted in China with a ratio of 0.46 (i.e., x%2 NO_x growth along with 0.46x% VOC growth on a basis of 1990 emission level). The modeled 3 increases in both maximum and average of DM8 O₃ values in China during this period are 4 significant. The almost linear response seen in the trajectory of modeled O₃ changes in the region 5 6 over the past two decades, suggests that control strategies that focus on combined control of NO_x and VOC emissions with a ratio of 0.46 may provide the most effective means for O_3 reductions 7 for the region devoid of non-linear response potentially associated with NO_x or VOC limitation 8 9 resulting from alternate strategies. The ratio suggested is less than 1 indicating greater sensitivity of ozone to NO_x emissions than VOC emissions. It's also obvious to see that the rate of O_3 10 increase was much smaller during 1995-2002 which was the period when VOC emission growth 11 12 was much greater than that of NO_x emissions in China.

In contrast, trends in emissions over the eastern US indicate significant reduction in VOC 13 emissions compared to NO_x prior to 2000. NO_x emission increased slightly during 1996-2000, 14 and then decreased significantly resulting from regional control measures. Change of O_3 during 15 the first decade (1990-2000) when VOC controls were dominant (reduction ratio of VOC and 16 NO_x is -42% and -4% respectively) is smaller (-2%) than that in the subsequent decade (2000-17 2010) when NO_x controls were dominant (reduction ratio of VOC and NO_x is -13% and -33%, 18 respectively) leading to an estimated reduction of -11% in ambient O₃. Additionally, model 19 simulations also show an increase in O₃ during 1997-1999 when NO_x emissions were estimated 20 to increase. Thus, the response of O_3 is more sensitive to changes in NO_x emissions in the eastern 21 US. The relative abundance of biogenic VOC emissions that tend to reduce the effectiveness of 22 23 VOC controls, contributes to this differing response.

1 In Europe, simultaneous control of NO_x and VOC with a ratio of 1.8 during 1990-2010 result in systematic reduction in ambient O_3 levels. Interestingly, the reductions in the annual 2 maximum of the regionally-averaged DM8 O_3 are much greater than those of the corresponding 3 annual mean DM8 O₃, indicating the impact of emission reductions in the region on reducing 4 peak O₃ during regional pollution episodes. During the period 2000-2007 when solely VOC 5 emissions reduced (-10%), no significant reduction in either annual maximum or average of 6 DM8 O₃ occurred. Reductions in NO_x (-10%) with VOC (-5%) emissions in the subsequent 2007 7 to 2010 period lead to reductions in both maximum and average of DM8 O₃. 8

9 4.2 PM chemistry

The nonlinear response of NO_3^- concentration to SO_2 , NO_x and NH_3 emissions are well 10 documented (e.g., Mathur and Dennis, 2003; Tsimpidi et al., 2007; Makar et al., 2009). Fig. 11 11 12 attempts to summarize the changes in emissions and factors driving the NO_x -SO_x-NH_x system and its influence on changing inorganic particulate matter composition for the three regions. 13 Contrasting trends in emissions over the past two decades in the three regions are apparent: while 14 China and many growing regions of Asia have witnessed significant increases in emissions of 15 NO_x, SO₂, and NH₃, significant reductions in emissions of all these species have occurred in 16 Europe. In contrast in the eastern U.S., while combustion related emissions of NO_x and SO₂ have 17 declined, growth in agricultural animal husbandry have resulted in significant increases in NH₃ 18 emissions. To examine the impact of the varying emissions patterns on inorganic particulate 19 matter formation and composition in these regions, we examined trends in two metrics relative to 20 their 1990 values: (i) the degree of sulfate neutralization, an estimate of the neutralization of 21 sulfate by ammonium (Pinder et al. (2008b); $DSN=([NH_4^+] - [NO_3^-]) / [SO_4^{2-}])$, and (ii) a new 22 23 metric, the "nitration ratio (NR)" (i.e., NO_3^- concentration divided by NO_x emission) to represent the relative amount of oxidized-N emissions that is eventually transformed to aerosol NO_3^- , changes in the ratio could thus be viewed as an indicator of the relative effectiveness of NO_x controls for given conditions. Fig. 11 presents the response of PM chemistry to the changes in emissions as indicated by the trends in these metric during the period 1990-2010.

In eastern China, simultaneous growth of NH₃ emission with SO₂/NO_x plays a very 5 important role in the increases of SO₄²⁻ and NO₃⁻ concentrations (Wang et al., 2011b). During the 6 period 1993-2002 the rate of increase in NH₃ emissions is greater than that of $NO_x+2\times SO_2$ 7 emissions (representing the amount of NH₃ needed for complete neutralization) with a ratio of 8 9 1.1 (i.e., x% (NO_x+2SO₂) growth along with 1.1x% NH₃ growth on a basis of 1990 emission level). In these NH₃-rich conditions, both DSN and NR consequently exhibit an increasing trend, 10 suggesting that sufficient NH₃ was available to neutralize the available and increasing aerosol 11 SO_4^{2-} and also enable formation of particulate NO_3^{-} . The increasing trend in NR for this region 12 also indicate that the simultaneous growth in emissions of both reduced and oxidized nitrogen 13 results in greater fraction of NO_x being eventually transformed to particulate NO₃⁻. After 2002, 14 both DSN and NR decline when the growth of $NO_x+2\times SO_2$ emissions is faster than that of NH_3 15 (ratio of 0.9), resulting in the decline of the DSN and NR and eventually back to the 1990-levels. 16 17 In contrast, in the eastern US, both DSN and NR exhibit a steady-increase during the entire 21 year period, suggesting progressively NH₃-rich conditions stemming from both the increased 18 NH₃ emissions as well as more free NH₃ being available due to reduced SO₄²⁻ levels associated 19 20 with declining SO_2 emissions. Steadily increasing trends in NR values also suggest that increasing NH₃ levels offset the relative effectiveness of NO_x controls in reducing the relative 21 fraction of aerosol NO₃⁻ formed from declining NO_x emissions. 22

23 Interestingly, in Europe simultaneous control of NH_3 along with NO_x and SO_2 emissions

yields an emission change ratio of 0.6 (i.e., x% (NO_x+2SO₂) reduction along with 0.6x%
reduction of NH₃ on a basis of 1990 emission level). Though a slight increase of DSN is
simulated during 1992-2003 resulting from faster growth of NO_x and SO₂ compared to NH₃,
there is no discernable trend in the estimated NR suggesting comparatively greater control
effectiveness in this region compared to the other two, due to the simultaneous control of NH₃
with combustion related emissions of NO_x and SO₂.

7 5. Conclusion

Trends in air quality across the northern hemisphere from 1990 to 2010 have been 8 simulated by the WRF-CMAQ model driven with a representation of historical emission 9 10 inventories derived from the EDGAR. Thorough comparison with several surface observation 11 networks mostly in Europe and North America has been conducted. Significant contrasting changes in emissions have occurred across the northern hemisphere over the past two decades 12 with reductions in North America and Western Europe resulting from control measures on 13 14 combustion related sources and increases across large parts of Asia associated with economic and population growth. Model calculations show associated contrasting trends in air pollution across 15 the northern hemisphere emphasizing the changing tropospheric composition of trace pollutants 16 17 as well as the potentially changing background pollution levels in different regions resulting from changes in the amounts of long-range transported pollution. The model is generally able to 18 capture the observed trends in air pollution and performance statistics are comparable with 19 results from other studies in regions across the northern hemisphere. However, the model 20 21 estimates still suffer from uncertainties in emissions (in regards to temporal variation and speciation), coarse spatial resolution, and subsequent impacts on representation of non-linear 22 atmospheric chemistry. The lightening NO_x emissions used in this studies (Price et al, 1997) are 23

likely overestimated by 0.5 to 5 times compared to a more recent study (Schumann and 1 Huntrieser et al., 2007) and may contribute to some extent to the overestimation of NO_x , O_3 and 2 nitrate concentrations. The trend of biogenic emissions, which hasn't been considered in this 3 study, might also impact the analysis. The lack of long-term observations in Asia, particularly 4 over China and India, limits a robust model performance evaluation as well as O₃ and PM 5 chemistry assessment in these polluted areas. To future explore the limitation of coarse spatial 6 resolution, we are currently conducting a study with a finer-scale simulation over the CONUS 7 domain for the same simulated period as from 1990 to 2010. A detailed description and 8 9 comparison will be provided in a separate paper (Gan et al., in preparation).

Model simulated air quality trends over the past two decades largely agree with those 10 derived from observations. Significant reduction in ambient levels of most pollutants is seen in 11 the U.S. and Europe resulting from emission controls implemented during 1990-2010, while 12 levels of all pollutants in China show pronounced increasing trends during the same period. 13 Examining the simulated and observed historical trends in atmospheric chemistry can help guide 14 development of future air pollution abatement strategies. Model calculations over the 1990-2010 15 period suggest that in the relative amounts of VOC and NO_x emission controls in different 16 regions across the northern hemisphere (east U.S., Europe, and China), have led to significantly 17 different trends in tropospheric O_3 in these regions. In particular, steady increase in NO_x and 18 VOC emissions (with a ratio of 0.46 relative to 1990 emissions) in China have resulted in a near-19 20 linear increase in surface O_3 concentrations in the region, suggesting that possible control strategies that maintain this relative ratio could potentially be most effective in avoiding non-21 linear response resulting from VOC-limitation of alternate approaches. Differences in the 22 23 historical changes in the relative amounts of NH₃, NO_x, and SO₂ emissions in these regions also

1 impact the trends in inorganic particulate matter amounts and composition in these regions. In particular, the amount of particulate nitrate formed per unit of NO_x emissions is influenced by 2 changing NH₃ emissions and could be important in assessing the relative effectiveness of 3 different control strategies. Simultaneous growth of NH_3 emission along with those of NO_x and 4 SO₂ in China over the past 2 decades has resulted in the increasing particulate nitrate formation 5 6 trends in the region. In contrast, in the eastern U.S. the relative fraction of NO_x converted to particulate nitrate exhibits a steady increase over the past two decades suggesting an offset in the 7 relative effectiveness of control measures on particulate nitrate levels in the region. Simultaneous 8 9 reductions in NH_3 emissions along with those of NO_x , and SO_2 in west Europe over the past two decades resulted in no significant trend in nitration ratio, suggesting effectiveness of the overall 10 measures in terms of particulate nitrate levels in the region. 11

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1 References

3	Akimoto, H., Ohara, T., Kurokawa, J. I., and Horii, N.: Verification of energy consumption in
4	China during 1996–2003 by using satellite observational data. Atmospheric Environment,
5	40(40), 7663-7667. 2006.
6	Appel, K. W., Gilliland, A. B., Sarwar, G., and Gilliam, R. C.: Evaluation of the Community
7	Multiscale Air Quality (CMAQ) model version 4.5: sensitivities impacting model
8	performance: part I-ozone. Atmospheric Environment, 41(40), 9603-9615, 2007.
9	Appel, K. W., Bhave, P. V., Gilliland, A. B., Sarwar, G., and Roselle, S. J.: Evaluation of the
10	community multiscale air quality (CMAQ) model version 4.5: sensitivities impacting
11	model performance; Part II-Particulate matter. Atmospheric Environment, 42(24), 6057-
12	6066, 2008.
13	Binkowski, F. S. and Roselle, S.J.: Community Multiscale Air Quality (CMAQ) model aerosol
14	component, I: Model description, Journal of Geophysical Research, 108, 4183,
15	doi:10.1029/2001JD001409, 2003.
16	Blanchard, C. L., Tanenbaum, S., Hidy, G. M.: Effects of Sulfur Dioxide and Oxides of Nitrogen
17	Emission Reductions on Fine Particulate Matter Mass Concentrations: Regional
18	Comparisons. J. Air Waste Manage. Assoc., 57, 1337–1350, 2007.
19	Butler, T. J., and Lakens, G. E.: The impact of changing regional emissions on precipitation
20	chemistry in the eastern United States. Atmospheric Environment. Part A. General Topics,
21	25(2), 305-315, 1991
22	Byun, D. and Schere, K. L.: Review of the governing equations, computational algorithms, and
23	other components of the Models-3 Community Multiscale Air Quality (CMAQ) modeling

system, Appl. Mech. Rev., 59, 51-77, 2006.

2	Chameides, W. L., Fehsenfeld, F., Rodgers, M. O., Cardelino, C., Martinez, J., Parrish, D.,
3	Lonneman, W., Lawson, D.R., Rasmussen, R.A., Zimmerman, P., Greenberg, J., Mlddleton,
4	P. and Wang, T.: Ozone precursor relationships in the ambient atmosphere. Journal of
5	Geophysical Research: Atmospheres (1984–2012), 97(D5), 6037-6055, 1992.
6	Civerolo, K., Hogrefe, C., Zalewsky, E., Hao, W., Sistla, G., Lynn, B., Rosenzweig, C., Kinney, P.
7	L.: Evaluation of an 18-year CMAQ simulation: Seasonal variations and long-term
8	temporal changes in sulfate and nitrate. Atmos. Environ., 44, 4745–3752, 2010.
9	Colette, A., Granier, C., Hodnebrog, Ø., Jakobs, H., Maurizi, A., Nyiri, A., Bessagnet, B.,
10	D'Angiola, A., D'Isidoro, M., Gauss, M., Meleux, F., Memmesheimer, M., Mieville, A.,
11	Rouïl, L., Russo, F., Solberg, S., Stordal, F., and Tampieri, F.: Air quality trends in Europe
12	over the past decade: a first multi-model assessment, Atmos. Chem. Phys., 11, 11657-
13	11678, doi:10.5194/acp-11-11657-2011, 2011.
14	Cong, Z., Kang, S., Gao, S., Zhang, Y., Li, Q., and Kawamura, K.: Historical trends of
15	atmospheric black carbon on Tibetan Plateau as reconstructed from a 150-year lake
16	sediment record. Environmental science & technology, 47(6), 2579-2586, 2013.
17	Ding, A. J., Wang, T., Thouret, V., Cammas, JP., and Nédélec, P.: Tropospheric ozone
18	climatology over Beijing: analysis of aircraft data from the MOZAIC program, Atmos.
19	Chem. Phys., 8, 1-13, doi:10.5194/acp-8-1-2008, 2008.
20	Eder, B. and Yu, S.: A performance evaluation of the 2004 release of Models-3 CMAQ,
21	Atmospheric Environment, 40, 4811-4824, 2006.
22	EEA (European Environment Ageney); European Union emission inventory report 1000-2010
	EEA (European Environment Agency). European Onion emission inventory report 1990–2010

1	Technical report, 30 Jul, Copenhagen, Denmark, doi:10.2800/5219, 2012.
2	European Commission: Joint Research Centre (JRC)/Netherlands Environmental Assessment
3	Agency (PBL). Emission Database for Global Atmospheric Research (EDGAR), release
4	version 4.2., available at: http://edgar.jrc.ec.europa.eu (last access: 25 September 2014),
5	2011.
6	Foley, K. M., Roselle, S. J., Appel, K. W., Bhave, P. V., Pleim, J. E., Otte, T. L., Mathur, R.,
7	Sarwar, G., Young, J. O., Gilliam, R. C., Nolte, C. G., Kelly, J. T., Gilliland, A. B., and Bash,
8	J. O.: Incremental testing of the Community Multiscale Air Quality (CMAQ) modeling
9	system version 4.7, Geosci. Model Dev., 3, 205–226, doi:10.5194/gmd-3-205-2010, 2010.
10	Gan, CM., Pleim, J., Mathur, R., Hogrefe, C., Long, C. N., Xing, J., Roselle, S., and Wei, C.:
11	Assessment of the effect of air pollution controls on trends in shortwave radiation over the
12	United States from 1995 through 2010 from multiple observation networks, Atmos. Chem.
13	Phys., 14, 1701–1715, 2014.
14	Gan, CM., Pleim, J., Mathur, R., Hogrefe, C., Long, C.N., Xing, J., Wong, D., Gilliam, R.,
15	Roselle, S.J. and Wei, C.: Assessment of long-term simulations with various observations
16	for better understanding of aerosol effects on radiation "brightening" in the United States,
17	in preparation.
18	Guenther, A., Hewitt, C. N., Erickson, D., Fall, R., Geron, C., Graedel, T., Harley, P., Klinger, L.,
19	Lerdau, M., McKay, W. A., Pierce, T., Scholes, B., Steinbrecher, R., Tallamraju, R., Taylor,
20	J., and Zimmerman, P.: A Global Model of Natural Volatile Organic Compound Emissions,
21	J. Geophys. Res., 100, 8873–8892, 1995.
22	He, K. B.: Multi-resolution Emission Inventory for China (MEIC): model framework and 1990-
23	2010 anthropogenic emissions, International Global Atmospheric Chemistry Conference,

	1	
	L	

17–21 September, Beijing, China, S1-I-2, 2012.

2	Heald, C. L., J. L. Collett Jr., Lee, T., Benedict, K. B., Schwandner, F. M., Li, Y., Clarisse, L.,
3	Hurtmans, D. R., Van Damme, M., Clerbaux, C., Coheur, PF., Philip, S., Martin, R. V.,
4	and Pye, H. O. T.: Atmospheric ammonia and particulate inorganic nitrogen over the
5	United States, Atmos. Chem. Phys., 12, 10295-10312, doi:10.5194/acp-12-10295-2012,
6	2012.
7	Hogrefe, C., Lynn, B., Goldberg, R., Rosenzweig, C., Zalewsky, E., Hao, W., Doraiswamy, P.,
8	Civerolo, K., Ku, JY., Sistla, G., and Kinney, P.L.: A combined model-observation
9	approach to estimate historic gridded fields of PM2.5 mass and species concentrations.
10	Atmospheric Environment 43, 2561-2570, 2009.
11	Hogrefe, C., Hao, W., Zalewsky, E. E., Ku, JY., Lynn, B., Rosenzweig, C., Schultz, M. G., Rast,
12	S., Newchurch, M. J., Wang, L., Kinney, P. L., and Sistla, G.: An analysis of long-term
13	regional-scale ozone simulations over the Northeastern United States: variability and trends,
14	Atmos. Chem. Phys., 11, 567-582, doi:10.5194/acp-11-567-2011, 2011.
15	Iacono, M.J., Delamere, J.S., Mlawer, E.J., Shephard, M.W., Clough, S.A., and Collins, W.D.:
16	Radiative forcing by long-lived greenhouse gases: Calculations with the AER radiative
17	transfer models, J. Geophys. Res., 113, D13103, doi:10.1029/2008JD009944, 2008.
18	Irie, H., Sudo, K., Akimoto, H., Richter, A., Burrows, J.P., Wagner, T., Wenig, M., Beirle, S.,
19	Kondo, Y., Sinyakov, V.P. and Goutail, F.: Evaluation of long-term tropospheric NO2 data
20	obtained by GOME over East Asia in 1996-2002. Geophysical Research Letter 32(11),
21	L11810, doi: 10.1029/2005GL022770, 2005.

Jaffe, D., and Ray, J.: Increase in surface ozone at rural sites in the western US. Atmospheric
 Environment, 41(26), 5452-5463, 2007.

1	Janssens-Maenhout, G., Dentener, F., Van Aardenne, J., Monni, S., Pagliari, V., Orlandini, L.,
2	Klimont, Z., Kurokawa, J., Akimoto, H., Ohara, T., Wankmueller, R., Battye, B., Grano, D.,
3	Zuber, A., and Keating, T.: EDGAR-HTAP: a Harmonized Gridded Air Pollution Emission
4	Dataset Based on National Inventories, European Commission Publications Office, Ispra,
5	Italy, EUR report No EUR 25229, 2012.
6	Järvi, L., Junninen, H., Karppinen, A., Hillamo, R., Virkkula, A., Mäkelä, T., Pakkanen, T., and
7	Kulmala, M.: Temporal variations in black carbon concentrations with different time scales
8	in Helsinki during 1996–2005, Atmos. Chem. Phys., 8, 1017-1027, doi:10.5194/acp-8-
9	1017-2008, 2008.
10	Jiang, D.H., Zhang, Y., Hu, X., Zeng, Y., Tan, J.G., Shao, D.M.: Progress in developing an ANN
11	model for air pollution index forecasting, Atmospheric Environment, 28, pp. 7055-7064,
12	2004.
13	Koumoutsaris, S. and Bey, I.: Can a global model reproduce observed trends in summertime
14	surface ozone levels? Atmos. Chem. Phys., 12, 6983–6998, doi:10.5194/acp-12-6983-2012,
15	2012.
16	Kukkonen, J., Olsson, T., Schultz, D. M., Baklanov, A., Klein, T., Miranda, A. I., Monteiro, A.,
17	Hirtl, M., Tarvainen, V., Boy, M., Peuch, VH., Poupkou, A., Kioutsioukis, I., Finardi, S.,
18	Sofiev, M., Sokhi, R., Lehtinen, K. E. J., Karatzas, K., San José, R., Astitha, M., Kallos, G.,
19	Schaap, M., Reimer, E., Jakobs, H., and Eben, K.: A review of operational, regional-scale,
20	chemical weather forecasting models in Europe, Atmos. Chem. Phys., 12, 1-87,
21	doi:10.5194/acp-12-1-2012, 2012.
22	Logan, J. A., Staehelin, J., Megretskaia, I. A., Cammas, JP., Thouret, V., Claude, H., De Backer,
23	H., Steinbacher, M., Scheel, H. E., St"ubi, R., Fr"ohlich, M., and Derwent, R.: Changes in

1	ozone over Europe: analysis of ozone measurements from sondes, regular aircraft
2	(MOZAIC) and alpine surface sites, J. Geophys. Res., 117, D09301,
3	doi:10.1029/2011JD016952, 2012.
4	Luo, C., Wang, Y.H., Mueller, S., and Knipping, E.: Diagnosis of an underestimation of
5	summertime sulfate using the Community Multiscale Air Quality model. Atmos Environ,
6	45, 5119–30, 2011.
7	Makar, P. A., Moran, M. D., Zheng, Q., Cousineau, S., Sassi, M., Duhamel, A., Besner, M.,
8	Davignon, D., Crevier, LP., and Bouchet, V. S.: Modelling the impacts of ammonia
9	emissions reductions on North American air quality, Atmos. Chem. Phys., 9, 7183-7212,
10	doi:10.5194/acp-9-7183-2009, 2009.
11	Mathur, R., and Dennis, R. L.: Seasonal and annual modeling of reduced nitrogen compounds
12	over the eastern United States: Emissions, ambient levels, and deposition amounts. Journal
13	of Geophysical Research: Atmospheres (1984–2012), 108, 4481,
14	doi:10.1029/2002JD002794, 2003.
15	Mathur, R., Yu, S., Kang, D., and Schere, K. L.: Assessment of the wintertime performance of
16	developmental particulate matter forecasts with the Eta - Community Multiscale Air
17	Quality modeling system. Journal of Geophysical Research: Atmospheres (1984 - 2012),
18	113, D02303, doi:10.1029/2007JD008580, 2008.
19	Mathur, R., Gilliam, R., Bullock, O.R., Roselle, S., Pleim, J., Wong, D., Binkowski, F., and
20	Streets, D.: Extending the applicability of the community multiscale air quality model to
21	hemispheric scales: motivation, challenges, and progress. In: Steyn DG, Trini S (eds) Air
22	pollution modeling and its applications, XXI. Springer, Dordrecht, pp 175–179, 2012.
23	Mathur, R., Roselle, S., Young, J. and Kang, D.: Representing the Effects of Long-Range 33

1	Transport and Lateral Boundary Conditions in Regional Air Pollution Models, Air
2	Pollution Modeling and its Application XXII NATO Science for Peace and Security Series
3	C: Environmental Security, Springer, Heidelburg, Germany, Chapter 51, 303-308, 2014.
4	Matthias, V., Aulinger, A., Bieser, J., Cuesta, J., Geyer, B., Langmann, B., Serikov, I., Mattis, I.,
5	Minikin, A., Mona, L., Quante, M., Schumann, U., and Weinzierl, B.: The ash dispersion
6	over Europe during the Eyjafjallajökull eruption-comparison of CMAQ simulations to
7	remote sensing and air-borne in-situ observations. Atmos. Environ. 48, 184–194, 2012.
8	Ott, L. E., Pickering, K. E., Stenchikov, G. L., Allen, D. J., De-Caria, A. J., Ridley, B., Lin, RF.,
9	Lang, S., and Tao, WK.: Production of lightning NO(x) and its vertical distribution
10	calculated from three-dimensional cloud-scale chemical transport model simulations, J.
11	Geophys. ResAtmos., 115, D04301, doi:10.1029/2009jd011880, 2010.
12	Parrish, D. D., Millet, D. B., and Goldstein, A. H.: Increasing ozone in marine boundary layer
13	inflow at the west coasts of North America and Europe, Atmos. Chem. Phys., 9, 1303-1323,
14	doi:10.5194/acp-9-1303-2009, 2009.
15	Parrish, D. D., Law, K. S., Staehelin, J., Derwent, R., Cooper, O. R., Tanimoto, H., Volz-Thomas,
16	A., Gilge, S., Scheel, HE., Steinbacher, M., and Chan, E.: Long-term changes in lower
17	tropospheric baseline ozone concentrations at northern mid-latitudes, Atmos. Chem. Phys.,
18	12, 11485-11504, doi:10.5194/acp-12-11485-2012, 2012.
19	Parrish, D. D., Lamarque, J. F., Naik, V., Horowitz, L., Shindell, D. T., Staehelin, J., Derwent, R.,
20	Cooper, O.R., Tanimoto, H., Volz-Thomas, A., Gilge, S., Scheel, HE., Steinbacher, M.,
21	and Fröhlich, M.: Long - term changes in lower tropospheric baseline ozone concentrations:
22	Comparing chemistry - climate models and observations at northern midlatitudes. Journal
23	of Geophysical Research: Atmospheres, 119(9), 5719-5736, 2014. 34

1	Price, C., Penner, J., and Prather, M.: NOx from lightning 1: Global distribution based on
2	lightning physics, J. Geophys. Res., 102(D5), 5929-5941, 1997.
3	Pleim, J. E. and Xiu, A.: Development and testing of a surface flux and planetary boundary layer
4	model for application in mesoscale models. J. Appl. Meteor., 34, 16–32, 1995.
5	Pleim, J. E. and Xiu, A.: Development of a land surface model. Part II: Data Assimilation. J.
6	Appl. Meteor., 42, 1811–1822, 2003.
7	Pleim, J. E.: A Combined Local and Nonlocal Closure Model for the Atmospheric Boundary
8	Layer. Part I: Model Description and Testing, J. Appl. Meteorol. Climatol., 46, 1383–1395,
9	doi:10.1175/JAM2539.1, 2007a.
10	Pleim, J. E.: A Combined Local and Nonlocal Closure Model for the Atmospheric Boundary
11	Layer. Part II: Application and Evaluation in a Mesoscale Meteorological Model, J. Appl.
12	Meteorol. Climatol., 46, 1396–1409, doi:10.1175/JAM2534.1, 2007b.
13	Pleim J. E. and R. Gilliam: An indirect data assimilation scheme for deep soil temperature in the
14	Pleim-Xiu land surface model. J. Appl. Meteor. Clim., 48, 1362-1376, 2009.
15	Richter, A., Burrows, J. P., Nues, H., Granier, C., and Niemeier, U.: Increase in tropospheric
16	nitrogen dioxide over China observed from space, Nature, 437, 129–132, 2005.
17	Pandis, S. N. and Seinfeld, J. H.: Sensitivity analysis of a chemical mechanism for aqueous -
18	phase atmospheric chemistry. Journal of Geophysical Research: Atmospheres (1984 -
19	2012), 94(D1), 1105-1126, 1989.
20	Pinder, R. W., Gilliland, A. B. and Dennis, R. L.: Environmental impact of atmospheric NH3
21	emissions under present and future conditions in the eastern United States, Geophys. Res.
22	Lett., 35, L12808, doi:10.1029/2008GL033732, 2008a.
23	Pinder, R. W., Dennis, R. L., and Bhave, P. V.: Observable indicators of the sensitivity of PM<

1	sub> 2.5 nitrate to emission reductions—Part I: Derivation of the adjusted gas ratio
2	and applicability at regulatory-relevant time scales. Atmospheric Environment, 42(6),
3	1275-1286, 2008b.
4	Schumann, U. and Huntrieser, H.: The global lightning-induced nitrogen oxides source, Atmos.
5	Chem. Phys., 7, 3823-3907, doi:10.5194/acp-7-3823-2007, 2007.
6	Sharma, S., Lavoué, D., Cachier, H., Barrie, L. A., and Gong, S. L.: Long - term trends of the
7	black carbon concentrations in the Canadian Arctic. Journal of Geophysical Research:
8	Atmospheres (1984 - 2012), 109, D15203, doi:10.1029/2003JD004331, 2004.
9	Sickles, J.E., II, and Shadwick, D.S.: Changes in air quality and atmospheric deposition in the
10	eastern United States: 1990-2004. J.Geophys. Res., 112, D17302, doi:
11	10.1029/2006JD007843, 2007.
12	Sillman, S.: The relation between ozone, NOx and hydrocarbons in urban and polluted rural
13	environments, Atmos. Environ., 33, 1821–1845, 1999.
14	Simpson D., Fagerli, H., Jonson, J.E., Tsyro, S., Wind, P., and Tuovinen, J.: "Transboundary
15	Acidification, Eutrophication, and Ground Level Ozone in Europe – Part I: Unified EMEP
16	Model Description." EMEP Status Report 2003, The Norwegian Meteorological Institute,
17	Oslo, 25 Norway, 2003.
18	Tesche, T.W., Morris, R., Tonnesen, G., McNally, D., Boylan, J., Brewer, P.: CMAQ/CAMx
19	annual 2002 performance evaluation over the eastern United States. Atmospheric
20	Environment 40, 4906–4919, 2006.
21	Trainer, M., Parrish, D. D., Buhr, M. P., Norton, R. B., Fehsenfeld, F. C., Anlauf, K. G.,
22	Bottenheim, J.W., Tang, Y.Z., Wiebe, H.A., Roberts, J.M., Tanner, R.L., Newman, L.,
23	Bowersox, V.C., Meagher, J.F., Olszyna, K.J., Rodgers, M.O., Wang, T., Berresheim, H., 36

1	Demerjian, K.L. and Roychowdhury, U. K.: Correlation of ozone with NOy in
2	photochemically aged air. Journal of Geophysical Research: Atmospheres (1984-2012),
3	98(D2), 2917-2925, 1993
4	Tsimpidi, A.P., Karydis, V.A., and Pandis, S.N.: Response of Inorganic Fine Particulate Matter to
5	Emission Changes of Sulfur Dioxide and Ammonia: The Eastern United States as a Case
6	Study, J. Air & Waste Manage. Assoc. 57:1489–1498, DOI:10.3155/1047-3289.57.12.1489,
7	2007.
8	Wakamatsu, S., Morikawa, T. and Ito, A.: Air Pollution Trends in Japan between 1970 and 2012
9	and Impact of Urban Air Pollution Countermeasures, Asian Journal of Atmospheric
10	Environment, Vol. 7-4, pp.177-190, December, 2013
11	Wang, K., Zhang, Y., Jang, C., Phillips, S., and Wang, B.: Modeling intercontinental air pollution
12	transport over the trans - Pacific region in 2001 using the Community Multiscale Air
13	Quality modeling system. Journal of Geophysical Research: Atmospheres (1984 - 2012),
14	114, D04307, doi:10.1029/2008JD010807, 2009
15	Wang, S., Xing, J., Chatani, S., Hao, J., Klimont, Z., Cofala, J., and Amann, M.: Verification of
16	anthropogenic emissions of China by satellite and ground observations. Atmospheric
17	Environment, 45(35), 6347-6358, 2011a.
18	Wang S., Xing J, Jang C, Zhu Y, Fu J, and Hao J.: Impact assessment of ammonia emissions on
19	inorganic aerosols in east China using response surface modeling technique. Environmental
20	Science and Technology, 45, 9293–9300, DOI: 10.1021/es2022347, 2011b.
21	Wang, S., Xing, J., Zhao, B., Jang, C., and Hao, J.: Effectiveness of national air pollution control
22	policies on the air quality in metropolitan areas of China. Journal of Environmental
23	Sciences, 26(1), 13-22, 2014.
	27

1	Wang, Y., Konopka, P., Liu, Y., Chen, H., Müller, R., Plöger, F., Riese, M., Cai, Z., and Lü, D.:
2	Tropospheric ozone trend over Beijing from 2002-2010: ozonesonde measurements and
3	modeling analysis, Atmos. Chem. Phys., 12, 8389-8399, doi:10.5194/acp-12-8389-2012,
4	2012.
5	Wild, M.: Global dimming and brightening: A review, Journal of Geophysical Research, 114,
6	D00D16, doi: 10.1029/2008JD011470, 2009.
7	Xing, J., Zhang, Y., Wang, S., Liu, X., Cheng, S., Zhang, Q., Chen, Y., Hao, J. and Wang, W.:
8	Modeling study on the air quality impacts from emission reductions and atypical
9	meteorological conditions during the 2008 Beijing Olympics. Atmospheric Environment,
10	45(10), 1786-1798, 2011a.
11	Xing, J., Wang, S. X., Jang, C., Zhu, Y., and Hao, J. M.: Nonlinear response of ozone to
12	precursor emission changes in China: a modeling study using response surface
13	methodology, Atmos. Chem. Phys., 11, 5027-5044, doi:10.5194/acp-11-5027-2011, 2011b.
14	Xing, J., Pleim, J., Mathur, R., Pouliot, G., Hogrefe, C., Gan, CM., and Wei, C.: Historical
15	gaseous and primary aerosol emissions in the United States from 1990 to 2010, Atmos.
16	Chem. Phys., 13, 7531-7549, doi:10.5194/acp-13-7531-2013, 2013.
17	Xiu, A. and J.E. Pleim: Development of a Land Surface Model. Part I: Application in a
18	Mesoscale Meteorological Model. J. Appl. Meteor. 40:192-209, 2001.
19	Xu, X., and Lin, W.: Trends of Tropospheric Ozone over China Based on Satellite Data (1979-
20	2005), Advances in Climate Change Research, 2, 43-48, 2011.
21	Yamaji, K., Ohara, T., Uno, I., Tanimoto, H., Kurokawa, J. I., and Akimoto, H.: Analysis of the
22	seasonal variation of ozone in the boundary layer in East Asia using the Community Multi-
23	scale Air Quality model: What controls surface ozone levels over Japan?. Atmospheric

Environment, 40(10), 1856-1868, 2006.

2	Zhang, H., Chen, G., Hu, J., Chen, SH., Wiedinmyer, C., Kleeman, M., and Ying, Q.:
3	Evaluation of a seven-year air quality simulation using the Weather Research and
4	Forecasting (WRF)/Community Multiscale Air Quality (CMAQ) models in the eastern
5	United States, Science of The Total Environment, Volumes 473-474, 1 March, Pages 275-
6	285, 2014.

- Zhang, X., Zhang, P., Zhang, Y., Li, X., and Qiu, H.: The trend, seasonal cycle, and sources of
 tropospheric NO2 over China during 1997–2006 based on satellite measurement. Science
 in China Series D: Earth Sciences, 50(12), 1877-1884, 2007.
- Zhang, X., van Geffen, J., Liao, H., Zhang, P., and Lou, S. J.: Spatiotemporal variations of
 tropospheric SO2 over China by SCIAMACHY observations during 2004–2009. Atmos.
 Environ., 60, 238–246, 2012.
- Zhang, Y., Vijayaraghavan, K., Wen, X.-Y., Snell, H. E., and Jacobson, M. Z.: Probing into
 regional ozone and particulate matter pollution in the United States: 1. A 1 year CMAQ
 simulation and evaluation using surface and satellite data, J. Geophys. Res., 114, D22304,
 doi:10.1029/2009JD011898, 2009.
- Zhao B, Wang S X, Dong X Y, Wang J D, Duan L, Fu X, Hao, J., and Fu, J.: Environmental
 effects of the recent emission changes in China: Implications for particulate matter
 pollution and soil acidification. Environmental Research Letters, 8, 24031,
 doi:10.1088/1748-9326/8/2/024031, 2013.

Species	Network Region		Number of sites (at least 18-year available with >75% annual coverage)	Time period	record frequency
Gaseous s					
	CASTNET	United States	38 selected from 133	1990-2010	Weekly
	AQS	United States	280 selected from 1177	1990-2010	Annual
SO_2	AIRBASE	Europe	126 selected from 510	1990-2010	Annual
	EMEP	Europe	44 selected from 237	1990-2010	Monthly
	API	China	7	2005-2010	Annual
	AQS	United States	181 selected from 714	1990-2010	Annual
NO	AIRBASE	Europe	160 selected from 440	1990-2010	Annual
NO ₂	EMEP	Europe	39 selected from 237	1990-2010	Monthly
	API	China	7	2005-2010	Annual
	CASTNET*	United States	25 selected from 133	1990-2010	Daily
	AIRBASE	Europe	147 selected from 315	1990-2010	Annual
O_3	EMEP	Europe	69 selected from 190	1990-2010	Daily
	WDCGG	Global(Japan used only)	3 selected from 102	1990-2010	Hourly
Particles					
	CASTNET	United States	38 selected from 133	1990-2010	Weekly
SO_4^{2-}	IMPROVE	United States	27 selected from 197	1990-2010	Semi-weekly
	EMEP	Europe	39 selected from 237	1990-2010	Monthly
	CASTNET	United States	38 selected from 133	1990-2010	Weekly
NO_3^-	IMPROVE	United States	27 selected from 197	1990-2010	Semi-weekly
	EMEP	Europe	12 selected from 237	1990-2010	Monthly
NH.+	CASTNET	United States	38 selected from 133	1990-2010	Weekly
1 N Π4	EMEP	Europe	6 selected from 237	1990-2010	Monthly
EC	IMPROVE	United States	26 selected from 197	1990-2010	Semi-weekly

Table 1 Summary of long-term observations used for trends analysis in this study

* There're few O₃ records from CASTNET in winter, thus criteria is set as at least 15 available

years with >75% coverage from March to November for each year

Table 2 Model performance

1	
2	

(a) Gaseous species

Spacias	Notwork		Obs	D	MB	NMB	RMSE	NME	Ν
species	INCLWOIK		(µg m ⁻³)	K	(µg m ⁻³)	(%)	(µg m ⁻³)	(%)	pairs
		Spring	5.0	0.73	-1.1	-21.8	3.2	72.4	2316
		Summer	3.3	0.74	0.2	5.3	2.4	93.4	2352
	US-CASTNET	Fall	4.5	0.78	1.6	36.0	3.8	118.0	2348
		Winter	8.1	0.67	-2.7	-33.4	6.0	81.7	2317
		Annual	5.2	0.67	-0.5	-9.4	4.1	91.5	9333
	US-AQS	Annual	12.2	0.2	-4.6	-37.5	10.6	135.3	2628
SO_2	EU- AIRBASE	Annual	8.7	0.3	-1.5	-17.7	9.6	98.8	580
		Spring	2.4	0.43	2.0	82.2	5.0	239.8	2399
		Summer	1.6	0.44	2.4	150.1	4.7	325.0	2355
	EU-EMEP	Fall	2.2	0.48	2.2	102.7	4.9	324.1	2344
		Winter	3.8	0.50	0.1	3.6	5.2	177.6	2363
		Annual	2.5	0.43	1.7	67.0	5.0	266.3	9461
_	CN-API	Annual	50.8	0.33	-18.4	-36.3	28.4	42.2	42
	US-AQS	Annual	29.0	0.2	-13.9	-47.9	22.6	63.4	1616
_	EU- AIRBASE	Annual	32.0	0.4	-17.1	-53.5	22.5	55.9	747
-	EU-EMEP	Spring	6.5	0.65	-0.1	-1.6	5.6	79.5	2049
NO		Summer	5.0	0.56	-0.7	-14.1	4.7	73.8	2066
NO_2		Fall	7.1	0.67	1.0	14.4	7.0	84.1	2084
		Winter	9.7	0.68	1.3	13.9	7.9	91.6	2068
		Annual	7.1	0.68	0.4	5.6	6.4	82.3	8267
-	CN-API	Annual	46.6	0.08	-31.5	-67.5	36.1	66.2	42
		Spring	168.1	0.52	-22.8	-13.6	29.7	16.1	1269
	US-CASTNET	Summer	176.8	0.59	-14.3	-8.1	30.5	14.5	1512
		Fall	155.3	0.60	-3.9	-2.5	23.5	12.4	1071
		Winter	112.5	0.51	-3.6	-3.2	10.1	7.6	217
_	EU-AIRBASE	Annual	169.4	0.40	14.4	8.5	38.9	17.4	2776
_		Spring	140.9	0.56	-2.1	-1.5	22.7	14.2	4145
O_3^*		Summer	152.3	0.60	6.5	4.3	30.5	18.4	4161
	EU-EMEP	Fall	108.5	0.66	18.4	16.9	25.4	25.9	4151
		Winter	92.5	0.29	3.1	3.4	16.1	16.6	4111
_		Spring	165.4	0.68	-8.9	-5.4	26.1	14.4	175
		Summer	157.3	0.83	10.8	6.9	34.0	21.4	172
	WDCGG-JP	Fall	128.5	0.62	17.4	13.5	31.4	21.9	173
		Winter	109.2	0.49	3.2	2.9	15.1	12.6	172

3 * Comparison of O₃ concentration is computed on the basis of annual or seasonal maximum of

4 DM8 (daily 8-hour maxima) value, except that for AIRBASE which is computed on the basis of

5 annual maxima of DM1 (daily 1-hour maxima)

(b) Fine particles

C	N - 4		Obs	л	MB	NMB	RMSE	NME	Ν
Species	Network		(µg m ⁻³)	K	(µg m ⁻³)	(%)	(µg m ⁻³)	(%)	pairs
		Spring	3.1	0.87	-0.2	-7.5	0.8	29.2	2316
		Summer	5.3	0.86	-2.4	-45.2	3.1	44.7	2352
	US-CASTNET	Fall	3.7	0.86	-1.0	-26.5	1.8	34.3	2348
		Winter	2.3	0.63	-0.8	-35.6	1.2	53.1	2316
_		Annual	3.6	0.81	-1.1	-30.8	1.9	40.3	9332
		Spring	1.4	0.89	0.3	22.5	0.7	70.3	1602
		Summer	2.2	0.90	-0.6	-28.9	1.8	37.8	1596
SO_4^{2-}	US- IMPROVE	Fall	1.3	0.90	0.2	15.7	0.7	68.4	1605
		Winter	0.9	0.76	0.1	16.3	0.6	106.7	1605
-		Annual	1.4	0.85	0.0	0.7	1.1	70.8	6408
		Spring	2.6	0.68	0.3	12.5	1.4	52.3	2099
		Summer	2.4	0.68	0.1	3.7	1.3	41.4	2071
	EU- EMEP	Fall	2.2	0.64	0.0	1.9	1.4	55.9	2042
		Winter	2.4	0.53	-0.7	-28.6	1.9	58.3	2058
		Annual	2.4	0.61	-0.1	-2.4	1.5	51.9	8270
		Spring	1.1	0.69	1.0	92.9	2.1	195.5	2316
		Summer	0.4	0.31	-0.2	-48.2	0.4	76.1	2352
	US-CASTNET	Fall	0.7	0.68	0.1	13.8	0.7	99.3	2348
		Winter	1.6	0.71	1.2	75.2	1.9	262.0	2316
_		Annual	0.9	0.72	0.5	56.4	1.5	157.7	9332
	US- IMPROVE	Spring	0.4	0.72	0.4	106.9	1.0	164.8	1602
		Summer	0.2	0.10	-0.1	-40.5	0.2	93.0	1596
NO ₃ -		Fall	0.3	0.66	0.0	11.4	0.4	125.7	1604
		Winter	0.5	0.66	0.5	94.8	1.1	226.9	1605
-		Annual	0.3	0.66	0.2	59.1	0.8	152.7	6407
		Spring	3.0	0.75	0.3	10.8	2.0	75.2	679
	EU- EMEP	Summer	1.8	0.74	-1.2	-67.0	1.5	74.7	656
		Fall	2.3	0.72	-0.4	-15.0	1.5	64.4	659
		Winter	2.6	0.64	0.6	23.1	2.1	91.2	671
		Annual	2.4	0.70	-0.2	-6.3	1.8	76.4	2665
		Spring	1.2	0.68	0.3	22.6	0.8	52.0	2316
		Summer	1.6	0.77	-0.8	-53.7	1.1	50.5	2352
	US-CASTNET	Fall	1.2	0.72	-0.3	-21.4	0.6	31.7	2348
		Winter	1.1	0.76	0.2	19.0	0.6	54.1	2316
NH_4^+		Annual	1.3	0.52	-0.2	-12.9	0.8	47.0	9332
-		Spring	1.4	0.69	0.7	51.3	1.4	101.4	335
		Summer	1.2	0.64	-0.2	-15.2	0.9	43.9	330
	EU- EMEP	Fall	1.2	0.67	0.3	28.2	1.0	73.7	332
		Winter	1.1	0.62	0.8	68.4	1.4	110.4	328
		Annual	1.2	0.62	0.4	33.7	1.2	82.4	1325
		Spring	0.2	0.79	-0.1	-62.5	0.2	62.7	1536
		Summer	0.3	0.54	-0.2	-73.5	0.3	92.7	1532
EC	US- IMPROVE	Fall	0.3	0.81	-0.2	-64.4	0.3	65.9	1548
		Winter	0.2	0.85	-0.1	-59.4	0.2	55.7	1542
		Annual	0.2	0.74	-0.2	-65.1	0.3	69.2	6158

	Eastern C	hina	Eastern	US	Europe		
Emission	kg km ⁻² yr ⁻¹	% yr-1	kg km ⁻² yr ⁻¹	% yr ⁻¹	kg km ⁻² yr ⁻¹	% yr ⁻¹	
SO_2	20.2	3.2	-16.1	-5.4	-20.4	-5.4	
NO _x	8.5	4.3	-3.7	-1.8	-3.0	-1.5	
VOC	18.6	2.3	-22.5	-3.3	-26.7	-3.3	
NH ₃	6.5	2.6	1.7	1.6	-2.6	-1.0	
PM_{10}	2.1	0.3	-4.5	-4.6	-10.0	-4.8	
Concentration	µg m ⁻³ yr ⁻¹	% yr ⁻¹	µg m ⁻³ yr ⁻¹	% yr ⁻¹	µg m ⁻³ yr ⁻¹	% yr ⁻¹	
SO_2	0.265	2.70	-0.175	-5.71	-0.178	-5.06	
NO_2	0.119	4.14	-0.048	-1.38	-0.040	-1.16	
*O ₃	2.566	1.49	-1.028	-0.66	-0.875	-0.54	
PM _{2.5}	0.481	2.21	-0.097	-1.21	-0.253	-2.62	
SO_4^{2-}	0.185	2.82	-0.072	-3.17	-0.109	-3.73	
NO ₃ -	0.097	5.40	0.014	1.61	-0.030	-1.84	
NH_4^+	0.081	3.44	-0.006	-0.72	-0.041	-2.91	
EC	0.005	0.99	-0.004	-3.39	-0.005	-2.46	

Table 3 Simulated trends in three regions (grid-averaged)

Colored entries are significant at p=0.05 level: green=significant decrease; orange=significant increase.

 \ast Trend in O₃ is computed on the basis of annual or seasonal maximum of DM8 (daily 8-hour maxima) value

1Table 4 Comparison of observed and simulated trend2(unit: μ g m⁻³ yr⁻¹, computed on the basis of annual and seasonal means over the 1990-20103period with a linear least square fit method) and the annual change rate (x%, i.e., concentration in4the year Y (C_Y) will be fit as C_Y=C₁₉₉₀×(1+x)^{Y-1990})

$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	C	N a trave alla		Spi	ring	Sun	nmer	Fa	all	Wi	nter	Ar	nnual
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	species	INCLWOIK		obs	sim								
$\frac{103-CASTNET}{6} - \frac{100}{6} - \frac{4.74}{6} - 6.26 - 4.91 - 6.13 - 5.61 - 6.63 - 4.79 - 7.01 - 4.98 - 6.57}{-0.626 - 0.467} - \frac{100}{6} -$		US CASTNET	µg m⁻³	-0.228	-0.238	-0.152	-0.204	-0.234	-0.385	-0.368	-0.366	-0.245	-0.298
$\begin{array}{c c c c c c c c c c c c c c c c c c c $		US-CASINEI	%	-4.74	-6.26	-4.91	-6.13	-5.61	-6.63	-4.79	-7.01	-4.98	-6.57
$SO_{2} = \frac{10000}{EU-AIRBASE} = \frac{\mu g m^{-3}}{\%} = \frac{0.873 - 0.441}{0.873} = \frac{-0.873 - 0.441}{0.873} = \frac{-0.873 - 0.441}{0.873} = \frac{-0.873 - 0.441}{0.873} = \frac{-0.873 - 0.441}{0.886} = \frac{-0.873 - 0.441}{0.886} = \frac{-0.873 - 0.441}{0.886} = \frac{-0.873 - 0.441}{0.202} = \frac{-0.873 - 0.441}{0.202} = \frac{-0.873 - 0.441}{0.202} = \frac{-0.873 - 0.441}{0.204} = \frac{-0.204 - 0.204}{0.204} = \frac{-0.204}{0.204} $			µg m⁻³									-0.626	-0.467
$\begin{array}{c c c c c c c c c c c c c c c c c c c $		US-AQS	%									-5.31	-6.45
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	50.		µg m⁻³									-0.873	-0.441
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	302	EU-AIKDASE	%									-8.86	-5.86
$\begin{array}{c c c c c c c c c c c c c c c c c c c $		ELLEMED	µg m⁻³	-0.187	-0.282	-0.108	-0.225	-0.180	-0.279	-0.339	-0.264	-0.204	-0.262
$\begin{tabular}{c c c c c c c c c c c c c c c c c c c $		EU-EIVIEF	%	-7.03	-6.16	-5.95	-5.53	-7.28	-6.23	-8.04	-6.28	-7.26	-6.05
		CN ADI	µg m⁻³									0.376	1.230
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		CIV-AI I	%									0.66	4.02
$\frac{1000 - 2.3}{6} = \frac{1000}{6} = \frac{1000}{6}$		US AOS	µg m⁻³									-0.629	-0.311
EU-AIRBASE $\frac{\mu g m^{-3}}{\%}$ -0.640 -0.136		US-AQS	%									-2.3	-2.2
-1.88 -0.86		ELL AIDRASE	µg m⁻³									-0.640	-0.136
	NO	EU-AIKDASE	%									-1.88	-0.86
$\mu g m^{-3} -0.087 -0.113 -0.115 -0.137 -0.150 -0.194 -0.150 -0.195 -0.126 -0.160$	INO2	ELLEMED	µg m⁻³	-0.087	-0.113	-0.115	-0.137	-0.150	-0.194	-0.150	-0.195	-0.126	-0.160
% -1.29 -1.64 -2.26 -3.03 -2.00 -2.30 -1.46 -1.70 -1.69 -2.04		EU-EIVIEF	%	-1.29	-1.64	-2.26	-3.03	-2.00	-2.30	-1.46	-1.70	-1.69	-2.04
-0.454 = -0.454 = 0.868		CN ADI	μg m ⁻³									-0.454	0.868
-0.97 = 5.94		CIN-AFI	%									-0.97	5.94
на състъчит µg m ⁻³ -1.187 -0.903 -1.860 -1.010 -1.220 -0.527 -0.029 -0.134 -1.859 -0.952			µg m⁻³	-1.187	-0.903	-1.860	-1.010	-1.220	-0.527	-0.029	-0.134	-1.859	-0.952
$\frac{100}{7} -0.73 -0.65 -1.14 -0.68 -0.83 -0.36 -0.02 -0.13 -1.10 -0.64$		US-CASTNET		-0.73	-0.65	-1.14	-0.68	-0.83	-0.36	-0.02	-0.13	-1.10	-0.64
-1.348 -2.129		EU-AIRBASE	ug m ⁻³									-1.348	-2.129
EU-AIRBASE $\frac{-1.13}{\%}$			%									-0.79	-1.13
$\mu g m^{-3} -0.651 -1.281 -1.207 -1.365 -0.157 -0.184 0.124 -0.048 -1.067 -1.313$		EU-EMEP	µg m ⁻³	-0.651	-1.281	-1.207	-1.365	-0.157	-0.184	0.124	-0.048	-1.067	-1.313
EU-EMEP % -0.46 -0.92 -0.85 -0.91 -0.13 -0.15 0.14 -0.05 -0.74 -0.87	^ *		%	-0.46	-0.92	-0.85	-0.91	-0.13	-0.15	0.14	-0.05	-0.74	-0.87
$\frac{1}{1000} = \frac{1}{1000} \frac{1}{10$	03	WDCGG-	µg m⁻³	0.485	-0.029	-1.131	-0.083	-0.688	0.090	-0.416	0.413	0.232	-0.126
Minamitorishima % 0.35 -0.02 -1.19 0.01 -0.70 0.09 -0.31 0.38 0.18 -0.11		Minamitorishima	%	0.35	-0.02	-1.19	0.01	-0.70	0.09	-0.31	0.38	0.18	-0.11
$\mu g m^{-3}$ 1.305 0.372 0.549 0.259 -0.638 0.308 0.166 0.217 0.702 0.440		WDCGG- Ryori	µg m⁻³	1.305	0.372	0.549	0.259	-0.638	0.308	0.166	0.217	0.702	0.440
wDCGG-Ryon $\frac{10}{\%}$ 0.79 0.24 0.44 0.18 -0.47 0.25 0.24 0.23 0.41 0.29			%	0.79	0.24	0.44	0.18	-0.47	0.25	0.24	0.23	0.41	0.29
WDCGG- µg m ⁻³ -1.073 -0.019 -4.015 -0.375 0.581 -1.017 -0.368 0.861 -3.299 -0.022		WDCGG-	µg m⁻³	-1.073	-0.019	-4.015	-0.375	0.581	-1.017	-0.368	0.861	-3.299	-0.022
Tsukuba % -0.60 -0.02 -1.78 -0.18 0.52 -0.56 -0.31 0.74 -1.40 -0.01		Tsukuba	%	-0.60	-0.02	-1.78	-0.18	0.52	-0.56	-0.31	0.74	-1.40	-0.01
μg m ⁻³ -0.070 -0.073 -0.161 -0.125 -0.112 -0.098 -0.054 -0.046 -0.099 -0.086			µg m ⁻³	-0.070	-0.073	-0.161	-0.125	-0.112	-0.098	-0.054	-0.046	-0.099	-0.086
US-CASTNET $\frac{-100}{\%}$ -2.30 -2.49 -3.25 -4.45 -3.31 -3.75 -2.25 -3.01 -2.87 -3.46		US-CASTNET	%	-2.30	-2.49	-3.25	-4.45	-3.31	-3.75	-2.25	-3.01	-2.87	-3.46
μg m ⁻³ -0.023 -0.021 -0.049 -0.043 -0.036 -0.041 -0.024 -0.016 -0.033 -0.030	10^{2}		µg m⁻³	-0.023	-0.021	-0.049	-0.043	-0.036	-0.041	-0.024	-0.016	-0.033	-0.030
SO_4^2 US-IMPROVE $\frac{10}{\%}$ -1.76 -1.24 -2.45 -2.86 -2.87 -2.69 -2.76 -1.59 -2.43 -2.11	5042	US-IMPROVE	%	-1.76	-1.24	-2.45	-2.86	-2.87	-2.69	-2.76	-1.59	-2.43	-2.11
μg m ⁻³ -0.119 -0.086 -0.111 -0.112 -0.097 -0.085 -0.090 -0.060 -0.104 -0.086			µg m⁻³	-0.119	-0.086	-0.111	-0.112	-0.097	-0.085	-0.090	-0.060	-0.104	-0.086
EU-EMEP $\frac{10}{\%}$ -4.28 -2.84 -4.35 -4.49 -4.27 -3.93 -3.39 -3.29 -4.06 -3.62		EU-EMEP	%	-4.28	-2.84	-4.35	-4.49	-4.27	-3.93	-3.39	-3.29	-4.06	-3.62
μg m ⁻³ -0.009 0.023 -0.011 0.005 -0.015 0.023 0.009 0.057 -0.006 0.027			µg m⁻³	-0.009	0.023	-0.011	0.005	-0.015	0.023	0.009	0.057	-0.006	0.027
$\frac{100}{\%} -0.94 1.19 -3.17 3.38 -2.27 3.33 0.61 2.35 -0.73 2.10$		US-CASTNET	%	-0.94	1.19	-3.17	3.38	-2.27	3.33	0.61	2.35	-0.73	2.10
μg m ⁻³ -0.002 0.012 -0.004 0.000 -0.005 0.010 -0.002 0.024 -0.003 0.012	NO -		µg m⁻³	-0.002	0.012	-0.004	0.000	-0.005	0.010	-0.002	0.024	-0.003	0.012
NO ₃ US-IMPROVE $\frac{10}{\%}$ -0.70 1.93 -2.13 0.14 -1.97 3.73 -0.28 2.99 -1.04 2.53	NO ₃	US-IMPROVE	%	-0.70	1.93	-2.13	0.14	-1.97	3.73	-0.28	2.99	-1.04	2.53
$\mu g m^{-3} -0.015 -0.086 -0.019 -0.032 -0.009 -0.043 0.013 -0.002 -0.008 -0.041$			µg m⁻³	-0.015	-0.086	-0.019	-0.032	-0.009	-0.043	0.013	-0.002	-0.008	-0.041
EU-EMEP % -0.47 -2.49 -1.06 -5.38 -0.51 -2.19 0.50 -0.13 -0.33 -1.74		EU-EMEP	%	-0.47	-2.49	-1.06	-5.38	-0.51	-2.19	0.50	-0.13	-0.33	-1.74
μg m ⁻³ -0.023 -0.002 -0.038 -0.010 -0.032 -0.006 -0.013 0.012 -0.026 -0.002			µg m⁻³	-0.023	-0.002	-0.038	-0.010	-0.032	-0.006	-0.013	0.012	-0.026	-0.002
US-CASINEI % -2.04 -0.19 -2.60 -1.54 -2.86 -0.68 -1.24 0.97 -2.19 -0.18	NTT	US-CASTNET	%	-2.04	-0.19	-2.60	-1.54	-2.86	-0.68	-1.24	0.97	-2.19	-0.18
NH4' μg m ⁻³ 0.003 -0.055 0.000 -0.049 0.020 -0.035 -0.002 -0.018 0.005 -0.039	$NH4^{T}$		µg m ⁻³	0.003	-0.055	0.000	-0.049	0.020	-0.035	-0.002	-0.018	0.005	-0.039
EU-EMEP % 0.80 -2.22 0.30 -4.52 1.75 -2.21 0.16 -0.87 0.70 -2.19		EU-EMEP	%	0.80	-2.22	0.30	-4.52	1.75	-2.21	0.16	-0.87	0.70	-2.19
μg m ⁻³ -0.005 -0.002 -0.002 -0.009 -0.004 -0.008 -0.003 -0.006 -0.003			µg m ⁻³	-0.005	-0.002	-0.003	-0.002	-0.009	-0.004	-0.008	-0.003	-0.006	-0.003
EC US-IMPROVE $\frac{1}{\%}$ -2.46 -2.77 -1.34 -3.42 -3.30 -3.67 -3.41 -3.32 -2.64 -3.32	EC	US-IMPKOVE	%	-2.46	-2.77	-1.34	-3.42	-3.30	-3.67	-3.41	-3.32	-2.64	-3.32

- 1
- 2 Colored entries are significant at p=0.05 level: green=significant decrease; orange=significant increase.
- 3 * Trend in O₃ is computed on the basis of annual or seasonal maximum of DM8 (daily 8-hour
- 4 maxima) value, except that for AIRBASE which is computed on the basis of annual maximum of
- 5 DM1 (daily 1-hour maxima)



3 Fig. 1 Processes of gridded emissions for northern hemispheric WRF-CMAQ simulation

(a) SO₂





2

Fig. 2 EDGAR emission trend over 1990 to 2010 for SO₂, NO_x, NMVOC and NH₃
(unit: kg km⁻² yr⁻¹, computed on the basis of annual means over the 1990-2010 period with a linear least square fit method)





Fig. 3 (a) simulated SO₂ trend from WRF-CMAQ (unit: μgm⁻³ yr⁻¹); (b) upper-color map: simulated SO₂ trend in East China overlaid with observed SO₂ trend from China-API, dot represents each observation site, computed on the basis of annual means over the 2005–2010 period with a linear least square fit method, dot size is determined by the significance of trend, i.e., larger symbols denote more significant trends at 0.05 level (unit: μg m⁻³ yr⁻¹); lower-scatter

6 plot: observed and simulated SO₂ concentration, network-mean for each year corresponding grid

1	cells from model simulation are selected for comparison (unit: $\mu g m^{-3}$); (c) same as (b) for
2	Europe – AIRBASE; (d) same as (b) for Europe – EMEP; (e) same as (b) for the U.S. – AQS; (f)
3	same as (b) for the U.S. – CASTNET
4	



Fig. 4 Same as Fig. 3 for NO₂



Fig. 5 Same as Fig. 3 for O₃ (unit: μg m⁻³, computed on the basis of annual or seasonal maximum of DM8 (daily 8h maxima) value, except that for AIRBASE which is computed on the basis of annual maximum of DM1 (daily 1h maxima); three sites of WDCGG are S1- Minamitorishima, lat: 24.28, lon: 153.98, S2- Ryori, lat: 39.03, lon: 141.82, S3-Tsukuba, lat: 36.05, lon:140.13)



Fig. 6 Same as Fig. 3 for SO₄²⁻















3

Fig. 11 Changes in PM chemistry from modeling results

4 (calculation based on molecular units; grid-averaged for three regions; (NO_x+2*SO₂) represents
 5 the amount of NH₃ needed for complete neutralization; DSN- degree of sulfate neutralization;
 6 Nitration ratio = NO₃⁻ concentration/NO_x emission)