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# Inverse modeling of Texas NO<sub>x</sub> emissions using space-based and ground-based NO<sub>2</sub> observations

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Inverse modeling of nitrogen oxide (NO<sub>x</sub>) emissions using satellite-based NO<sub>2</sub> observations has become more prevalent in recent years, but has rarely been applied to regulatory modeling at regional scales. In this study, OMI satellite observations of NO<sub>2</sub> column densities are used to conduct inverse modeling of NO<sub>x</sub> emission inventories for two Texas State Implementation Plan (SIP) modeling episodes. Addition of lightning, aircraft, and soil NO<sub>x</sub> emissions to the regulatory inventory narrowed but did not close the gap between modeled and satellite observed NO<sub>2</sub> over rural regions. Satellitebased top-down emission inventories are created with the regional Comprehensive Air Quality Model with extensions (CAMx) using two techniques: the direct scaling method and discrete Kalman filter (DKF) with Decoupled Direct Method (DDM) sensitivity analysis. The simulations with satellite-inverted inventories are compared to the modeling results using the a priori inventory as well as an inventory created by a ground-level NO<sub>2</sub> based DKF inversion. The DKF inversions yield conflicting results: the satellitebased inversion scales up the a priori NO<sub>x</sub> emissions in most regions by factors of 1.02 to 1.84, leading to 3-55% increase in modeled NO2 column densities and 1-7 ppb increase in ground 8h ozone concentrations, while the ground-based inversion indicates the a priori NO<sub>x</sub> emissions should be scaled by factors of 0.34 to 0.57 in each region. However, none of the inversions improve the model performance in simulating aircraft-observed NO<sub>2</sub> or ground-level ozone (O<sub>3</sub>) concentrations.

#### Introduction

Nitrogen oxides  $(NO_x = NO + NO_2)$  in the troposphere are primary air pollutants, emitted from both anthropogenic sources like fossil-fuel combustion and biomass burning, and natural sources such as soil microbial processes and lightning. NO<sub>x</sub> also acts as a precursor of a secondary air pollutant, tropospheric O<sub>3</sub>, when it reacts with the oxidation products of volatile organic compounds (VOCs) in the presence of sunlight. Ox-

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idation with hydroxyl (OH) radical is the dominant sink of NO<sub>x</sub>, leading to atmospheric nitric acid (HNO<sub>3</sub>) formation. The atmospheric lifetime of tropospheric NO<sub>x</sub> varies from a few hours in summer to a couple of days in winter (Seinfeld and Pandis, 2006).

NO<sub>x</sub> emission inventories used in air quality modeling are typically developed by a bottom-up approach based on estimated activity rates and emission factors for each category. Due to inaccuracies in determining these rates and factors, the uncertainty in NO<sub>x</sub> emission inventories has been suggested to be as high as a factor of two and classified as one of the top uncertainties in ozone simulations and sensitivity analysis (Hanna et al., 2001; Xiao et al., 2010).

Inverse modeling techniques can be used with atmospheric models to estimate model variables that may not be directly measurable (Gilliland and Abbitt, 2001). Inverse modeling generates an optimized "top-down" NO<sub>x</sub> emission inventory for air quality models by minimizing the difference between observed and modeled NO2 concentrations, providing an opportunity to identify possible biases in the bottom-up NO<sub>x</sub> emission inventory (Napelenok et al., 2008). However, as uncertainties may also associate with the measurement data and the inverse methods themselves, inverse modeling has its own limitations. Hence, it is valuable to compare both bottom-up and top-down NO<sub>x</sub> emission inventories in order to improve the understanding of NO<sub>x</sub> emissions.

Several inverse modeling studies have used surface NO<sub>2</sub> measurements (Mendoza-Dominguez and Russell, 2000; Quélo et al., 2005; Pison et al., 2007) or aircraft NO<sub>2</sub> measurements (Brioude et al., 2011) to constrain NO<sub>x</sub> emissions. Compared to ground and aircraft measurements, satellite-based observations generate greater spatial coverage of NO<sub>2</sub>. Studies on combining satellite NO<sub>2</sub> measurements with inverse modeling techniques to create the top-down NO<sub>x</sub> emission inventories also have been conducted recently in both global scale (Martin et al., 2003; Müller and Stavrakou, 2005; Jaeglé et al., 2005; Lin et al., 2010) and regional scale modeling (Konovalov et al., 2006, 2008; Deguillaume et al., 2007; Napelenok et al., 2008; Kurokawa et al., 2009; Zhao and Wang, 2009; Chai et al., 2009).

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Discrete Kalman filter (DKF) (Prinn, 2000) is an inverse modeling method that solves the inverse problem iteratively, and can be applied to the cases with linear or weakly non-linear relationships between emissions and pollutants. It has been used in several studies to constrain emissions of carbon monoxide (Mulholland and Seinfeld, 1995), chloroflourocarbons (Haas-Laursen et al., 1996), isoprene (Chang et al., 1996) and ammonia (Gilliland et al., 2003). Most recently, Napelenok et al. (2008) applied the DKF method to the regional Community Multiscale Air Quality (CMAQ) model, generating a top-down NO<sub>x</sub> emission inventory for the southeastern United States using Scanning Imaging Absorption Spectrometer for Atmospheric Chartography (SCIAM-CHY) (Bovensmann et al., 1999) satellite NO<sub>2</sub> data.

Despite the growing number of scientific studies conducting satellite-based inversions of NO<sub>x</sub> emissions, the applicability of these methods to state-level regulatory attainment modeling has not been widely explored. In this work, the DKF method introduced by Napelenok et al. (2008) is applied with finer resolution satellite NO2 data now available from the Ozone Monitoring Instrument (OMI) as well as ground-level NO<sub>2</sub> observations, to constrain NO<sub>3</sub> emissions for actual regulatory modeling episode in Texas. Lightning and aircraft NO<sub>x</sub> emissions are added to the base case NO<sub>x</sub> emission inventory to address the bias noted by Napelenok et al. (2008) of regional models underestimating upper tropospheric NO<sub>x</sub>. The DKF inverted a posteriori emissions are compared to the base case emissions, the a priori emissions and a posteriori emissions derived by the inversion method of Martin et al. (2003).

# Methodology

## Model inputs and configurations

Base case model inputs were taken from episodes developed by the Texas Commission on Environmental Quality (TCEQ) for Texas ozone attainment planning. CAMx version 5.3 (ENVIRON, 2010) was used in this study to simulate two modeling episodes 13, 17479–17517, 2013

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in 2006 with high ozone concentrations in the Dallas-Fort Worth (DFW) region, from 31 May to 1 July, and in the Houston-Galveston-Brazoria (HGB) region, from 13 August to 15 September (Fig. 1). The NCAR/Penn State (National Center for Atmospheric Research/Pennsylvania State University) Mesoscale Model, Version 5, release 3.7.3 (MM5v.3.7.3) (Grell et al., 1994), conducted with the Eta-PBL scheme, was used to generate the meteorological fields with 43 vertical layers. The preprocessor MM5CAMx was used to convert MM5 outputs into CAMx-ready meteorology inputs. The vertical configuration of CAMx modeling consists of 17 vertical layers for the August-September modeling episode, whereas 28 vertical layers were used for the June modeling episode. Modeling was conducted with the Carbon Bond version 2005 (CB-05) chemical mechanism, PPM advection scheme, and K-theory vertical diffusion scheme (TCEQ, 2010, 2011). Boundary conditions for the 36 km eastern US domain were generated by the Model for Ozone and Related Chemical Tracers (MOZART) global model (ENVIRON, 2008).

### **Emission inventory**

Base case emission inventories were provided by TCEQ. The point source emissions were from the State of Texas Air Reporting System (STARS) database which collects emission information from approximately 2000 point sources annually, and the EPA's acid rain database (ARD) which contains emissions from electric generating units (EGUs). The on-road mobile emission inventory was generated by Motor Vehicle Emission Simulator 2010a (MOVES2010a), and the non-road mobile inventory was developed by National Mobile Inventory Model (NMIM) and the Texas NONROAD (TexN) mobile source model. The area source inventory was projected by the EPA Economic Growth Analysis System (EGAS) model based on 2005 emissions from the Texas Air Emissions Repository (TexAER) database. The Emission Processing System, version 3 (EPS3) (ENVIRON, 2007) was used for processing the point, mobile, and area emissions to the model-ready format (TCEQ, 2010, 2011). Biogenic emissions were generated by the Global Biosphere Emissions and Interactions System (GloBEIS) biogenics **ACPD** 

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emissions model, version 3.1 (Yarwood et al., 1999), with soil NO<sub>x</sub> emissions estimated by the Yienger and Levy method (Yienger and Levy, 1995).

Lightning and aircraft NO<sub>x</sub> emissions in the upper troposphere were missing in the base case emission inventories and should be added before conducting inversions. In this study, lightning NO emissions were developed based on National Lightning Detection Network (NLDN) data obtained from Vaisala Inc., following the approach of Kaynak et al. (2008). Intra-cloud lightning flashes were treated as three times of cloud-to-ground lightning flashes with 500 mol NO emission per flash. Lightning NO was placed into the model to match the time and location of NLDN flashes, and then distributed vertically based on the profile obtained from the mean April to September 2003–2005 vertical distribution of VHF sources from the Northern Alabama Lightning Mapping Array (Koshak et al., 2004). Global aircraft NO<sub>v</sub> emissions of year 2005 in 0.1° × 0.1° resolution were obtained from the Emission Database for Global Atmospheric Research (EDGAR) v4.1 (http://edgar.jrc.ec.europa.eu/datasets\_grid\_list41. php?v=41&edgar\_compound=NOx) and mapped to our modeling domain and placed at 9 km altitude.

### **Inversion regions**

Five urban areas: Houston-Galveston-Brazoria (HGB), Dallas-Fort Worth (DFW), Beaumont-Port Arthur (BPA), Northeast Texas (NE Texas), and Austin and San Antonio; plus two surrounding rural areas: North Rural area (N rural) and South Rural area (S rural) (Fig. 1) were designed as inversion regions for the DKF inversions of NO<sub>x</sub> emissions. The five urban regions are all air quality planning areas included in Texas SIP development (Gonzales and Williamson, 2011). HGB and DFW were classified by US EPA as ozone nonattainment areas for violating the 1997 ozone National Ambient Air Quality Standard (NAAQS) of 84 ppb. BPA was designated as an ozone maintenance area, and NE Texas, Austin and San Antonio were designated as ozone early action compact areas under that standard. However, the recent tightening of the NAAQS to 75 ppb has heightened interest in ozone reduction in all of these regions.

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The sensitivities of NO<sub>2</sub> concentrations to boundary conditions and to NO<sub>x</sub> emissions from each inversion region and the border region (the area between model boundary and inversion regions) were computed through DDM. The border region minimizes the impacts from boundary conditions on the inversion regions to the level of only 2%. The DDM sensitivities show that NO<sub>x</sub> emissions from each urban region has the most impact on NO<sub>2</sub> concentrations within that region, and has less than 10% influence on other regions.

#### Inversion methods 2.4

Two methods are applied for inverse modeling: a direct scaling method introduced by Martin et al. (2003), and the DKF method.

#### **Direct Scaling (DS) inversion method** 2.4.1

The DS method applies the ratio between satellite NO<sub>2</sub> observations and modeled NO<sub>2</sub> concentrations to scale the bottom-up NO<sub>x</sub> emissions in each grid cell:

$$E_{\rm t} = E_{\rm b} \times \frac{\Omega_{\rm s}}{\Omega_{\rm m}} \tag{1}$$

where  $E_t$  is the top-down NO<sub>x</sub> emission rate;  $E_b$  is the bottom-up NO<sub>x</sub> emission rate;  $\Omega_s$  and  $\Omega_m$  are the satellite and modeled NO<sub>2</sub> column densities, respectively.

This method was developed in a global model with coarse grid resolution and assumes that the NO<sub>2</sub> concentration in each model grid will not be affected by the NO<sub>x</sub> emitted from surrounding grids. However, in a regional model with relatively small grid size, this assumption may fail, generating a spatial smearing error when NO<sub>x</sub> lifetime is longer than the horizontal transport time (Martin et al., 2003; Boersma et al., 2008; Lamsal et al., 2010; Turner et al., 2012). Martin et al. (2003) indicated that the spatial smearing error can be neglected if the grid length is greater than 100 km. Therefore, smoothing kernels (Toenges-Schuller et al., 2006; Boersma et al., 2008; Lamsal et al.,

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2010) need to be applied in order to alleviate the spatial smearing error in CAMx by accounting for the emissions from adjacent grid cells in developing the top-down NO<sub>v</sub> emissions. The smoothing kernel is defined as

# $\mathbf{K} = \frac{1}{k+8} \begin{pmatrix} 1 & 1 & 1 \\ 1 & k & 1 \\ 1 & 1 & 1 \end{pmatrix}$ (2)

where k is a smoothing parameter, and is determined by applying the smoothing kernel (K) to each grid cell in the bottom-up  $NO_x$  emission inventory with different k values until the correlation between smoothed bottom-up NO<sub>x</sub> emissions and corresponding CAMx modeled NO<sub>2</sub> column density reaches a maximum. The smoothing kernel (**K**) is then applied to Eq. (1) to form Eq. (3),

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$$E_{i,j}^{t} = \frac{E_{i,j}^{b}}{\sum_{n=-1}^{1} \sum_{l=-1}^{1} \mathbf{K}_{l,n} E_{i+l,j+n}^{b}} \frac{\Omega_{s}}{\Omega_{m}} \times E_{i,j}^{b}$$
 (3)

where *i* and *j* represent column and row in horizontal model grids.

#### **DKF** inversion method 2.4.2

The direct scaling inversion approach, as described above, creates spatial smearing errors when applies to the regional models with fine resolution. Meanwhile, it assumes concentrations scale proportionally with emissions; hence, the nonlinearity between NO<sub>2</sub> concentrations and NO<sub>x</sub> emissions becomes problematic because NO<sub>x</sub> may influence its own lifetime by influencing concentrations of OH radicals (Martin et al., 2003). The DKF inversion (Fig. 2), however, solves the spatial smearing problem by taking the spatial relationship between NO<sub>2</sub> concentrations and NO<sub>x</sub> emissions directly from model simulations, and also reduces the non-linearity issue by performing the inversion iteratively.

$$\mathbf{G}_{k} = \mathbf{P}_{\mathrm{NO}_{x},k}^{-} \mathbf{S}_{k}^{T} \left( \mathbf{S}_{k} \mathbf{P}_{\mathrm{NO}_{x},k}^{-} \mathbf{S}_{k}^{T} + \mathbf{R}_{k} \right)^{-1}$$
(4)

$$\widehat{\mathbf{E}}_{NO_{x},k} = \mathbf{E}_{NO_{x},k}^{-} + \mathbf{G}_{k} \left( \mathbf{C}_{NO_{x},k}^{\text{measured}} - \mathbf{C}_{NO_{x},k}^{\text{modeled}} \right)$$
(5)

$$\widehat{\mathbf{P}}_{\mathrm{NO}_{x},k} = (\mathbf{I} - \mathbf{G}_{k} \mathbf{S}_{k}) \mathbf{P}_{\mathrm{NO}_{x},k}^{-}$$
(6)

**S** represents the  $NO_2$  sensitivity to  $NO_x$  emissions. **R** is the measurement error covariance, and it relates to the uncertainties in OMI and ground  $NO_2$  measurements. In here, the uncertainty for the AQS ground  $NO_2$  measurements was set to 0.15 (US EPA, 2006) and for the NASA standard OMI  $NO_2$ , version 2, was set to 0.3 (Bucsela et al., 2013) for all diagonal elements in **R**. The error covariance (**P**) relates to the uncertainty in the  $NO_x$  emission inventory, and the uncertainty value of 2.0 (Hanna et al., 2001; Napelenok et al., 2008) was chosen here for all diagonal elements in **P**. To simplify, off-diagonal elements in **R** and **P** were set to zero, because we assume each inversion region is an independent element.

In the time update process at time step k, the inversion process predicts the emission  $(\mathbf{E}_{\mathrm{NO}_{\mathrm{x}},k+1}^{-})$  and the error covariance  $(\mathbf{P}_{\mathrm{NO}_{\mathrm{x}},k+1}^{-})$  for the measurement update process at time step k+1, based on the corrected emission  $(\widehat{\mathbf{E}}_{\mathrm{NO}_{\mathrm{x}},k})$  and error covariance  $(\widehat{\mathbf{P}}_{\mathrm{NO}_{\mathrm{x}},k})$ 

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$$\mathbf{E}_{\mathrm{NO}_{x},k+1}^{-} = \mathbf{M}_{k} \widehat{\mathbf{E}}_{\mathrm{NO}_{x},k} + \boldsymbol{\varepsilon}_{k} \tag{7}$$

$$\mathbf{P}_{\mathrm{NO}_{x},k+1}^{-} = \mathbf{M}_{k} \widehat{\mathbf{P}}_{\mathrm{NO}_{x},k} \mathbf{M}_{k}^{T} + \mathbf{Q}_{k}$$
(8)

M represents a transition matrix;  $\varepsilon$  and  $\mathbf{Q}$  are process errors which relate to errors in modeling processes, and are difficult to estimate. Since we assume the bias between modeled and measured NO<sub>2</sub> is mostly from errors in NO<sub>x</sub> emissions (Prinn, 2000; Napelenok et al., 2008),  $\varepsilon$  and  $\mathbf{Q}$  were set to zero.

CAMx-DDM (Koo et al., 2007) calculates a semi-normalized  $NO_2$  sensitivity to  $NO_x$  emissions (unitless), as shown in Eq. (9), replacing sensitivity elements in **S** in Eq. (4),

$$S_{\text{NO}_2 \text{toNO}_x} = \tilde{E}_{\text{NO}_x} \frac{\partial C_{\text{NO}_2}}{\partial E_{\text{NO}_x}} = \tilde{E}_{\text{NO}_x} \frac{\partial C_{\text{NO}_2}}{\partial ((1+x)\tilde{E}_{\text{NO}_x})} = \frac{\partial C_{\text{NO}_2}}{\partial (1+x)} = \frac{\partial C_{\text{NO}_2}}{\partial x}$$
(9)

where  $\widetilde{E}$  represents the unperturbed NO<sub>x</sub> emission field; x is the perturbation factor. Hence, in this study, the DKF inversion actually seeks the optimal perturbation factor (x) at each iteration. The inversion processes will repeat iteratively until the perturbation factor for each emission region converges within a prescribed criterion,  $\delta$  (Fig. 2), for which the value of 0.01 was chosen in this study.

#### 2.5 NO<sub>2</sub> observations

### 2.5.1 Satellite NO<sub>2</sub> measurements

The Dutch–Finnish Ozone Monitoring Instrument (OMI) aboard NASA's EOS Aura satellite, launched on 15 July 2004, is a nadir-viewing UV-Vis spectrometer that measures solar backscattered irradiance in the range of 270 nm to 500 nm. It has been utilized to retrieve atmospheric  $NO_2$  in the spectral range from 405 nm to 465 nm with spatial resolution down to scales of  $13\,\mathrm{km} \times 24\,\mathrm{km}$  at nadir view point (Levelt et al.,

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2006a,b). The EOS Aura satellite follows a Sun-synchronous polar orbit at approximately 705 km altitude with local equator crossing time around 13:40 LT (Levelt et al., 2006b; Boersma et al., 2007). In this study, the NASA standard product, version 2 (Bucsela et al., 2013) retrieval of OMI NO<sub>2</sub>, gridded at 0.1° × 0.1° resolution, was obtained from NASA Goddard Space Flight Center and mapped to the 12 km CAMx modeling domain. OMI pixels with cloud radiance fraction greater than 0.5 and sizes of more than 20 km × 63 km at swath edges were excluded in the dataset. The OMI averaging kernels (Eskes and Boersma, 2003) were interpolated into each CAMx model layer and then applied to the modeled NO2 column density (Eq. 10), to account for the influence of the a priori NO<sub>2</sub> vertical profile used in the OMI retrieval and the OMI measurement sensitivities at each altitude:

$$C_{\text{NO}_2}^{\text{modeled}} = \sum A_i \cdot X_i \tag{10}$$

where  $A_i$  is the averaging kernel at pressure level i, and  $X_i$  is the CAMx modeled partial NO<sub>2</sub> subcolumn density at the corresponding pressure level.

In order to reduce the OMI measurement uncertainties and effects from invalid data points, monthly averaged OMI NO<sub>2</sub> column densities were used in the DKF inversions.

#### Ground and other NO<sub>2</sub> measurements 2.5.2

The US EPA Air Quality System (AQS) NO<sub>2</sub> ground monitoring network data (Fig. 1) (http://www.epa.gov/ttn/airs/airsags/) were also used for inverse modeling. AQS monitors are equipped with a heated molybdenum catalytic converter that first transforms NO<sub>2</sub> to NO, and then measures the resultant NO using a chemiluminescence analyzer. NO<sub>2</sub> is then calculated by subtracting NO measured in a separate NO mode from the resultant NO (US EPA, 1975). Studies (US EPA, 1975; Demerjian, 2000; Lamsal et al., 2008) indicate that the catalytic converter also converts fractions of other reactive nitrogen species (e.g. HNO<sub>3</sub>, PAN) into NO during this measurement. Therefore, correction factors computed from CAMx modeled concentrations by the method

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of Lamsal et al. (2008) (Eq. 11) are applied before deploying the AQS NO<sub>2</sub> data in the DKF inversion:

$$CF = \frac{NO_2}{NO_2 + \sum AN + (0.95PAN) + (0.35HNO_3)}$$
(11)

In Eq. (11), \(\sumeq AN\) represents the sum of all alkyl nitrates and PAN is peroxyacetyl nitrate. The CAMx model with CB05 mechanism does not output alkyl nitrates specifically, so the difference between modeled total organic nitrates and PAN was used to represent modeled alkyl nitrates.

The NOAA P-3 aircraft NO<sub>2</sub> data (http://www.esrl.noaa.gov/csd/tropchem/ 2006TexAQS/) and the Texas Radical and Aerosol Measurement Program (TRAMP) NO<sub>2</sub> data, measured at Moody Tower (Fig. 1), (http://geossun2.geosc.uh.edu/web/ blefer/TRAMP/Final%20data/) were used to evaluate the inverse modeling results. The Moody Tower measurement site located at the University of Houston campus is approximately 70 m above the ground (Luke et al., 2010), corresponding to the CAMx modeling layer 2, with hourly NO<sub>2</sub> data available for the whole August-September episode, but no coverage for the June episode. The P-3 aircraft measurement was made from ground level to around 5000 m height with 1 s resolution, but only available on 4 days (31 August, 11 September, 13 September, and 15 September 2006) during our modeling period. Hourly averaged aircraft NO<sub>2</sub> data was used to compare with the hourly modeled NO<sub>2</sub> at corresponding grid cells. Both P-3 aircraft and Moody Tower NO<sub>2</sub> measurements were made by using a photolytic converter, and hence did not require corrections via Eq. (11).

#### 3.1 Pseudodata test for the DKF inversion with CAMx-DDM

To evaluate the performance of the DKF inversion technique, a controlled pseudodata test was performed for 10 modeling days (31 May to 9 June, and 13 August to 22 August) for each modeling episode. The 10 day averaged modeled NO $_2$  columns at 1–2 p.m. from the base case were used as pseudo-observations, and the model was rerun with NO $_x$  emissions from each region perturbed by known factors ranging from 0.5 to 2.0 (Table 1). Appling the DKF inversion successfully adjusted the perturbed NO $_x$  emissions from each region back to their base values, converging in 4 iterations (Fig. 3). The robustness of the DKF inversion was tested by varying the uncertainty parameters, which were set to 2.0 for emissions and 0.3 for observations in the initial pseudodata test. While higher levels of the emission uncertainty parameter and lower levels of the observation uncertainty parameter led to more rapid adjustments, the final results of the DKF inversion were insensitive to the assumed uncertainty parameters, and also to the off-diagonal elements in the error covariance matrix.

#### 3.2 Additional NO<sub>x</sub> emissions

Since DKF inversions scale emissions from their original levels, an appropriate a priori  $NO_x$  emission inventory is essential for obtaining reasonable results. The NASA Intercontinental Chemical Transport Experiment (INTEX-A) air quality study (Singh et al., 2006) found large discrepancies between aircraft measurements and CMAQ simulations of  $NO_2$  concentrations in the upper troposphere. Possible explanations could be upper tropospheric  $NO_x$  sources such as lightning and aircraft  $NO_x$  emissions that are often neglected in emission inventories. Missing  $NO_x$  sources in the upper troposphere may bias the inversion on the remaining emissions (Napelenok et al., 2008). At ground level, Hudman et al. (2010) found that the soil  $NO_x$  emissions estimated by the widely used Yienger and Levy method (Yienger and Levy, 1995) were underestimated

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by a factor of 2 over the United States. Therefore, in this study, the lightning and aircraft NO<sub>x</sub> emissions were added in the upper troposphere as described in the Sect. 2.2, and the soil NO<sub>x</sub> emissions were doubled from base case levels. The emission inventory with added lightning and aircraft NO<sub>x</sub>, and doubled soil NO<sub>x</sub> (hereafter referred to as the a priori emission inventory) was used for the following inversion studies. Inclusion of these NO<sub>x</sub> sources improves the performance of the model in simulating satellite observed NO<sub>2</sub> column densities, especially in the rural areas (Figs. 4c and 5c), and reduces the bias and error by around 15% (Table 3).

#### 3.3 Top-down NO<sub>x</sub> emissions using OMI NO<sub>2</sub>

#### DS inversion 3.3.1

The DS inversion method was performed with OMI NO<sub>2</sub> column densities to create topdown NO<sub>x</sub> emissions for the 12 km modeling domain. The monthly averaged (3 June to 1 July, and 16 August to 15 September) NO<sub>2</sub> column densities at 1-2 p.m. were used to calculate the ratio of OMI to CAMx (Eq. 1). The first three modeling days were discarded for both modeling episodes to avoid the influence of initial conditions. The monthly 24 h averaged NO<sub>x</sub> emissions and modeled NO<sub>2</sub> column densities were used to determine the value of the smoothing parameter, k. In this case, k equals to 2.0 for both episodes, indicating large influence of NO<sub>x</sub> emissions transported from surrounding grid cells.

Results (Table 2) show the DS inversion scales up the NO<sub>x</sub> emissions in all seven regions, leading to higher estimates of modeled NO<sub>2</sub> column densities (Figs. 4d and 5d) in most of the domain. However, especially in urban areas, the simulated NO2 column densities with inverted NO<sub>x</sub> emissions overshoot those observed by OMI. This indicates that the ability of NO<sub>v</sub> to influence its own lifetime via changes in OH radical concentrations results in significant nonlinearity between NO<sub>2</sub> concentration and NO<sub>x</sub> emission that are neglected by the DS method. Use of inverted NO<sub>x</sub> emissions does reduce bias and error in simulating OMI observed column densities, while  $R^2$  gets worse

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(Table 3), indicating no improvement in the spatial distribution. The comparisons with AQS ground measurements (Table 4) indicate that the inverted  $NO_x$  emissions actually deteriorate the simulations of ground-level  $NO_2$ , with bias and error increasing by 70 %. Similar results are shown in evaluating model performance against Moody Tower and P-3 measurements: the DS inversion increases bias and error by approximately 30 % and 20 %, respectively.

#### 3.3.2 DKF inversion

DKF inversion using the OMI NO<sub>2</sub> measurements was conducted to constrain NO<sub>x</sub> emissions from the seven designated regions. The monthly averaged (3 June to 1 July, and 16 August to 15 September) OMI and CAMx NO<sub>2</sub> column densities at 1–2 p.m. were used in the inversion. All modeling grids in the inversion area were covered by the OMI NO<sub>2</sub> measurement data. The DKF inversions were performed with 2116 data points in one time step (1–2 p.m.). The scaling factors generated by inversion for each region were applied to the NO<sub>x</sub> emission inventory hourly, since we assume that the 1-2 p.m. NO<sub>2</sub> column density is contributed by the NO<sub>x</sub> emissions from all previous hours, and the uncertainty in the bottom-up NO<sub>x</sub> emission inventory should be the same for every time step. The satellite-based DKF inversions scale a priori NO<sub>x</sub> emissions by factors ranging from 1.02 to 1.84 in almost all regions in both episodes (Table 2), adhering to the specified uncertainty range of 0.5 to 2.0. The scaling factors tend to be larger over the rural and small urban regions than over the urban DFW and HGB ozone nonattainment regions, where the inversions scale up emissions only slightly (factors of 1.02 to 1.14). It results from the inversion attempts to compensate for the large gap between higher observed than modeled NO2 over rural regions, despite varied patterns over urban grid cells. One exception occurs in the NE Texas region in the August—September episode (Table 2), which shows downward scaling (factor of 0.56). This reflects the inversion shifting emissions between NE Texas and the much larger surrounding N rural region (Fig. 1); taken together, the net scaling factor for the two regions in August-September is 1.72, consistent with the upward scaling of rural emis-

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sions throughout the two episodes. Apart from this anomaly, scaling factors for most regions were consistent across the two episodes, varying by less than 15%.

CAMx modeled NO<sub>2</sub> column densities with the inverted NO<sub>x</sub> emissions (Figs. 4e and 5e) are increased by 3-55 % in all regions, but the increments are much more moderate compared to the DS method inversion. The statistical results (Table 3) indicate that the DKF inversed NO<sub>2</sub> are closer to OMI observations than the a priori case in terms of 20% less in bias and 10% less in error, but without improvements in the spatial distribution. The DS method scales up NO<sub>x</sub> emissions more than the DKF inversion (Table 2), making the inversed NO<sub>2</sub> concentrations have slightly less bias and error (Table 3). However, the DKF inversed  $NO_2$  has better  $R^2$  than that of inversed by the DS method, indicating the DKF inversion method has better ability to retain the spatial structure of NO<sub>x</sub> emissions. Each of the inversions using OMI NO<sub>2</sub> data actually worsens the model performance in simulating ground level NO<sub>2</sub> concentrations (Table 4), since the modeled ground NO2 using the base case emission inventory had already been overestimated (Fig. 6). However, the DKF inversed NO<sub>2</sub> only increases the bias and error by 30% in simulating AQS NO<sub>2</sub> (Table 4), by 5% in simulating P-3 NO<sub>2</sub> (Table 5), and by 8% in simulating Moody Tower NO<sub>2</sub>, whereas greater deterioration resulted from the DS inversion.

# Top-down NO<sub>x</sub> emissions using ground AQS NO<sub>2</sub>

Ground-level AQS NO<sub>2</sub> measurements were also used to drive DKF inversions of NO<sub>x</sub> emissions for the two modeling episodes. There are 37 ground measurement sites in the designated inversion regions (Fig. 1), mostly located in the urban cores. The N rural and S rural regions were excluded in this case because they contain too few measurement sites. Correction factors from Eq. (11) were applied to the ground NO<sub>2</sub> before using the data in the inversion.

The base case simulations strongly overpredicted observed NO<sub>2</sub> in the early morning and late afternoon during both modeling episodes (Fig. 6), when the model may underestimate planetary boundary layer (PBL) heights (Kolling et al., 2013). To allevi-

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ate the influence from PBL heights, daily 24 h averaged NO<sub>2</sub> levels were used in the inversions.

To address the overprediction of ground-level NO<sub>2</sub>, the ground-based inversions sharply reduce a priori NO<sub>x</sub> emissions by applying scaling factors of 0.30 to 0.57 (Table 2). The reductions in NO<sub>x</sub> emissions reduce model error relative to the AQS (Table 4) and Moody Tower NO2 observations on an hourly basis, as well as NO2 observed by the P-3 aircraft (Table 5), but may be too sharp, as they lead negative bias in predicting NO<sub>2</sub> from the AQS monitors (Table 4) and the P-3 aircraft measurements (Table 5). More moderate scaling factors are obtained if the inversion is conducted with data only from a midday window (9 a.m.-2 p.m.) when PBL heights are less problematic (not shown). However, scaling factors still remain far below 1.0 and show up to factor of two inconsistencies between the two episodes.

#### 3.5 Impacts on O<sub>3</sub> simulations

O<sub>3</sub> concentrations and their sensitivities to changes in emissions are calculated for both modeling episodes using the a priori and each of the a posteriori emission inventories. The scaled up NO<sub>x</sub> emissions from the satellite-based DKF inversion (Table 2) lead to 1-7 ppb higher modeled 8 h (10 a.m.-6 p.m.) O<sub>3</sub> concentrations over most of the domain in the June episode (Fig. 7, top row). Largest increases occur over NE Texas and N rural regions (Fig. 1), where the a priori simulation shows O<sub>3</sub> to be most sensitive to NO<sub>x</sub> (Fig. 7, middle row) and where the satellite-based DKF inversion scaled up emissions by large amounts.

The a priori simulation shows O<sub>3</sub> to be primarily sensitive to NO<sub>x</sub> over most of the domain, but VOC-limited in the core of the Houston region and with joint sensitivity to NO<sub>x</sub> and VOC in Dallas, Austin, and San Antonio (Fig. 7, left column). The satellitebased inversion increases NO<sub>x</sub> emissions and thus shifts the O<sub>3</sub> formation chemistry toward being more VOC sensitive (Fig. 7, middle column). Over much of the domain, O<sub>3</sub> sensitivity to VOC increases by a factor of about 1.5. The slight increases in O<sub>3</sub> sensitivity to NO<sub>x</sub> occur because the semi-normalized sensitivity coefficients represent the 13, 17479–17517, 2013

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local slope of O<sub>3</sub>-emissions response scaled to a 100% change in emissions. As the satellite-based inversion scales up NO<sub>x</sub> emissions, these semi-normalized coefficients increase, even though the impacts per ton of NO<sub>x</sub> decrease.

The ground-based DKF inversion leads to O<sub>3</sub> reductions of 3–8 ppb over urban regions (Fig. 7, top right), where it scales down emissions (Table 2), and less changes over rural regions where emissions were left unchanged due to lack of NO<sub>2</sub> monitors. The reduction in urban NO<sub>x</sub> makes O<sub>3</sub> less sensitive to VOC emissions as expected (Fig. 7, bottom right). However, the impact on sensitivity to NO<sub>x</sub> is mixed. In urban areas which are transitional between NO<sub>x</sub>-limited and NO<sub>x</sub>-saturated conditions, the reduction in NO<sub>x</sub> emissions pushes the chemistry toward more NO<sub>x</sub>-limited conditions and thus increases the sensitivities. In downwind regions which are already NO<sub>x</sub>-limited, the sensitivities decline because there are now less NO<sub>x</sub> emissions contributing to the semi-normalized coefficients.

Model performance in simulating hourly AQS ground-level observations of O<sub>3</sub> indicates that the bias and error slightly worsened when each of the a posteriori inventories are used in place of the a priori inventory (Table 6). The largest deterioration comes from the DS inversion as the bias and error increase by around 10%, likely because this inversion method does not retain the spatial structure of emissions from the a priori inventory. For the other inversions, the changes in bias and error are too slight to determine if performance is meaningfully impacted.

#### **Conclusions**

Inverse modeling has been performed using either NO<sub>2</sub> column densities observed by OMI satellite or ground-level NO2 concentrations observed by AQS monitors to constrain the NO<sub>x</sub> emissions for two regulatory attainment modeling episodes in Texas. Two inversion methods, DS and DKF, are applied to the OMI NO2 data, and the DKF method is also applied to the ground-level NO<sub>2</sub> data. Pseudodata test results validate that the DKF method effectively captures known perturbations in CAMx simulations.

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Two missing NO<sub>x</sub> sources in the upper troposphere, lightning and aircraft NO<sub>x</sub> emissions, are added into the base case NO<sub>x</sub> emission inventory, contributing 14% and 6% to the total NO<sub>x</sub> emissions for the June episode, and 7% and 6% for the August– September episode, respectively. The underestimated soil NO<sub>x</sub> emissions are doubled from the base case, adding an additional 8 % NO<sub>x</sub> emission to the base case for both episodes. The additional NO<sub>x</sub> emissions increase the modeled NO<sub>2</sub> column densities mostly at rural areas and improve the inversion performance with the OMI NO<sub>2</sub>, but not with the ground  $NO_2$ .

The DS inversions tend to overshoot the OMI-observed NO<sub>2</sub> column densities since this linear inversion method ignores the nonlinear influence of NO<sub>x</sub> on its own lifetime. The iterative approach of the DKF inversion avoids this problem, but fails to substantially improve the spatial correlation of modeled and observed NO2 levels since it applies only a single scaling factor to each inversion region.

The overall tendency of the model to underpredict OMI observed NO<sub>2</sub> column densities and to overpredict AQS observed ground NO<sub>2</sub> concentrations leads to conflicting results between the inversions. It is difficult to determine which observations provide a more reliable basis for the inversions, since none of the inversions improve model performance against independent data such as aircraft-observed NO<sub>2</sub> or ground-level O<sub>3</sub> concentrations. Whether this indicates that the a priori inventory is the best available representation of NO<sub>x</sub> emissions, or that tuning of the base model led to its better performance, is impossible to determine. Nevertheless, this suggests that inverse modeling of NO<sub>x</sub> emissions should for now remain a complement to SIP modeling efforts rather than a substitute for traditional bottom-up inventories.

The AQS ground NO<sub>2</sub> measurements face limitations due to the inaccuracies of the molybdenum converter method, and because the mostly urban measurement sites may be unrepresentative of the entire region. In addition, model shortcomings in simulating PBL heights may strongly bias the inversions based on ground-level observations.

For the satellite data, several factors could explain the more spatially smeared and higher rural NO<sub>2</sub> in the satellite observations than the base model which drove the up-

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ward scaling of emissions. Our inclusion of lightning and aircraft NO<sub>x</sub> emissions and doubling of soil NO<sub>x</sub> emissions narrowed but did not eliminate the discrepancy. A higher resolution OMI NO<sub>2</sub> product (retrieved with small pixels and high resolution a priori profile) has been shown to enhance NO<sub>2</sub> column densities in urban areas and reduce them in rural areas (Russell et al., 2011), which would more closely resemble the modeled distribution. Lin et al. (2012) highlighted several uncertain model parameterizations that impact model predictions of NO<sub>2</sub> column density for a given emissions inventory. For example, lowering the rate constant of the  $NO_2$  + OH reaction to match the rate of Mollner et al. (2010) would lead to a longer NO<sub>x</sub> lifetime and reduce the gap between modeled urban and rural NO<sub>2</sub> concentrations. Henderson et al. (2011) suggested that better representation of acetone and organic nitrates in the CB05 mechanism could help address its underprediction of NO<sub>2</sub> in the remote upper troposphere. Future work could explore how combinations of these adjustments influence satellite-based inversions.

The upcoming DISCOVER-AQ campaign by NASA in fall 2013 will provide vertically resolved measurements of NO<sub>x</sub> from repeated aircraft spirals in the Houston region. This may help resolve some of the discrepancies noted here between inversions driven by ground-based and satellite-based NO<sub>2</sub> observations. The future Tropospheric Emissions: Monitoring of Pollution (TEMPO) mission, using a geostationary satellite with high spatial and temporal measurement capabilities, could provide a richer data source to drive the NO<sub>x</sub> inversions. Future work could also conduct inversions based on emission categories rather than emission regions, to explore potential errors in the emission inventory on a component rather than location basis.

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# **Table 1.** Arbitrary perturbation factors for pseudodata testing.

Source Region	Perturbation factor
HGB	1.8
DFW	0.6
BPA	1.6
NE Texas	0.7
Austin and San Antonio	1.4
N rural	1.5
S rural	0.8

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**Table 2.** Scaling factors for each region from different inversions.

Source Region			3 Jun to 1 Ju	ıl 2006		16 Aug to 15 Sep 2006				
	Base NO <sub>x</sub> emission (td <sup>-1</sup> )	Priori NO <sub>x</sub> emission <sup>a</sup> (td <sup>-1</sup> )	Scaling factor relative to priori (unitless)			Base NO <sub>x</sub> emission (td <sup>-1</sup> )	Priori NO <sub>x</sub> emission (td <sup>-1</sup> )	Scaling factor relative to priori (unitless)		
			Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion <sup>b</sup>			Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion
HGB	374	455	1.46	1.12	0.37	382	436	1.56	1.03	0.57
DFW	335	435	1.36	1.02	0.34	314	412	1.47	1.14	0.47
BPA	81	97	2.23	1.83	0.49	86	98	2.02	1.75	0.42
NE Texas	141	164	2.33	1.84	0.49	155	174	1.69	0.56	0.51
Austin and San Antonio	252	319	1.55	1.28	0.30	248	302	1.82	1.70	0.40
N rural	522	823	1.93	1.67	_	543	759	2.00	1.98	_
S rural	472	728	1.83	1.52	_	489	668	2.04	1.72	_

 $<sup>^{\</sup>rm a}$  Adds lightning and aircraft  $\rm NO_x$ , and doubled soil  $\rm NO_x$  emissions to the base case.  $^{\rm b}$  Conducted with 24 h averaged ground-level  $\rm NO_2$  data.

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**Table 3.** Performance of CAMx in simulating OMI-observed  $NO_2$  column densities.

Statistical Parameters		3	Jun to 1 Jul 200	06	16 Aug to 15 Sep 2006			
	Base case	Priori <sup>c</sup>	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Base	Priori	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion
R <sup>2</sup> NMB <sup>a</sup> NME <sup>b</sup>	0.62 -0.47 0.48	0.61 -0.30 0.32	0.42 0.087 0.22	0.54 -0.12 0.23	0.63 -0.54 0.55	0.48 -0.33 0.39	0.40 0.13 0.32	0.51 -0.12 0.28

<sup>&</sup>lt;sup>a</sup> Normalized mean bias.

b Normalized mean error.

 $<sup>^{\</sup>rm c}$  Adds lightning and aircraft NO $_{\rm x}$ , and doubled soil NO $_{\rm x}$  emissions to the base case.

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**Table 4.** Performance of CAMx in simulating AQS Ground-level NO<sub>2</sub><sup>a</sup>.

Statistical Parameters	3 Jun to 1 Jul 2006					16 Aug to 15 Sep 2006				
	Base case	Priori	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion	Base case	Priori	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion
R <sup>2</sup> NMB NME	0.55 0.82 0.96	0.54 0.91 1.03	0.49 1.66 1.71	0.51 1.30 1.37	0.52 -0.16 0.48	0.50 0.35 0.63	0.49 0.42 0.67	0.43 1.10 1.21	0.44 0.72 0.91	0.47 -0.24 0.48

<sup>&</sup>lt;sup>a</sup> Hourly AQS data was used to compare with modeled NO<sub>2</sub> at corresponding locations.

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**Table 5.** Performance of CAMx in simulating P-3 aircraft-observed NO<sub>2</sub>.

Statistical Parameters	16 Aug to 15 Sep 2006 <sup>a</sup>							
	Base case	Priori	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion			
R <sup>2</sup> NMB NME	0.23 0.10 0.99	0.23 0.10 0.99	0.25 0.46 1.24	0.22 0.15 1.02	0.22 -0.14 0.86			

<sup>&</sup>lt;sup>a</sup> Comparison available for only four days (31 August, 11 September, 13 September, and 15 September 2006).

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**Table 6.** Performance of CAMx in simulating AQS hourly ground-level O<sub>3</sub>.

Statistical Parameters	3 Jun to 1 Jul 2006					16 Aug to 15 Sep 2006				
	Priori	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion	Priori	Posteriori OMI-based DS inversion	Posteriori OMI-based DKF inversion	Posteriori Ground-based DKF inversion		
$R^2$	0.61	0.61	0.63	0.58	0.50	0.52	0.51	0.46		
NMB	0.01	0.12	0.02	0.04	0.38	0.49	0.41	0.40		
NME	0.29	0.37	0.30	0.30	0.47	0.58	0.50	0.48		

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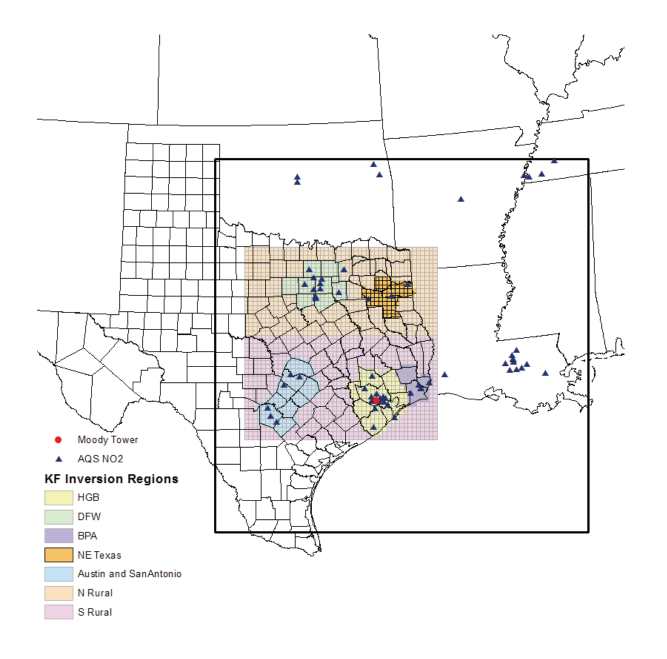


Fig. 1. 12 km CAMx modeling domain for eastern Texas (black square), inversion regions (shaded), ground AQS NO<sub>2</sub> monitoring sites (blue triangles), and Moody Tower (red circle).

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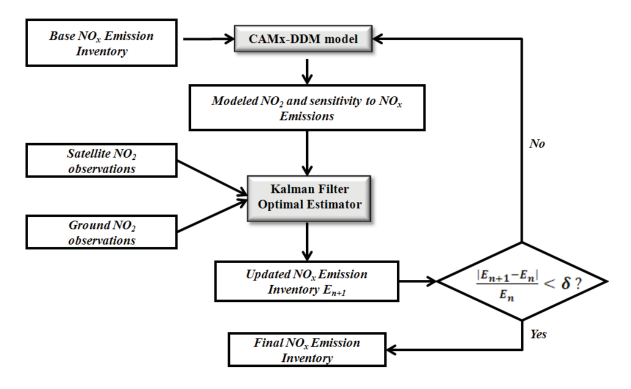
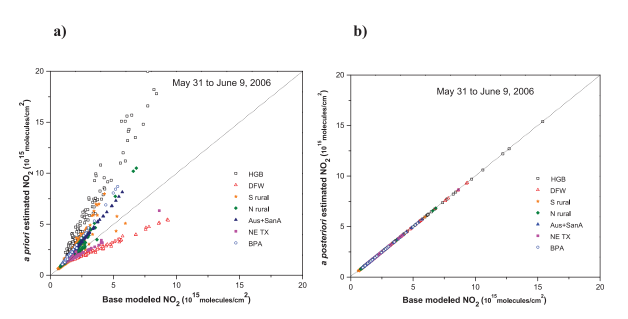


Fig. 2. Schematic diagram of Kalman filter inversion process.



**Fig. 3.** Pseudodata test showing that the DKF inversion accurately adjusts the  $NO_x$  emissions from the perturbed case (a) to the a posteriori case (b) to match the desired base  $NO_2$  column densities. Similar performance is found for the 13–22 August test period.

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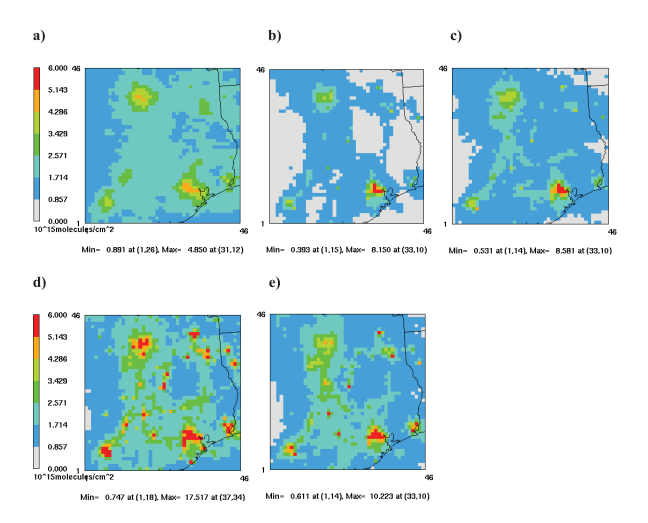


Fig. 4. Monthly averaged (3 June to 1 July) tropospheric NO<sub>2</sub> vertical columns at 1–2 p.m. from (a) OMI observations, and from CAMx simulations using (b) base case emissions inventory, (c) a priori emission inventory (with additional lightning, aircraft, and soil NO<sub>x</sub>), and OMI-based inverted NO<sub>x</sub> emissions using (d) DS and (e) DKF methods.

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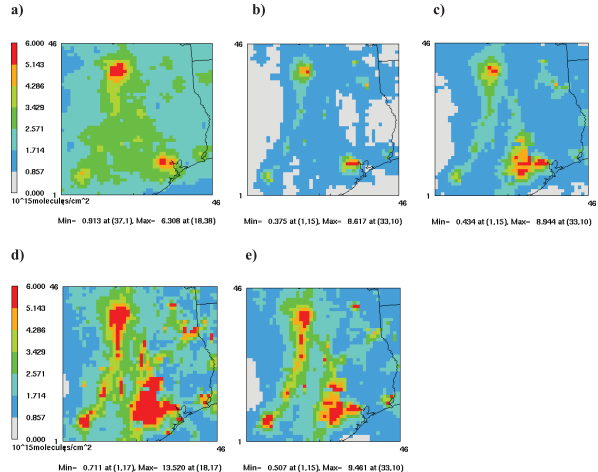


Fig. 5. Same as Fig. 4, but for the August–September episode.

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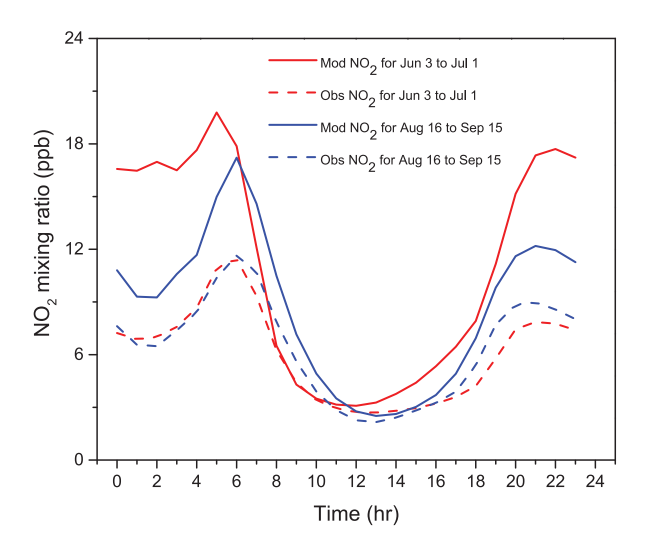


Fig. 6. Daily variations of modeled (solid line) and observed (dashed line) ground NO2 concentrations for the June (red) and August-September (blue) episodes. Note: NO2 concentrations were taken by averaging monthly data for all sites.

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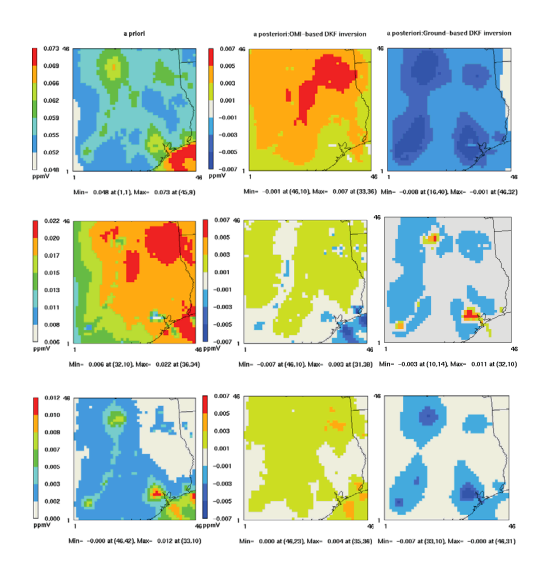


Fig. 7. Monthly 8 h (10 a.m.-6 p.m.) averaged ground-level O<sub>3</sub> concentrations (top), O<sub>3</sub> sensitivity to NO<sub>x</sub> (middle), and O<sub>3</sub> sensitivity to VOC (bottom) for the a priori case (left column), and differences (a posteriori minus a priori) for the OMI-based (middle column) and ground-based (right column) DKF inversions in the June episode. The August-September episode shows similar results.

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