Observation and modeling of the evolution of Texas power plant plumes

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Abstract

During the second Texas Air Quality Study 2006 (TexAQS II), a full range of pollutants was measured by aircraft in eastern Texas during successive transects of power plant plumes (PPPs). A regional photochemical model is applied to simulate the physical and chemical evolution of the plumes. The observations reveal that SO₂ and NOₓ were rapidly removed from PPPs on a cloudy day but not on the cloud-free days, indicating efficient aqueous processing of these compounds in clouds. The model reasonably represents observed NOₓ oxidation and PAN formation in the plumes, but fails to capture the rapid loss of SO₂ (0.37 hour⁻¹) and NOₓ (0.24 hour⁻¹) in some plumes on the cloudy day. Adjustments to the cloud liquid water content (QC)
and the default metal concentrations in the cloud module could explain some of the SO$_2$ loss. However, NO$_y$ in the model was insensitive to QC. These findings highlight cloud processing as a major challenge to atmospheric models. Model-based estimates of ozone production efficiency (OPE) in PPPs are 20-50% lower than observation-based estimates. Possible explanations for this discrepancy include the observed rapid NO$_y$ loss which biases high some observation-based OPE estimates, and the model’s under-prediction of isoprene emissions.

1 Introduction

Power plants are the leading point source emitters of SO$_2$ and oxides of nitrogen (NO$_x$=NO+NO$_2$) (EPA, 2009). The large amount of SO$_2$ and NO$_x$ emitted from power plants has been linked to a series of environmental issues, such as acid deposition, photochemical O$_3$ and particulate matter (Srivastava et al., 2004; Ryerson et al., 2001; Brock et al., 2003; Flues et al., 2002). Various regulations and market-based policies have been implemented to reduce these emissions, including the Acid Rain Program (EPA, 2005) and the NO$_x$ State Implementation Plan Call (NO$_x$ SIP Call) (EPA, 2004) in the United States. Power plants are among the most accurately measured emission sources in the U.S. national emission inventory due to direct smoke stack measurements by Continuous Emission Monitoring Systems (CEMS). Good agreement has been found in comparing power plant emissions reported by CEMS with airborne measurements of power plant plumes (PPPs) (Frost et al., 2006) and with satellite measurements of NO$_2$ (Kim et al., 2006).

The emissions, transport, and chemical evolution of pollutants from power plants have been investigated by multiple observational and modeling methods (Ryerson et al., 1998; Neuman et al., 2004; Godowitch et al., 2008; Frost et al., 2006; Kim et al., 2006; Sillman, 2000). Airborne
measurement of chemical composition and meteorological parameters in PPP transects have been conducted in several field campaigns over North America (Trainer et al., 1995; Ryerson et al., 1998; Springston et al., 2005; Neuman et al., 2009).

SO$_2$ freshly emitted from power plant stacks is quickly diluted and undergoes chemical evolution during plume transport. Previous aircraft measurements in PPPs have revealed that gas-phase SO$_2$ oxidation is the key pathway for the SO$_2$ removal and the particle growth in PPPs in the absence of clouds (Brock et al., 2002, 2003; Springston et al., 2005). SO$_2$ can also readily dissolve in cloud water and then convert to sulfate via aqueous reactions. Previous studies indicate that SO$_2$ in anthropogenic plumes has relatively long lifetime (10 hours to a few days) in the lower troposphere (Ryerson et al., 1998; Lee et al., 2011) and can undergo intercontinental transport if lifted into the middle and upper troposphere by deep convection (Fiedler et al., 2009).

Numerical simulations suggest that cloud droplet cannot effectively remove SO$_2$ plume when plume passes through or interactive with intensive clouds, which has been rarely evaluated with the observational data (Crutzen and Lawrence, 2000; Kreidenweis et al., 1997).

Several previous field studies have investigated the chemical evolution and lifetime of NO$_x$, ozone production efficiency, and the loss rate of reactive nitrogen in PPPs (Ryerson et al., 1998; Springston et al., 2005; Neuman et al., 2009). Even though numerical models have been utilized to simulate the plume chemistry and regional transport of PPPs (Frost et al., 2006; Zaveri et al., 2010), detailed model evaluation of pollutant concentrations at plume transects has been rarely done due to the scarcity of the comprehensive airborne measurement of plume pollutants. Some studies have reported the rapid loss of NO$_y$ and HNO$_3$ in PPPs (Neuman et al., 2004; Nunnermacker et al., 2000), but others have not (Ryerson et al., 2003).
In eastern Texas, power plants have significant contribution to the primary emission and high ozone concentration in the region. While the several airborne observations have been used to characterize the emissions of power plants and investigated the ozone formation in PPPs, the modeling of plume transport with 3-D photochemical models and chemical evolution and the detailed evaluation of plume simulation with the high spatial and temporal airborne observations are not available yet. In several flights during the summer 2006 Second Texas Air Quality Study (TexAQS II) (Parrish et al., 2009a), a NOAA WP-3 aircraft performed successive downwind transects of PPPs in eastern Texas. The instruments aboard the WP-3 measured a full range of chemical species, aerosol, and meteorological parameters at high time resolution and spatial resolution. This study utilizes the rich data source to examine whether a 3D photochemical model with a fine spatial resolution but without subgrid plume treatment can effectively simulate the chemical and physical evolution of PPPs as they disperse and transport downwind. We focus on the evolution of sulfur, reactive nitrogen, and O₃ in the plumes.

2 Airborne measurement

TexAQS II was a comprehensive observational campaign in eastern Texas from August to October, 2006, which aimed to improve scientific understanding of the sources and atmospheric processes responsible for the formation and distribution of O₃ and aerosols in the region (Parrish et al., 2009a). PPPs observed during TexAQS II originated from eastern Texas coal-fired power plants with a large range of reported NOₓ and SO₂ emission rates (Table 1 and Fig. 1).

The measurements and operational characteristics of the NOAA WP-3 have been summarized elsewhere (Parrish et al., 2009a). Instruments aboard the WP-3 measured numerous reactive nitrogen species (NO, NO₂, HNO₃, NO₃, N₂O₅, PAN, peroxy propionyl nitrate (PPN),
methacrylol peroxynitrate (MPAN), isoprene, CO₂, CO, SO₂, HCHO, major aerosol parameters, UV-VIS actinic flux, relative humidity, and temperature (Tables A1a and A1b of Parrish et al. (2009a) and the references therein). The instruments used in measuring major gas-phase species (O₃, CO, CO₂, SO₂, NO, NO₂, HNO₃, NOₓ, PAN, and isoprene) are summarized in Table S2 (in supplementary materials). The time resolution of most instruments was 1 second, equal to approximately 100m spatial resolution at typical WP-3 flying speeds.

Coal-fired power plants are major sources of SO₂ and NOₓ, so their plumes can be identified by elevated concentrations of SO₂ and NOₓ (Ryerson et al., 1998; Ryerson et al., 2003). SO₂ enhancement can be a more reliable diagnostic of PPPs than NOₓ since there are numerous sources of NOₓ, but coal-fired power plants are dominant sources of SO₂ in eastern Texas (Ryerson et al., 2003; Neuman et al., 2009). Background SO₂ levels were consistently below 1 ppb, so this level of SO₂ is chosen as a threshold value for identifying PPPs.

In rural areas of northeastern Texas, power plants are also leading sources of CO and anthropogenic CO₂ (Nicks et al., 2003), even though CO is not elevated in all PPPs. Airborne measurements in 2000 and 2006 showed that CO and CO₂ could be signatures of the Martin Lake, Monticello, and Welsh plumes, the concentration enhancements of which completely overlap SO₂ and NOₓ concentration enhancements at transects (Nicks et al., 2003). As the atmospheric lifetime of CO₂ is years, it is a conservative species in plumes. CO has a lifetime of one to two months in the atmosphere (Akimoto, 2003), thus serving as another conservative species in PPPs. CO emissions from Martin Lake, Monticello, and Big Brown, which were significantly underestimated in a previous emission inventory (1999), have been improved as indicated by the observed CO/CO₂ in TexAQS II (Peischl et al., 2010).
Of the 18 WP-3 flights during TexAQS II, the 16 September and 25 September flights measured successive cross-wind transects of PPPs from multiple power plants and the 19 September flight measured the Parish plume (Fig. 1 and Table S3). The 16 September flight (11:00 to 15:00 local time) observed transects of plumes from the Martin Lake, Pirkey, Monticello, and Welsh power plants at successive downwind distances (Fig. 2a). Since the three plumes transported northward through rural areas devoid of other large anthropogenic SO$_2$ sources, SO$_2$ concentration enhancements clearly denote plume locations (Fig. 2a). Pirkey is located just several km north-northeast (downwind) of Martin Lake, so their plumes cannot be distinguished on this flight after the first Martin Lake transect; we refer to the plume as Martin Lake (Ma-1) for simplicity. On 19 September, the WP-3 measured five plume transects of Parish in Houston-Galveston Brazoria (HGB) metropolitan region. On 25 September (13:00 to 16:00 local time), the WP-3 measured two plume transects of Big Brown and Limestone under northerly flow, and two plume transects of Parish in HGB (Fig. S5-S6 in supplementary materials). All transects on the three days occurred at altitudes of 600-700 m, well within the planetary boundary layer height of approximately 1500 m determined from measured temperature profiles. The exception was five transects (Ma-4 to Ma-8) of the Martin Lake plume at different heights but at the same downwind distance on 16 September.

3 Model setup and input parameters

Atmospheric chemistry for the episode was simulated by the Community Multiscale Air Quality (CMAQ) model (Byun and Schere, 2006) version 4.7 (Foley et al., 2010), using the CB05 chemical mechanism (Yarwood et al., 2005). Inline processing was applied to generate the meteorology dependent emissions properties (i.e., biogenic emissions) (Foley et al., 2010). After
accounting for plume rise, most of the power plant emissions were modeled to be released between 200 and 600 m elevation (Fig. S7).

The model was configured with 34 vertical layers and three one-way nested domains. The outer two domains cover the continental U.S. (148×112 with 36 km grid resolution) and the eastern U.S (279×240 with 12km grid resolution) including all of Texas, respectively. The rectangular frame in Fig. 1 shows the fine domain with 4km grid resolution. A full description of the modeling configuration and performance for the 12km domain can be found in Appel et al. (2009). The CMAQ modeling for the 4km domain was from September 1-25, 2006, which covers the days with WP-3 plume measurements.

Meteorology for the episode was simulated by the Fifth-Generation NCAR/Penn State Mesoscale Model (MM5) (Grell et al., 1994) version 3.7.4 for the 36 km domain. For the inner domains (12km and 4km modeling domains), the Weather Research and Forecasting Model (WRF) version 3.0 (Skamarock et al., 2008) has lower biases in simulated wind and temperature than MM5, so it was used for these domains. Thus, the meteorological field simulated by WRF has been used to drive the air quality model. Both models had 34 vertical layers extending from the surface up to 100 hPa. WRF was applied with Asymmetric Convective Model 2 PBL model (Pleim, 2007), Pleim-Xiu Land Surface Model (Xiu and Pleim, 2001), Dudhia shortwave radiation scheme (Dudhia, 1989), RRTM longwave radiation scheme (Mlawer et al., 1997), Kain-Fritsch 2 subgrid convective scheme (Kain, 2004), and the Thompson microphysics scheme (Thompson et al., 2004). MM5 used similar physical schemes. The consistency between MM5 and WRF for the modeling domains was tested and verified (Appel et al., 2009). Meteorological fields were converted to CMAQ-ready format by MCIP version 3.4.2 (Otte and Pleim, 2009).
Emission inputs for the three modeling domains were generated by Sparse Matrix Operator 1 Kernel Emissions (SMOKE) (EPA, 2006) based on the National Emission Inventory for 2005. Mobile emissions were projected to 2006 and actual CEMS data were used for point sources. BEIS3.12 (Environmental Protection Agency Biogenic Emissions Inventory System 3.12) (http://www.epa.gov/asmdnerl/biogen.html) was applied to compute the biogenic emissions.

NO$_y$ species in the CB05 chemical mechanism are NO, NO$_2$, NO$_3$, N$_2$O$_5$, HONO, HNO$_3$, PNA (peroxynitric acid), PAN (peroxyacetyl nitrate), PANX (C3 and higher peroxyacyl nitrates), and NTR (organic nitrate). The sum of all these species (with N$_2$O$_5$$\times$2) is the concentration of NO$_y$ from the model. CMAQ simulates CO but not CO$_2$.

To identify and analyze the impact of each power plant, a zero-out simulation is run with the emissions of that facility removed from the base emission inventory. The difference between concentrations in the base simulation and each zero-out simulation represents the zero-out-contribution (ZOC) of that power plant, which indicates the overall effect due to its emission and is not influenced by the nonlinear feature of plume chemistry. One base simulation and five zero-out simulations for the five power plants were performed. We focus our analyses on SO$_2$ and NO$_y$ species, whose concentration are greatly elevated in PPPs.

The aqueous processing module in CMAQ (Walcek and Taylor, 1986) processes the absorption of gas-phase species and accumulation-mode aerosols separately. The gas-phase absorption into liquid water content of clouds depends on the thermodynamic equilibrium, whereas accumulation-mode aerosols are considered to be absorbed completely into the cloud water. The dissociation of compounds into ions, oxidation of S(IV) to S(VI) by aqueous H$_2$O$_2$, O$_3$, Fe(III) and Mn(II) etc, and wet deposition are also processed in the model. For computational efficiency, CMAQ does not transport cloud-aqueous concentrations separately
from gas-phase concentrations between model grids. At the end of the cloud processing module, the cloud concentrations are removed and the mass of each species is passed to either gas-phase or aerosol concentrations.

In this study, the advection schemes used in the processing pollutant transport by the CMAQ model are Piecewise Parabolic Method (PPM) (Colella and Woodward, 1984) and Yamartino-Blackman Cubic Scheme (YAM) (Yamartino, 1993). The Asymmetric Convective Model version 2 (ACM2) (Pleim, 2007) was used to simulate the vertical mixing of pollutants in CMAQ.

4 Results and Discussion

During airborne measurement on the three days, ground temperature was 24.4-35.5°C (average: 29.0 °C) and surface wind was 0-7.2 m/s (average: 3.1m/s) at ground-based monitors in eastern Texas. At 600-700m (WP-3 typical flying height), the observed ambient temperature was 23.7-30.3°C (average: 26.8 °C), wind speed was 1.6-12.0 m/s (average: 6.4 m/s) and no precipitation was observed. The height of the planetary boundary layer (PBL), determined from the vertical profiles of equivalent potential temperature for the three days, was about 1500 m on 16 September and about 1000 m in the HGB region on 19 and 25 September.

The CEMS-reported SO₂ and NOₓ emissions of big power plants in the eastern U.S. were previously evaluated based on with WP-3 measurements of PPPs in 2004 (Frost et al., 2006). Since the emitted NOₓ in PPPs can quickly be oxidized to NO₂ (NO₂ = NOₓ-NOₓ), the observed enhancements of NOₓ and SO₂ serve as the basis for evaluation. The strong correlation between NOₓ and SO₂ for all first plume transects (R²=0.68~0.98) suggests that the power plants were the
dominant sources of these gases there. The three ratios of these plants show strong consistency within the uncertainties of the measurements, although the model slightly under-predicts SO$_2$/NO$_x$ ratios (Table 2). Likewise, previous studies have reported strong consistency between CEMS(SO$_2$/NO$_x$) and OBS(SO$_2$/NO$_y$) (Frost et al., 2006; Ryerson et al., 2003; Ryerson et al., 2001).

### 4.1 Evaluation of plume dispersion and transport

On 16 September, the WP-3 observed mostly southerly winds with average wind speeds of 6.9 m/s. The southerly winds allowed PPPs of Monticello and Welsh to remain distinct in both model and observation (Fig. 2) but caused the Martin Lake and Pirkey plumes to coincide since the two plants are just 18.5 km apart. Maximal SO$_2$ enhancements for each plume transect were used to identify the plume centers in observations and modeling results to enable comparative analyses of observations and modeling results. The plumes produced by CMAQ mostly have similar spatial extent to the measured plumes on 16 September (Fig. 2), 19 and 25 September (Fig. S5-S6). The wind speed and direction in the model were more homogeneous than observed winds, resulting in slight differences between modeled and observed locations of the plumes and plume centers (Fig. 2).

The high-resolution aircraft observations were compared with the model outputs extracted from the corresponding grid cells, adjusted to align the modeled and measured plume peak locations as necessary. Since the aircraft was flying consistently at approximately 100 m/s at each plume transect, each gridline interval in Fig. 3 (40 seconds) is equal to the spatial distance of 4 km (one grid cell).
The 16 September flight path proceeded northward in 14 successive crosswind (east-west) transects, the first 12 of which intercepted the Martin Lake (and Pirkey) plumes (Ma-1 to Ma-12 in Fig. 2; Ma-4 to Ma-8 are increasing altitudes at the same transect) and the last four of which intercepted the Monticello and Welsh plumes (Fig. 2). The extensive observation of the Martin Lake plume provides a unique opportunity to examine plume evolution from the emission stack until dilution to background levels. Comparisons between modeled and observed SO$_2$, NO$_y$, and CO mixing ratios are shown for each successive plume transect of the two days in Fig. S1-S4.

At the first transect of the Martin Lake, Monticello, and Welsh plumes, the model generates lower peak SO$_2$ and NO$_y$ concentrations and wider plumes than was observed. This likely reflects the inability of the 4-km resolution model to resolve subgrid-scale plume structure in the initial formation of a plume. No subgrid or Plume-in-Grid (PinG) was used in the modeling.

The modeled CO captured the observed extent at each plume transect, slightly underestimating the peak values (Fig. 3). The modeled SO$_2$ (18ppb) at Ma-2 matched the observed peak (23 ppb) closely as subgrid effect weakened and the plume width was larger than one grid cell. As the plume transported to Ma-3, the modeled SO$_2$ (14ppb) was higher than the observed peak (7ppb). The modeled SO$_2$ at plume center was consistently higher than the observed while the background SO$_2$ matched observations.

The measured CO at the plume center declined only from 240 (Ma-1) to 150 ppb (Ma-3). However, SO$_2$ was observed to decline by more than a factor of 10 from Ma-1 to Ma-3, indicating rapid loss.

Ma-4 through Ma-8 observed the Martin Lake plume at the same downwind distance (53 km) but flew at different altitudes (Ma-4 to Ma-8 of Fig. S1; Table S3). SO$_2$ emission from Martin Lake was modeled to occur mostly at 400 m, accounting for the stack height and plume rise (Fig.
S7). At 1800 m (Ma-4), which was near the top of the PBL, no enhancement of SO$_2$, NO$_y$, or CO was simulated but a weak SO$_2$ plume was observed, implying that the model failed to capture some of the observed upward transport. At lower flight altitudes (between 660 and 300 m, corresponding Ma-6 to Ma-8 in Fig. S1), the model effectively simulated plume extent. The comparisons between the modeled and observed SO$_2$, CO and NO$_y$ species on 19 and 25 September are shown in Fig.S1-S4.

4.2 Correlations between conservative and non-conservative species

In this section, we explore the correlation between conservative and non-conservative species from the observed plume concentrations. The correlations are presented by the slopes and R$^2$ of the least-square-fit between conservative and non-conservative species. At the time scale of PPP transport (a few hours), CO and CO$_2$ are expected to experience similar dispersion and minimal loss to chemistry or deposition, leading to near constant slopes of CO to CO$_2$. CO and CO$_2$ concentrations were strongly correlated within the Martin Lake and Monticello plumes and the slopes of CO to CO$_2$ held steady as both plume aged (Fig. 4) (for Ma-1 to Ma-3 and Ma-6 to Ma-12, slopes of the least square fit: 0.58~0.71 ppb/ppm, R$^2$: 0.89~0.96; for Mo-1 to Mo-4, slopes of the least square fit: 4.3~5.3 ppb/ppm, R$^2$:0.77~0.94), indicating the same extent of dispersion exerting on the concentration evolution of CO and CO$_2$. For the Welsh and Big Brown plumes, only the first one or two transects had the strong correlation between CO and CO$_2$, with their later transects likely affected by nearby CO or CO$_2$ emissions. Due to the strong interference from HGB urban emissions of CO, no clear correlation between CO and CO$_2$ could be found in the Parish plume.
Concentrations of SO$_2$ and reactive nitrogen species in PPPs are strongly affected by chemical reactions, heterogeneous conversion, deposition, dispersion, and cloud processing. Dispersion is expected to have the same extent of impact on both conservative (e.g., CO$_2$ and CO) and non-conservative species (e.g., SO$_2$, NO$_x$, HNO$_3$, and PAN). Thus, the variations of slopes between non-conservative and conservative species reflect the impact of plume chemistry, deposition and heterogeneous processing on non-conservative species.

Given that the observed SO$_2$ and CO$_2$ in all plumes showed strong correlations, CO$_2$ could serve well as a signature of PPPs. However, since CO$_2$ is not modeled by CMAQ, CO is selected as the conservative species for the purpose of comparison between the model and the observations. CO is a signature emission of some but not all power plants in Texas. In observed PPPs, only in the Martin Lake and Monticello plumes could the strong correlations between the non-conservative species (SO$_2$, reactive nitrogen species) and CO be found at all transects. In the first one or two transects of the Big Brown and Welsh plumes, SO$_2$ strongly correlates with CO.

### 4.3 Evaluation of SO$_2$ plume evolution

On 19 September, under the clear-sky background (Fig. S9), the normalized SO$_2$ to CO ratio from the model and the normalized SO$_2$ to CO$_2$ ratio from the observation matched closely for the Parish plume, both of which shows the slow SO$_2$ loss (Fig. 8). At the plume age of 11 hours, only 25% SO$_2$ was removed in both the modeling and the observation. Thus, the model can capture SO$_2$ evolution when no cloud processing occurs.

However, plume observations demonstrate rapid loss of SO$_2$ in the 16 September plumes (Martin Lake, Monticello and Welsh) (Fig. 5). For the Martin Lake transects, the decreasing trend of SO$_2$/CO fits to an exponential function with a first-order loss rate of 0.38 hour$^{-1}$, the
inverse of which is a lifetime of 2.6 hours ($R^2=0.94$) (Fig. 5). SO$_2$/CO from the model decreases far slower as plume ages with a loss rate of 0.016 hour$^{-1}$ (lifetime of 62.5 hours), which suggests the model significantly underestimates SO$_2$ loss for the Martin Lake plume. Similarly, for the Monticello plume, the curve fit of observed SO$_2$/CO indicates an SO$_2$ lifetime of 2.7 hours (loss rate of 0.37 hour$^{-1}$) compared to a modeled SO$_2$ lifetime of 17.2 hours. Although SO$_2$ and CO were not strongly correlated in observations of the other plumes, diminishing SO$_2$/CO$_2$ ratios indicate that rapid SO$_2$ loss also occurred in the Welsh plume, but not in the Big Brown and Parish plumes during cloud-free days (Fig. 5 and 8).

The lifetime of SO$_2$ against gas-phase oxidation by OH is a few days to one week, and SO$_2$ lifetime against dry deposition approximates one day in the boundary layer. Thus, gas-phase oxidation and dry deposition are insufficient to explain the rapid loss of SO$_2$ in the 16 September plumes. The observed rapid loss of SO$_2$ in the 16 September plumes may indicate aqueous-processing by the scattered clouds that were presented on this day. In clouds, SO$_2$ can be substantially dissolved in water droplets and subsequently be oxidized to form H$_2$SO$_4$ in cloud water far more rapidly than the gas-phase process.

Observational data indicates scattered cloudiness on 16 September and clear skies on 19 and 25 September. MODIS images of cloud and aerosol optical depth (http://ladsweb.nascom.nasa.gov/browse_images/) show that there were scattered clouds over eastern Texas on 16 September and it was a clear-sky and sunny day on 19 and 25 September. On 16 September, the measured photolysis rates of NO$_2$ and NO$_3$ oscillated by a factor of 2 during the aircraft measurement, in contrast to less variability on 19 and 25 September (Fig. S8). The cloudy meteorological condition on 16 September, indicated by the variations of photolysis rates, agrees with scattered clouds captured by the satellite cloud image (Fig. S9). The relative
humidity measured from Ma-4 to Ma-8, when the WP-3 descended from 1800 to 300 m at the same downwind distance of Martin Lake, reached saturation between 1800~1000 m, implying clouds distributed at that altitude and potentially interacting with the plume (Ma-4 and Ma-5 in Fig. S1).

The model successfully simulated the distribution of scattered clouds over northeastern Texas on 16 September indicated by a MODIS cloud image (Fig. S9 and S10), but placed them predominately between 2500 and 4000 m altitude as indicated by the cloud bottom height (CLDB) in the model (Fig. S10), well above the PPPs which resided under the PBL (~1000-1500 m) as shown in the vertical distribution of plume SO$_2$ for Martin Lake, Monticello, and Welsh plumes (Fig. S11-S12). The liquid water mixing ratio (QC) from the layer 10 and above in the model was zero. Thus, no significant cloud processing was modeled to occur in the base modeling, so the observed rapid loss of SO$_2$ in the 16 September plumes could not be replicated.

Could CMAQ have simulated the rapid SO$_2$ loss on 16 September if the meteorological model had placed the clouds at lower altitudes in contact with the PPPs? The cloud module in CMAQ includes two mechanisms for removing pollutants: aqueous chemical reactions and scavenging and wet deposition. SO$_2$ absorption into cloud droplets and subsequent oxidation are explicitly represented. The absorption is governed by the thermodynamic equilibrium. The aqueous S(IV) is then oxidized to S(VI) by H$_2$O$_2$, O$_3$, metal ions (Fe(III) and Mn(II)), and methylhydroperoxide (MHP), and peroxyacetic acid (PAA). Since no precipitation was observed during the airborne measurements, pollutants were not expected to be scavenged.

Cloud parameters of meteorological inputs are perturbed to diagnose how efficiently pollutants such as SO$_2$ and NO$_y$ are removed from plumes. Specifically, the cloud bottom height in the meteorological field on 16 September is adjusted to 1000 meters so that the plumes
interact with clouds during their transport. Liquid water content QC is the cloud parameter determining the extent of the pollutant aqueous processing. The cloud aqueous module can be executed only if QC is larger than 0.01 g/kg. Initially, QC over northeastern Texas in the base model is zero and the maximum QC over the entire domain is about 0.4 g/kg. In the perturbation cases, we uniformly increase QC to the levels from modest to strong cloudy conditions (from 700 m to 1000 m over northeastern Texas, see Fig. S13). In the first perturbation case (denoted QC_0.05), QC is set to 0.05 g/kg = 0.05 g/m^3, equivalent to fog, a modest cloud. Then QC is increased to 0.5 g/kg = 0.5 g/m^3, equivalent to stratocumulus clouds) and 5 g/kg = 5 g/m^3, equivalent to cumulonimbus clouds), respectively, representing extremely cloudy conditions.

In the base modeling for the Martin Lake plume, only 11% SO_2 is removed in the model (the normalized SO_2/CO decreased to 0.89 from Ma-1 to Ma-12). In the QC_0.05 case, 25% of SO_2 is removed during that span (SO_2/CO decreases from 0.344 (Ma-1) to 0.257 (Ma-12)), far short of the observed 92% SO_2 removal (SO_2/CO decreased from 0.480 (Ma-1) to 0.041 (Ma-12) in observations) (Fig. 5). The cloudier scenarios yield 66% (QC=0.5 g/kg) and 81% (QC=5.0 g/kg) SO_2 removal, still below the observed rate.

Four S(IV) oxidation reactions are explicitly implemented in the cloud aqueous module, i.e. H_2O_2, O_3, metal (Fe(III) and Mn(II)), MHP, and PAA oxidations. In QC_0.05, S(IV) oxidation is dominated by H_2O_2 oxidation, with 96.2% of S(IV) oxidation occurring by H_2O_2 in the Martin Lake plume. Only about 1.7% of S(IV) oxidation was by the metal ions.

In the default CMAQ cloud module, Fe(III) and Mn(II) are uniformly set to 0.01 and 0.005 ug/m^3 over the domain. These values are assumed to represent the aqueous metal ion concentrations in the background atmosphere. However, power plants are major emission sources of particulate metals (Alexander et al., 2009). Fe(III) and Mn(II) in the power plant
plumes are expected to be higher than the background levels, thus potentially enhancing the aqueous oxidation of sulfur in PPPs. In another perturbation case, both Fe(III) and Mn(II) concentrations are increased by a factor of 10 with QC setting to 0.05 g/kg (called QC_METAL hereafter). The increase of Fe(III) and Mn(II) is within the range of metal ion concentrations measured in fogs and cloud water (Raja et al., 2005; Parazols et al., 2006). SO$_2$ removal in QC_METAL was more rapid than that of QC_0.05 (Fig. 5). At the last plume transect, SO$_2$ decreased by 33%, compared to the 25% SO$_2$ removal in QC_0.05, suggesting the increased metals in plume lead to more SO$_2$ removed in cloud water. Thus, some combination of enhancements in cloud liquid water content and metals concentrations may help explain the observed rapid SO$_2$ loss rates in the cloudy day plumes.

Few studies have observed rapid SO$_2$ loss in anthropogenic plumes, though similar rates of SO$_2$ loss have been found in volcanic plumes (Oppenheimer et al., 2010; Rodríguez et al., 2008). These studies proposed that cloud aqueous processing is the mechanism for the rapid SO$_2$ removal. The comprehensive airborne measurement of plume concentrations and meteorological parameters supported by satellite images in this study confirms that the cloud processing caused the rapid SO$_2$ loss. SO$_2$ taken up by cloud droplet and subsequent aqueous oxidation, as a complex process affected by QC, droplet pH, and the oxidant concentrations, and catalysis in the droplets etc, are a major challenge to models. Earlier studies have also found that models can underestimate SO$_2$ loss rates in clouds (Crutzen and Lawrence, 2000; Kreidenweis et al., 1997).

### 4.4 Evaluation of plume chemistry of reactive nitrogen

In PPPs, HNO$_3$, NO$_3$, N$_2$O$_5$, PAN, and other organic nitrates are formed via the NO$_x$ chemical reactions. Freshly emitted NO$_x$ titrates O$_3$ and consumes OH, resulting in slow
formation of HNO₃ and no formation of PAN in the initial plume (Karamchandani et al., 1998).

As a plume dilutes, OH levels recover and HNO₃ and other products form from NOₓ oxidation.

Previous daytime observations of PPPs concluded that HNO₃ and PAN were the major (more than 90%) products of NOₓ oxidation in PPPs (Neuman et al., 2006; Neuman et al., 2004; Ryerson et al., 2003; Ryerson et al., 2001). The observational data in this study also show that HNO₃ and PAN were the only two major oxidation products in PPPs, with NO₃ and N₂O₅ and other organic nitrates at least one order of magnitude lower in plume transects.

The measured and modeled NOₓ, HNO₃, and PAN are shown for comparison in Fig. S2 and S3. NOₓ was higher than HNO₃ until the plume transported 2.0 hours at Ma-7 and Ma-8. The model generally captured the observed evolution of reactive nitrogen species NOₓ, HNO₃, and PAN in the plume, simulating the transition from NOₓ to HNO₃ dominance and approximately matching the observed PAN levels. However, the simulated HNO₃ concentrations were higher than observed, implying over-prediction of HNO₃ formation or under-prediction of HNO₃ loss.

The oxidation of NOₓ by radicals approximates as a first-order reaction if radical concentrations are assumed to be constant in the plume. The observed NOₓ/CO fits to an exponential decay function (for Martin Lake, $R^2=0.85$; for Monticello, $R^2=0.86$; Fig. 6), corresponding to NOₓ lifetimes of 2.6 and 1.2 hours for the Martin Lake and Monticello plumes, respectively. The NOₓ lifetimes computed here are consistent with the NOₓ lifetimes (1.0~1.6 hours) estimated for both plants in TexAQS 2000 (Neuman et al., 2004). The declining trends of NOₓ/CO from the model and the observation closely match in the Martin Lake and Monticello plumes, with discrepancies only in the initial transects due to the inability of the model to resolve subgrid-scale plume structure (Fig. 6).
The ratios of \( \text{HNO}_3/\text{CO} \) and \( \text{PAN}/\text{CO} \) are compared between the model and observations to explore chemical evolution in the Martin Lake and Monticello plumes. We find that the model captures the PAN formation very well, closely matching observed trends as the plumes age (Fig. 6 and Fig. S2). The modeled \( \text{HNO}_3/\text{CO} \), however, was 0.7~6.6 times larger than observed. Given the good agreement between the modeled and observed \( \text{NO}_x \) oxidation and PAN formation, the \( \text{HNO}_3 \) gap between the model and the observation on the cloudy day implies that \( \text{HNO}_3 \), while being formed during plume transport, was rapidly removed from the atmosphere, which is not captured by the model.

Unexpectedly rapid loss of \( \text{NO}_y \) has also been reported by some measurement studies of biomass burning (Takegawa et al., 2003) and PPPs (Neuman et al., 2004), but not in others (Ryerson et al., 2003). When \( \text{NO}_x \) is oxidized to other reactive nitrogen species, the reactive nitrogen may be removed from the atmosphere via rain scavenging, dry deposition, heterogeneous conversion to aerosol, and cloud processing, resulting in the loss of \( \text{NO}_y \). Assuming a first-order decline of \( \text{NO}_y/\text{CO} \) (Fig. 7), the observed \( \text{NO}_y \) loss rate was 0.15 hour\(^{-1}\) for the Martin Lake plume whereas the modeled \( \text{NO}_y \) loss rate was lower by a factor of 6 (0.026 hour\(^{-1}\)). For the Monticello plume, the observed \( \text{NO}_y \) loss rate (0.24 hour\(^{-1}\)) was 2.3 times the modeled. The observed \( \text{NO}_y/\text{CO}_2 \) in Martin Lake, Monticello, and Welsh plumes had the similar extent of \( \text{NO}_y \) loss, especially during the early plume age (≤2 hours) when \( \text{NO}_y/\text{CO} \) declined by 40~50\% (Fig. 7).

On 19 September, a cloud-free day, the model effectively simulates the observed slow removal of \( \text{NO}_y \) (Fig 8). \( \text{NO}_y \) loss on the cloudy day likely reflects deposition of highly soluble \( \text{HNO}_3 \), since the other main \( \text{NO}_y \) constituents (\( \text{NO}_x \) and PAN) have low water solubility, cannot directly convert to aerosol, and have negligible dry deposition in plume. \( \text{NO}_x \) oxidation and
thermal decomposition of PAN do not shift the gas-phase NO\textsubscript{y} budget since their products are also gas-phase NO\textsubscript{y} constituents. The measured NO\textsubscript{3}\textsuperscript{-} was minor in the inorganic aerosol composition, indicating that the loss of HNO\textsubscript{3} to aerosol-NO\textsubscript{3}\textsuperscript{-} was negligible under the high ambient temperatures (the measured average temperature was 28.9°C) and the lack of ammonia enhancement beyond levels needed to neutralize the sulfate in the PPPs (Nowak et al., 2010). Given that no wet precipitation was reported on the flight days, no rain scavenging is expected to have occurred. HNO\textsubscript{3} may have rapidly dissolved in cloud droplets if the plume interacted with a cloud, as is possible under the 16 September scattered cloudiness conditions discussed earlier.

In contrast to the SO\textsubscript{2} results, the cloud perturbation scenarios did not significantly impact modeled concentrations of NO\textsubscript{y} species. Among NO\textsubscript{y} species, HNO\textsubscript{3} is the only one to be processed by the cloud module. Even raising QC to 5.0 g/kg, there is no scavenging removal of HNO\textsubscript{3} since no wet deposition happens in the absence of precipitation. At the end of the cloud module, the aqueous concentration of HNO\textsubscript{3} in cloud is passed to either gas-phase or aerosol species. Also, the ratio HNO\textsubscript{3} to total NO\textsubscript{y} is assumed to be constant. Thus, in the cloud aqueous modeling, HNO\textsubscript{3} is expected to be insensitive to QC.

4.5 Evaluation of O\textsubscript{3} simulation in PPPs

Various numerical models have been applied to simulate the O\textsubscript{3} chemistry of PPPs (Sillman, 2000; Springston et al., 2005; Frost et al., 2006; Zaveri et al., 2010). The simulation of 3-D models are as the most widely used tool to assess the effectiveness of emission controls of power plant pollutants while the model performance has been merely examined with ground concentrations (Mauzerall et al., 2005; Vijayaraghavan et al., 2009; Godowitch et al., 2008a; Godowitch et al., 2008b). In this study, the simulated plume concentration and evolution which essentially happens
at several hundred meters above ground have been compared to those aircraft measurement at plume transects. In this section, the O₃ concentration and OPE are compared with the observed at each plume transect in detail. The model overestimated background O₃ by 8–15 ppb during the flights (Table S4). Sensitivity modeling shows that boundary conditions were the biggest contributor to background O₃ levels. Thus, we focus on the differences (ΔO₃) between plume and background O₃ mixing ratios to assess model performance for O₃ formation from power plant plumes (Table S4).

The model accurately simulates that the Monticello and Welsh plumes shift from being depleted to being enriched in O₃ between transect 1 and 2, and predicts the transition to occur one transect sooner than observed for Martin Lake. All of these plumes traversed rural regions of northeastern Texas where biogenic isoprene is abundant. However, the model underestimates the amount of O₃ enrichment downwind by 20–70% (Ma-9 to Ma-12, Mo-2 to Mo-4, We-2 to We-4). The model also underestimates titration in the initial transects, reflecting the more rapid dilution of NOₓ in the model.

OPE illustrates the number of O₃ molecules formed per molecule of NOₓ irreversibly oxidized to NO₂ species (Liu et al., 1987). Box and 2D Lagrangian models driven by the observational data have previously computed OPE of pollution plumes and at ground-based monitors (Sillman, 2000; Zaveri et al., 2003). Three dimensional global models have been applied to calculate the global and regional OPE averaged at coarse scale (Fang et al., 2010; Hudman et al., 2009). It is more relevant to apply the regional 3D model at fine resolution in deriving the ozone sensitivities and OPE for regional and urban air quality strategies. OPE from 3D regional models, however, has rarely been evaluated with the observation-based results at the plume-transect scale due to the scarcity of measurements (Yu et al., 2010; Godowitch et al., 2008b). This
study computes the OPE at each plume transect from the model and then compares it with the corresponding observational results. In the model, OPE is determined from the ratio ZOC$_{O_3}$ to ZOC$_{NO_x}$. The observation-based OPE is typically derived from the least square slope of O$_3$ versus NO$_y$-NO$_x$ (NO$_z$) (Trainer et al., 1993; Kleinman et al., 2002; Griffin et al., 2004; Ryerson et al., 2003).

For Martin Lake, Monticello, and Welsh, while O$_3$ production evolves from being depleted to being formed, OPE exhibits a steady increase, consistent with OPE trends from PPPs in the southeast U.S. (Ryerson et al., 2001) and in Texas in 2000 (Springston et al., 2005; Ryerson et al., 2003). OPEs from Martin Lake (Ma-6), Monticello (Mo-4), and Welsh (We-4) plumes at similar plume ages are compared in Fig. 9. OPEs for Monticello and Welsh were remarkably similar (Fig. 9), reflecting approximately equal O$_3$ formation potentials of these facilities with similar NO$_x$ emission rates (Table 1). Martin Lake emitted about two times as much NO$_x$ as Monticello and Welsh, and thus exhibited a smaller OPE (7.25). OPE in the Big Brown plume (Bi-1) was 1.7 at a plume age of 1.3 hours, lower than the similar-plume-age OPE of Martin Lake (2.6, Ma-3) and Welsh (4.6, We-2), but close to the OPE of Monticello (1.4, Mo-2). OPE could not be quantified in the subsequent Big Brown transect due to lack of correlation between O$_3$ and NO$_z$.

The Parish plume exhibited an OPE of 4.4 at a plume age of just 0.6 hours, suggesting rapid O$_3$ formation under the influence of Houston region anthropogenic VOCs.

For the Martin Lake, Monticello, and Welsh plumes, the modeled OPEs steadily increase from O$_3$ titration (negative OPE) to rapid O$_3$ formation, showing similar trends to the observed OPEs. The modeled maximum OPEs are systematically about a factor of 2 lower than the observed for these plumes likely due to the rapid loss of NO$_y$ observed during the cloudy day (Table S4). Since the definition of OPE implicitly assumes that NO$_y$ is conservative in plumes,
the accuracy of the observation-based OPE may be undermined due to the rapid loss of NOy in this study.

It should also be noted that the model tended to under-predict measured isoprene concentrations. Observed isoprene concentrations averaged over all transects, is higher than the modeled average by 51.3%. We perturb domain-wide isoprene emission rates by this factor in the model to investigate how much impact the isoprene discrepancy has on the O3 formation in plumes. After perturbation, the simulated ZOC O3 has a maximum increase of 3 ppb (Ma-2). The maximum O3 increase for Monticello and Welsh is 1.5 and 2 ppb, respectively. OPEs of Martin Lake, Monticello, and Welsh plumes would increase to 7.0, 7.8, and 5.9, respectively, closing roughly half of the gap between modeled and observed OPEs. For Big Brown, the OPE would increase by a factor of 1.4, and for Parish, the OPE would increase to 6 (Pa-2), exceeding the observed OPE (4.4).

5 Discussion and Conclusions

A regional 3D photochemical model was applied with fine-grid resolution to simulate PPPs during three days of airborne measurement by NOAA’s WP-3 aircraft in TexAQS II. In comprehensive evaluation of the model performance, the modeled and airborne observed concentrations are compared in detail at each plume transect, which has rarely been done due to the scarcity of the airborne observation of PPPs. Under steady wind meteorological conditions, the fine-scale (4km) CMAQ demonstrated its ability to simulate the transport and dispersion of PPPs despite lacking a plume-in-grid module.
SO$_2$ and NO$_x$ show strong consistencies among the CEMS-reported emission data. In the Martin Lake and Monticello plumes, CO was strongly correlated with SO$_2$ and NO$_y$ and could serve as a conservative tracer species to track plume evolution; CO$_2$ was strongly correlated with SO$_2$ and NO$_y$ in all plumes but was not modeled by CMAQ. The trend in the least square slopes of pollutants relative to CO (CO$_2$) was used to assess species lifetime.

On clear-sky days (19 and 25 September), SO$_2$ and NO$_y$ experienced slow evolution (loss) in the Parish and Big Brown plumes. Both the model and the observation were closely correlated in the ratios of SO$_2$ and NO$_y$ to conservative species, suggesting the model well captured SO$_2$ and NO$_y$ evolution in the plumes.

SO$_2$ was observed to be rapidly lost in the Martin Lake, Monticello, and Welsh plumes under scattered cloudiness on 16 September. The observation-based SO$_2$ lifetime was 2.6 and 2.7 hours for the Martin Lake and Monticello plumes, respectively. The detailed examination of the photolysis rate and relative humidity data suggested cloud-processing of PPPs caused the rapid SO$_2$ loss on 16 September. The original simulation did not show the apparent SO$_2$ loss since PPPs resided below clouds in the model. Perturbing the cloud bottom heights to interact with the PPPs yielded modest rates of SO$_2$ removal via aqueous processing in the CMAQ cloud module. SO$_2$ removal in the model was still slower than the observed rapid loss, even after increasing cloud liquid water content and metals concentrations in cloud droplets to enhance SO$_2$ oxidation.

The simulation closely matched the observed NO$_x$ oxidation rates. The observed NO$_x$ lifetime for Martin Lake and Monticello plumes was 2.6 hours and 1.2 hours, respectively. The modeled PAN formation reflected the observed trend of PAN formation, while the modeled HNO$_3$ was 0.7~6.6 times higher than observed due to the rapid HNO$_3$ loss in observation on cloudy days. Martin Lake, Monticello, and Welsh plumes showed the similar extent of NO$_y$ loss.
For the Martin Lake plume, the loss rate of NO\textsubscript{y} has been quantified to be 0.148 hour\textsuperscript{-1} in observation, faster than the modeled NO\textsubscript{y} (0.026 hour\textsuperscript{-1}) by a factor of 6. In the model, NO\textsubscript{y} loss was insensitive to the aqueous processing when there was no precipitation happening. This study, together with modeling studies, shows that the numerical representation of cloud aqueous processing remains a major challenge.

The model effectively simulated the transition between ozone titration and formation but tended to under-predict the magnitude of O\textsubscript{3} production and the OPE indicated by observations. The discrepancies of OPEs between the model and the observations could be explained by the observed rapid NO\textsubscript{y} loss that biases high the observation-based OPE estimates, or under-prediction of isoprene emissions that leads the model to under-predict OPE.

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Table 1. Major power plants in eastern Texas

<table>
<thead>
<tr>
<th>Facility</th>
<th>NO\textsubscript{x} emission rate \textsuperscript{a} (tons/hour)</th>
<th>SO\textsubscript{2} emission rate \textsuperscript{a} (tons/hour)</th>
<th>Stack Height (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin Lake</td>
<td>2.02</td>
<td>10.37</td>
<td>138</td>
</tr>
<tr>
<td>Monticello</td>
<td>1.34</td>
<td>5.49</td>
<td>128</td>
</tr>
<tr>
<td>Welsh</td>
<td>0.95</td>
<td>2.21</td>
<td>172</td>
</tr>
<tr>
<td>Pirkey</td>
<td>0.58</td>
<td>0.21</td>
<td>160</td>
</tr>
<tr>
<td>Big Brown</td>
<td>0.84</td>
<td>13.09</td>
<td>122</td>
</tr>
<tr>
<td>Parish</td>
<td>0.33</td>
<td>2.74</td>
<td>183</td>
</tr>
<tr>
<td>Limestone</td>
<td>0.79</td>
<td>0.63</td>
<td>137</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Emission rate is the hourly averaged CEMS data for Martin Lake, Monticello, and Welsh on 16 September, 2006, and for Parish, Big Brown, and Limestone on 25 September, 2006.
Table 2. CEMS-reported E(SO$_2$)/E(NO$_x$) emission molar ratio, the observed SO$_2$/NO$_y$ and the modeled ZOC$_{SO2}$/ZOC$_{NOy}$ at the location of the first plume transect.

<table>
<thead>
<tr>
<th>Plant</th>
<th>CEMS SO$_2$/NO$_x$</th>
<th>OBS SO$_2$/NO$_y$</th>
<th>MODEL ZOC$<em>{SO2}$/ZOC$</em>{NOy}$</th>
<th>Plume age (hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martin Lake</td>
<td>3.05</td>
<td>3.94 (0.98)</td>
<td>3.30</td>
<td>0.7</td>
</tr>
<tr>
<td>Monticello</td>
<td>2.04</td>
<td>3.00 (0.86)</td>
<td>1.84</td>
<td>0.3</td>
</tr>
<tr>
<td>Welsh</td>
<td>1.10</td>
<td>1.20 (0.86)</td>
<td>1.08</td>
<td>0.4</td>
</tr>
<tr>
<td>Big Brown</td>
<td>8.94</td>
<td>10.95 (0.97)</td>
<td>9.73</td>
<td>1.3</td>
</tr>
<tr>
<td>Parish</td>
<td>5.28</td>
<td>6.83 (0.68)</td>
<td>5.18</td>
<td>0.6</td>
</tr>
</tbody>
</table>

$^a$ the values in brackets are the R$^2$ of least square fit of SO$_2$ versus NO$_y$

$^b$ ZOC$_{SO2}$ = SO$_2$ model, base - SO$_2$ model, zero-out that plant, ZOC$_{NOy}$ = NO$_y$ model, base - NO$_y$ model, zero-out that plant
Fig. 1. WP-3 flight tracks (16 September in blue, 19 September in yellow, and 25 September in pink,) and power plants in eastern Texas. NO\textsubscript{x} emission rates are shown by colors and SO\textsubscript{2} emission rates are indicated by size of circles. Rectangular frame shows the 4km modeling domain. Black stars are all other point sources in Texas. The Houston-Galveston-Brazoria (HGB) and Dallas-Forth-Worth (DFW) metropolitan areas are also shown.
Fig. 2. (a) Observed PPPs of Martin Lake, Monticello, Pirkey, and Welsh on 16 September, 2006. The black dots show the locations of the power plants. PPPs are identified by measured SO$_2$ enhancement (color gradient in the figure), as outlined by the green dash lines. Measured wind vectors are presented on the plume transect. (b) Simulated PPPs of Martin Lake, Monticello, Pirkey, and Welsh at 18:00 GMT (600~700 m) (local time: 12:00)
Fig. 3. The comparison of the modeled and observed SO$_2$, NO$_y$, and CO at plume transect of Ma-1, Ma-2, Ma-3 and Ma-6. The modeled SO$_2$, NO$_y$, and CO are labeled as yellow, green, and purple flat lines, respectively. The observed SO$_2$, NO$_y$, and CO are labeled as blue, red and black dots. The Horizontal coordinate is time scale in GMT (local time = GMT - 6 hours) and vertical coordinate is concentration (ppb). Transect names listed in Table S3 of the manuscript are labeled in each subplot. Fig. S1-S4 summarize the comparisons for all the plume transects.
Fig. 4. Scatter plot of CO (ppb) versus CO$_2$ (ppm) from plume transects (a) Martin Lake (Ma-1 to Ma-12), and (b) Monticello (Mo-1 to Mo-4). The unit of the slopes from the least square fits is ppb/ppm.
Fig. 5. (a) The observed least square slopes of SO$_2$ to CO (red square) and modeled ZOC$_{SO2}$/ZOC$_{CO}$ (blue diamond for the base case, green dot for the adjusted cloud case) as a function of plume age at each transect of the Martin Lake plume (16 September). The observed SO$_2$ loss rate was 0.38 hour$^{-1}$ (R$^2$=0.94, SO$_2$ lifetime: 2.6 hours); the modeled SO$_2$ loss rate was 0.016 hour$^{-1}$ (R$^2$=0.36, SO$_2$ lifetime: 62.5 hours). The SO$_2$ to CO slopes for each perturbation case are also plotted accordingly. (b) The observed least square slopes of SO$_2$ to CO$_2$ (ppb/ppm) for the five plumes; Martin Lake, Welsh, and Monticello plumes were observed on September 16 (cloudy day), Big Brown and Parish plumes were made on 25 September (sunny day).
Fig. 6. Observed (yellow circle) and modeled (green triangle) NO$_x$/CO (a) for the Martin Lake plume, (b) for the Monticello plume. The observed NO$_x$ oxidation rate was 0.38 hour$^{-1}$ ($R^2=0.85$) for the Martin Lake plume and 0.84 hour$^{-1}$ ($R^2=0.86$) for the Monticello plume. Observed and modeled PAN/CO and HNO$_3$/CO, (c) for the Martin Lake plume, (d) for the Monticello plume.
Fig. 7. (a) The observed least square slopes of NO$_y$ to CO (red square) and modeled ZOC$_{NOy}$/ZOC$_{CO}$ (orange diamond for the base case, green dot for the adjusted cloud case) as a function of plume age at each transect of the Martin Lake plume; the observed NO$_y$ loss rate was 0.145 hour$^{-1}$ ($R^2=0.69$) and the modeled NO$_y$ loss rate was 0.026 hour$^{-1}$ ($R^2=0.48$). (b) The observed least square slopes of NO$_y$ to CO$_2$ (ppb/ppm) for the Martin Lake, Monticello, and Welsh. NO$_y$ in the least-square fits was directly measured and not the sum of measured reactive nitrogen species.
Fig. 8. The least-square-fit slopes of SO$_2$ to CO$_2$ (from the observation) and SO$_2$ to CO (from the model) (a), and the least-square-fit slopes of NO$_y$ to CO$_2$ (from the observation) and NO$_y$ to CO (from the observation) (b). All slopes are normalized to the slope at the first transect.
Fig. 9. O₃ versus NOₓ-NOₓ from the transects of Martin Lake (Ma-6, plume age of 2.0 hours), Monticello (Mo-4, plume age of 2.4 hours), and Welsh (We-4, plume age of 2.7 hours). The slopes from the least square fits indicate the observation-based estimates of OPE from each plume transect.