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Improving regional ozone modeling through systematic evaluation of errors using the aircraft observations during the International Consortium for Atmospheric Research on Transport and Transformation

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[1] During the operational phase of the ICARTT field experiment in 2004, the regional air quality model STEM showed a strong positive surface bias and a negative upper troposphere bias (compared to observed DC-8 and WP-3 observations) with respect to ozone. After updating emissions from NEI 1999 to NEI 2001 (with a 2004 large point sources inventory update), and modifying boundary conditions, low-level model bias decreases from 11.21 to 1.45 ppbv for the NASA DC-8 observations and from 8.26 to -0.34 for the NOAA WP-3. Improvements in boundary conditions provided by global models decrease the upper troposphere negative ozone bias, while accounting for biomass burning emissions improved model performance for CO. The covariances of ozone bias were highly correlated to NO_z , NO_y , and HNO_3 biases. Interpolation of bias information through kriging showed that decreasing emissions in SE United States would reduce regional ozone model bias and improve model correlation coefficients. The spatial distribution of forecast errors was analyzed using kriging, which identified distinct features, which when compared to errors in postanalysis simulations, helped document improvements. Changes in dry deposition to crops were shown to reduce substantially high bias in the forecasts in the Midwest, while updated emissions were shown to account for decreases in bias in the eastern United States. Observed and modeled ozone production efficiencies for the DC-8 were calculated and shown to be very similar (7.8) suggesting that recurring ozone bias is due to overestimation of NO_x emissions. Sensitivity studies showed that ozone formation in the United States is most sensitive to NO_x emissions, followed by VOCs and CO. PAN as a reservoir of NO_x can contribute to a significant amount of surface ozone through thermal decomposition.

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1. Introduction

[2] Air pollution models have been used to predict air quality during numerous field campaigns [Bates et al.,

1998; Jacob et al., 2003; Menut et al., 2000; Ramana et al., 2004; Vautard et al., 2003], with the objective to both place air pollution in a geographical context for experimen-

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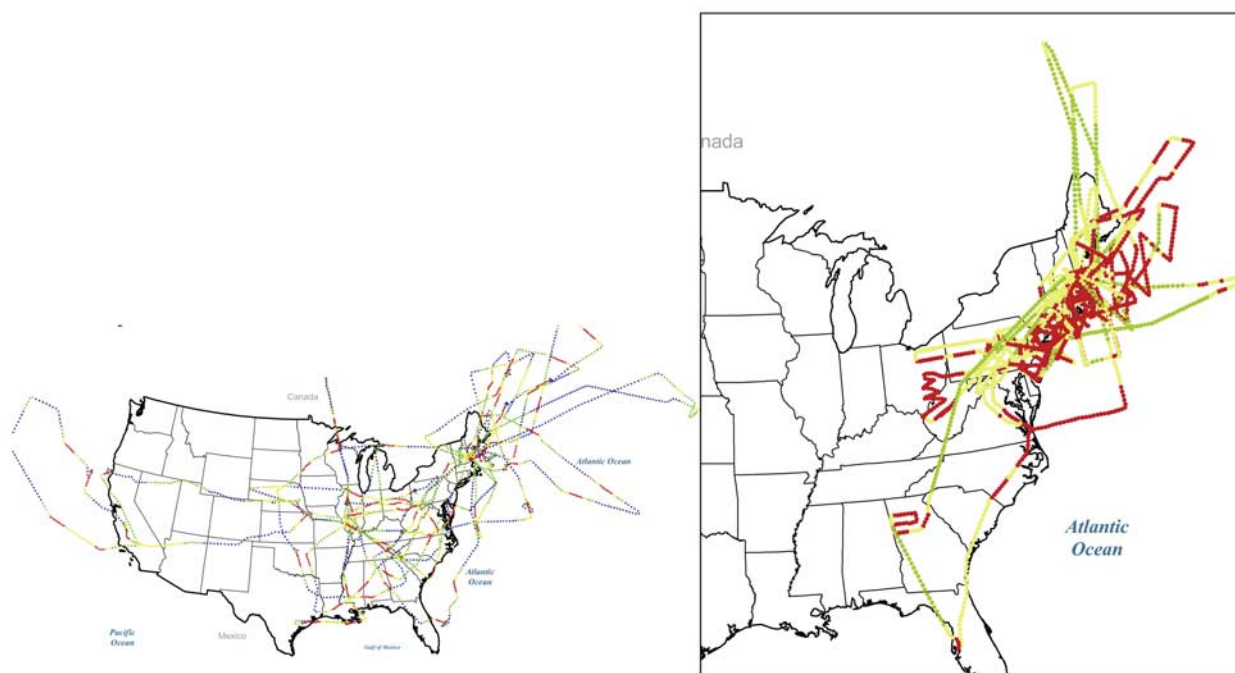


Figure 1. Flight tracks and altitude range for ICARTT. Red indicates 0–1 km, yellow indicates 1–4 km, green indicates 4–8 km, and blue indicates 8–12 km. (left) DC-8 and (right) WP-3.

tal design and, as the data is collected, evaluate our current understanding of atmospheric processes [Kiley *et al.*, 2003] and anthropogenic and biogenic emissions [Carmichael *et al.*, 2003]. During the summer of 2004, the International Consortium for Atmospheric Research on Transport and Transformation (ICARTT) field experiment was performed (<http://esrl.noaa.gov/csd/ICARTT>), which included a NASA experiment called INTEX-A (Intercontinental Chemical Transport Experiment–A), and a NOAA experiment called NEAQS/ITCT-2k4 (New England Air Quality Study–Intercontinental Transport and Chemical Transformation, 2004). During this period NASA DC-8 and NOAA WP-3 aircraft each performed 18 research flights over the continental United States, with a special focus on the northeastern United States. Figure 1 shows the flight tracks of the DC-8 and WP-3 during the mission, and the altitude at which the aircraft flew. The WP-3 aircraft flew at lower altitudes, mostly over the NE United states. More details about the aircraft measurements and main findings are given by Singh *et al.* [2006]. During the field experiment operations forecasts using the University of Iowa STEM model [Carmichael *et al.*, 1991; Tang *et al.*, 2003, 2004] were used (along with other models) in support for flight planning. Analysis of the forecasts has shown a persistent positive bias (modeled–observed) for ozone [McKeen *et al.*, 2006] in comparison to surface sites in the AIRMAP network (<http://airmap.unh.edu/>), and in comparison to the aircraft platform observations. The objective of this paper is to show that model performance in relation to ozone and its precursors was improved through systematic analysis of model prediction with the observed data to evaluate where model error persists, and how ozone model error is related to model error of other species. The paper also demonstrates how

geospatial interpolation through kriging can be used to provide geographical contextualization to model error, which in turn can be used to improve model performance.

2. Methodology

[3] Figure 2 shows the layout of this study, in which systematic data analysis is used to improve model performance by modifying boundary conditions and emissions inventories through an iterative process which included geospatial interpolation of model bias through kriging to provide qualitative support for regional modification of emissions. After selecting the best model run sensitivity studies are carried out to evaluate the contribution of different ozone precursors to ozone formation.

[4] In this study we used the STEM-2K3 model [Carmichael *et al.*, 1991]. The model features the lumped species SAPRC99 chemical mechanism [Carter, 2000] with an online photolysis solver, and the SCAPE II aerosol solver (Simulating Composition of Atmospheric Particles in Equilibrium) [Kim *et al.*, 1993a, 1993b; Kim and Seinfeld, 1995]. Meteorological inputs to the model came from MM5 [Grell *et al.*, 1995], using the AVN data [Huang *et al.*, 1997] during forecasting and NCEP FNL (Final Global Data Assimilation System) analyzed data during postanalysis. For this study the model domain was the continental United States, using a 60 km resolution, 62 cells in longitude, and 97 cells in latitude. The model had 21 vertical layers, extending from the surface to 100 hPa. The Grell cumulus parameterization [Grell *et al.*, 1995] and the MRF planetary boundary layer parameterization [Hong and Pan, 1996] were used for the MM5 runs.

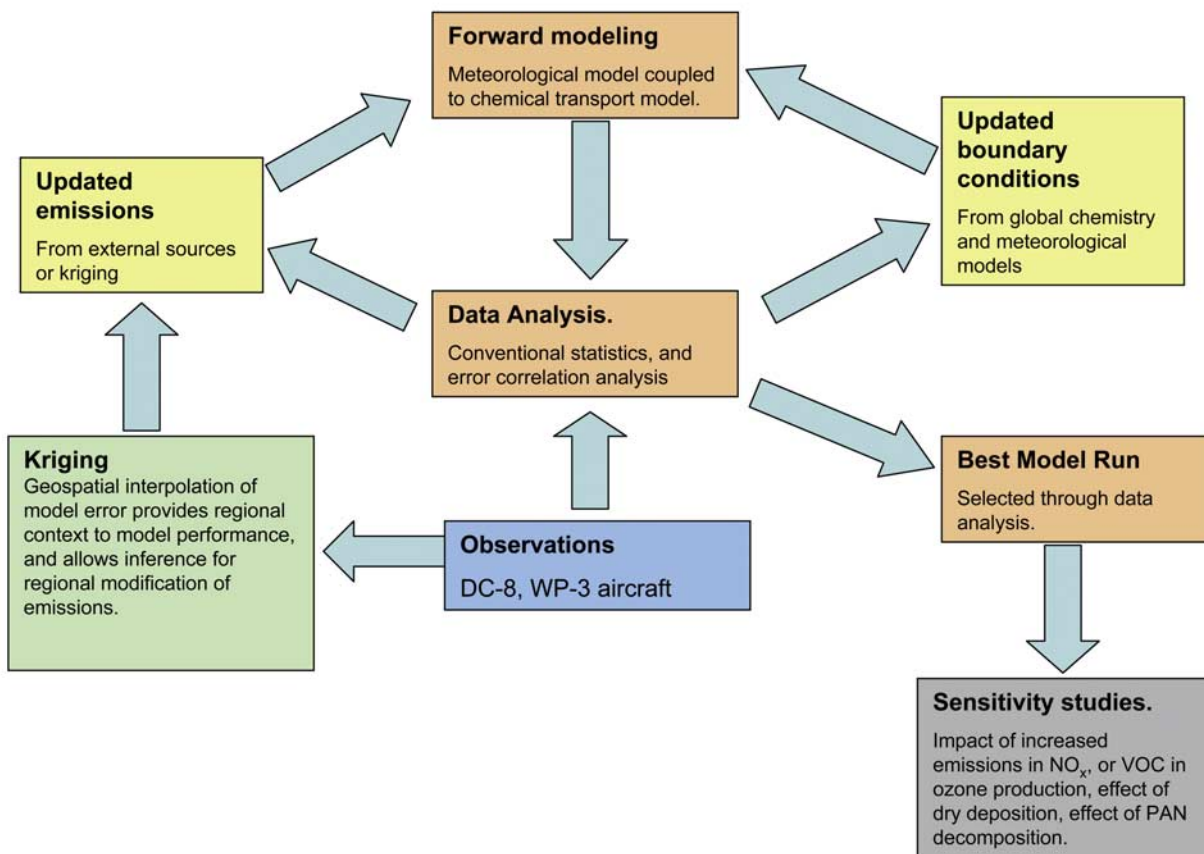


Figure 2. Study framework for data analysis and sensitivity studies.

[5] During the operational portion of the experiment, anthropogenic emissions were taken from the U.S. EPA National Emissions Inventory for the base year of 1999 (NEI 1999) [U.S. Environmental Protection Agency (U.S. EPA), 2006] while the 2001 update of the same (NEI 2001) was used for the postanalysis stage. It should be noted that NEI 2001 has lower emissions of CO, NO_x, and SO₂ than NEI 1999 (Table 1). Modifications were still needed since the simulations with NEI 2001 systematically underestimated light alkanes, and overpredicted aromatic species. The large point source emissions (LPS) used were from the updated inventory by Gregory Frost at NOAA Earth Systems Research Laboratory [Frost et al., 2006], which represents emissions for 2004, the year of the campaign. Upper troposphere lightning NO_x emissions were added to the model in postanalysis based on the National Lightning Detection Network (NLDN), relating emissions to signal strength and multiplicity of flashes. Further information about the lightning emissions used is given by Tang et al. [2007]. Biogenic emissions were estimated using BEIS 2 (Biogenic Emissions Inventory System) [Geron et al., 1994], which generates time-variable isoprene and monoterpene emissions driven by meteorological variables from MM5 simulations. Forest fires that occurred during the ICARTT period were largely outside our regional model domain (in Alaska and northwestern Canada). Their episodic influence on lateral boundary conditions was incor-

porated during postanalysis using MOZART NCAR boundary conditions [Pfister et al., 2005] with data assimilated CO concentrations from Measurements of Pollution in the Troposphere instrument on board the TERRA satellite (MOPITT) to constrain the fire emissions influence for the study period, while during the forecast MOZART GFDL [Horowitz et al., 2003] boundary conditions were used, which did not include episodic fire emissions, but climatological emissions.

[6] The postanalysis work focused on improving model performance by carefully comparing predictions with observations, and the use of the error information to identify aspects of the model in need of improvement. Model sensitivity studies were done for factors with significant uncertainty including boundary conditions, anthropogenic

Table 1. Total Column Emissions in Model Domain, in Tg/year, Including Point Source, Area, and Aviation Emissions^a

	Emissions, Tg/year		
	CO (as CO)	NO _x (as N)	VOCs (as C)
NEI 1999	40.7	7.1	15.7
NEI 2001	25.0	5.2	13.5
NEI 2001-Frost LPS	26.8	4.8	13.8
NEI 2001-Frost LPS*	26.8	4.2	13.8

^aCO emissions reported as Tg C/year, and NO_x emissions as Tg N/year.

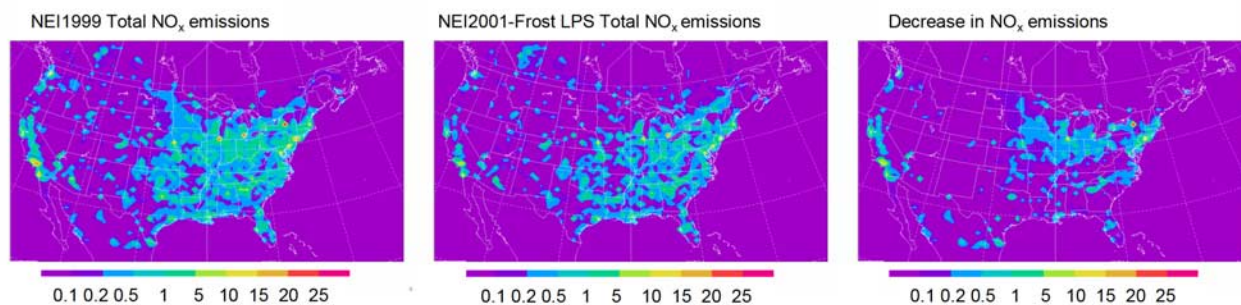


Figure 3. Total column NO_x emissions. (left) NEI 1999. (middle) NEI 2001-Frost LPS emissions. (right) Decrease in NO_x emissions from NEI 1999 to NEI 2001-Frost LPS. Scale is tonnes/ km^2 /year.

emissions inventories, sea salt emissions, lightning NO_x emissions, and dry deposition rates. From these various runs three were selected for detailed analysis in this paper. These are (1) operational forecasting conditions (Forecast, NEI 1999); (2) the most updated emissions inventory (NEI2001-Frost LPS); and (3) a modification of that emissions inventory (NEI2001-Frost LPS*) that intended to improve model performance by further decreasing the regional NO_x bias. Table 2 shows a summary of the model parameters for the different scenarios compared in this study. Table 1 shows that the total column anthropogenic CO emissions for the domain were reduced by $\sim 40\%$ from NEI 1999 to NEI 2001, and that the NEI2001-Frost LPS estimate increases CO with respect to NEI 2001. The surface NO_x emissions were reduced significantly from NEI 1999 to NEI 2001 ($\sim 30\%$). Figure 3 shows the domain column emissions of NO_x for NEI 1999 (Figure 3, left), NEI2001-Frost LPS (Figure 3, middle), and the decrease of emission from NEI 1999 to NEI 2001-Frost LPS (Figure 3, right).

[7] The results from runs with these different conditions for the period 1 July to 18 August 2004, which includes a model spin up period and which spans the times of the DC-8 and WP-3 flights 3–20 for ICARTT, are discussed in this paper. Merged data for both measurement platforms were resampled from a 1 s to a 3 min resolution, and compared to interpolated data from the 3 h, 60 km, and 21-level variable vertical resolution model output.

[8] Kriging is a geospatial interpolation technique [Oliver and Webster, 1990] that assumes that bias at a point without observations can be related to points with observations, by an expression which considers three components: an aver-

age term, a spatially correlated term, and a random error term. The spatially correlated component is calculated using a semivariogram, constructed by the semivariance among observations as a function of distance. Kriging has been previously used for interpolating surface measurements of ozone, and particulate matter for health studies, and estimation of exposure [Liao *et al.*, 2006], to generate maps of air pollution based on discrete measurements, such as the AIRNOW network [U.S. EPA, 1999], and to provide a geographical perspective to ozone analyses [Blond *et al.*, 2003]. Kriging produces a surface of predicted values and uncertainty using a semivariogram (in this case exponential), which relates percent bias (bias/mean observed*100) to distance among points. The analysis is limited to altitudes less than 4000 m. This assumes that the vertical variability in this range is smaller than the horizontal variability. The continuous surface output of kriging provides geospatial context to bias, and allows the comparison of related bias of different precursors to ozone.

3. Data Analysis

[9] The surface ozone forecasted during ICARTT has been compared with surface observations and showed a significant high bias for daytime values (~ 15 ppbv) [McKeen *et al.*, 2006]. We anticipated such bias would occur because of the fact that the experiment was conducted in 2004, and that the actual emissions would differ from the 1999 values used in the forecast. The summer of 2004 presented significant fires in Alaska and Canada which were misrepresented by climatological fires used in the forecast.

Table 2. Summary of Model Parameters for Scenarios Studied During Forecast and Postanalysis

Scenario	Parameters					
	National Emissions Inventory	Biogenic Emissions	Boundary Conditions	Lightning NO_x	Dry Deposition	Meteorology
Forecast	1999	BEIS 2	MOZART-GFDL, climatological fires	no	dormant agricultural lands	MM5 and forecasted AVN global model
NEI2001-Frost LPS ^a	2001, Frost Large Point Sources. VOCs adjusted by increasing alkanes and alkenes, and decreasing aromatics.	BEIS 2	MOZART NCAR, burned area biomass burning emissions [Pfister <i>et al.</i> , 2005]	yes	nondormant agricultural lands	MM5 and NCEP FNL reanalyses

^aParameters for NEI2001-Frost LPS* are the same but reducing area NO_x emissions by 60% for Alabama, Mississippi, Georgia, North Carolina, South Carolina, Tennessee, West Virginia, Indiana, and Ohio.

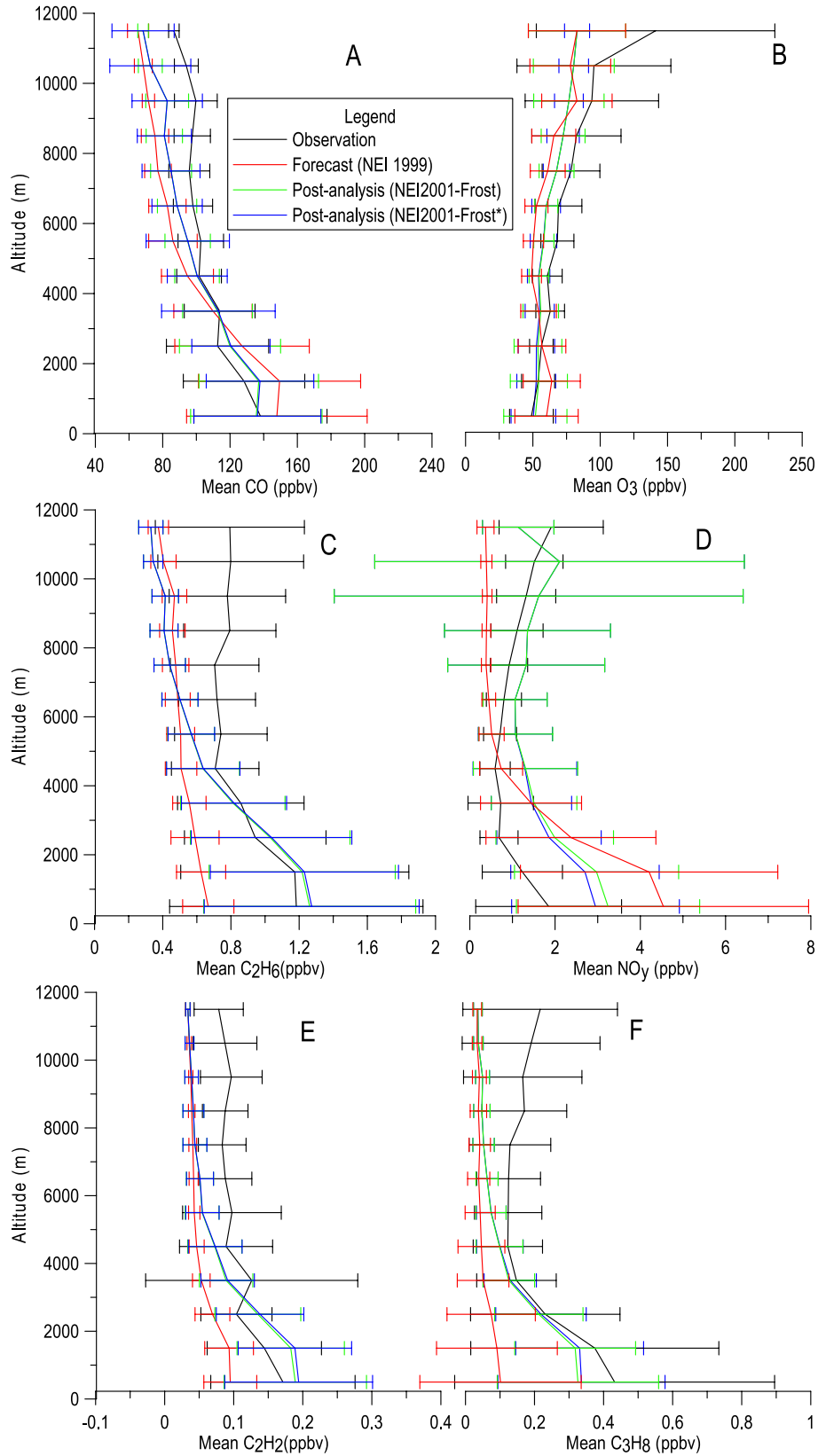


Figure 4. Observed and 60 km simulated O₃, CO, and NO_y vertical profiles and standard deviations for all DC-8 flights. (a) CO, (b) ozone, (c) ethane, (d) NO_y, (e) ethyne, and (f) propane. Blue indicates NEI 2001-Frost LPS*, red indicates forecast, NEI 1999, and black indicates observed.

Table 3. Model Performance Statistics for Selected Species^a

	O ₃				CO				NO _y			
	Forecast	Frost LPS	Frost LPS*	Obs	Forecast	Frost LPS	Frost LPS*	Obs	Forecast	Frost LPS	Frost LPS*	Obs
	<i>0–1 km</i>											
Mean modeled, ppbv	60.12	51.82	50.36	48.92	148	136	136	138	4.16	3.24	2.95	1.85
Mean bias, ppbv	11.21	2.90	1.45	–	9.75	–2.48	–1.84	–	2.70	1.40	1.11	–
S.D./mean modeled	0.39	0.34	0.33	0.33	0.36	0.29	0.27	0.28	0.77	0.66	0.67	0.92
R	0.71	0.70	0.72	–	0.61	0.68	0.68	–	0.59	0.59	0.56	–
	<i>1–4 km</i>											
Mean modeled, ppbv	58.97	54.68	53.32	57.34	131	127	125	119	2.76	2.26	2.09	0.93
Mean bias, ppbv	1.63	–2.66	–4.02	–	11.37	5.20	5.82	–	1.83	1.36	1.19	–
S.D./mean modeled	0.31	0.26	0.25	0.20	0.14	0.11	0.11	0.26	0.87	0.74	0.72	0.88
R	0.40	0.48	0.50	–	0.49	0.65	0.65	–	0.49	0.52	0.51	–
	<i>4–12 km</i>											
Mean modeled, ppbv	63.71	67.97	67.89	80.20	79	85	85	98	0.46	1.43	1.43	1.05
Mean bias, ppbv	–16.49	–12.23	–12.31	–	–19.37	–12.64	–12.60	–	–0.59	0.43	0.42	–
S.D./mean modeled	0.01	0.00	0.00	0.01	0.00	0.00	0.00	0.20	1.25	1.37	1.37	0.60
R	0.56	0.47	0.47	–	0.06	0.33	0.33	–	–0.13	0.10	0.10	–

^aModeled versus observed data, DC-8 platform. 0–12 km range. Frost LPS: NEI2001-FrostLPS. Frost LPS*: NEI2001-Frost LPS*. Modified area NO_x emissions. SD, standard deviation.

Additionally, it was found in postanalysis that the dry deposition velocities for agricultural crops were incorrectly set to low growing season conditions. These factors all contributed to forecast errors.

[10] Below we compare the forecasted values with aircraft observations. We also compare the results from model runs where the dry deposition velocity has been corrected, and where the emissions and boundary conditions have been updated.

3.1. Statistical Performance for All Flights

[11] The predictions of O₃, CO and NO_y for the various simulation cases are compared with the DC-8 observations in Figure 4 and Table 3. For these comparisons all data from flights 3–20 are combined together and analyzed by altitude. The predicted values are interpolated to the same spatial location of the observations using trilinear interpolation. In general the forecast values show a significant positive bias in predicted ozone at altitudes below ~4 km, and a high negative bias above this altitude. The mean bias below 1 km is ~11 ppbv, similar to, but slightly lower than the values found from the analysis of the surface AirMap observations. Comparable patterns are found in the forecast for CO and NO_y, with high values at low altitudes and low values at high altitudes. The postanalysis runs show significant improvements in the predictions at altitudes below 4 km (see for example the correlation coefficients for the 1–4 km range). In the case of ozone the NEI2001-Frost LPS case shows that the low altitude bias is reduced to less than 3 ppbv. The bias in the mid troposphere (4–12 km) is also reduced (by ~25%). Similar improvements are found for CO, where the bias above 4 km is reduced by updated global boundary conditions (MOZART-NCAR) that improve the correlation coefficient for CO because of their better representation of the biomass burning emissions from Alaska and northern Canada. Tang *et al.* [2007] evaluated the impact of boundary conditions on model performance using results from three different global models and found that they dominate the performance of the regional model at

these altitudes. The remaining bias reflects the performance skills of the global models used. For NO_y, the bias in the near surface regions is reduced, albeit at a smaller rate, while the negative bias at higher altitudes decreases significantly. The improvements at the higher altitudes reflect the importance of including lightning NO_x emissions. The comparison of the ratio of the standard deviation to the mean value of the predictions and those for the observations shows that the model exhibits a variability that is similar to that of the observations.

[12] Similar results are found for the WP-3 comparisons (Table 4). As shown by comparing Figures 1 and 2, the flight operations of the WP-3 and DC-8 were different, with the WP-3 focused largely on the northeast United States. This along with the flight altitude differences lead to differences in the statistics in the observed distributions of the DC-8 and WP-3 data. For example, the WP-3 low altitude concentrations on average are higher. In the case of ozone and CO the mean observed values from all flights were 56 and 158 ppbv for the WP-3 and 49 and 138 ppbv, respectively, for the DC-8. The mean bias in the forecast for ozone for the WP-3 was ~8 ppbv, compared to 11 ppbv for the DC-8 comparison. Correlation coefficients for both aircraft were very similar for the 0–1 km range (0.71 for DC-8, and 0.63 for WP-3). For the postanalysis simulation NEI2001-Frost LPS the bias in predicted ozone at low altitudes was reduced to 1.34 ppbv and the correlation was increased to 0.66. The mean biases for the lower altitude predicted for CO and NO_y were also reduced significantly (by ~90 and 70%, respectively). For the 1–4 km range the ozone correlation improved (R increased from 0.57 to 0.65) and the mean bias decreased from 4.91 to –0.58. CO predictions also improved with R increasing from 0.52 to 0.66, and bias decreasing from 28.37 to 8.79. Note that the CO bias remains significant for the WP-3 flights, from which we can infer that there is still a systematic over prediction of CO for the area that was sampled by WP-3, i.e., the NE United States. For the 4–6 km range emissions and boundary condition improvements significantly enhanced ozone

Table 4. Model Performance Statistics for Selected Species^a

	O ₃				CO				NO _y			
	Forecast	Frost LPS	Frost LPS*	Obs	Forecast	Frost LPS	Frost LPS*	Obs	Forecast	Frost LPS	Frost LPS*	Obs
	<i>0–1 km</i>											
Mean modeled, ppbv	64.52	57.60	55.93	56.27	202.81	163.56	164.49	158.60	7.77	5.03	4.79	3.91
Mean bias, ppbv	8.26	1.34	−0.34	–	44.24	4.96	5.89	–	3.87	1.12	0.88	–
S.D./mean modeled	0.38	0.31	0.30	0.35	0.31	0.24	0.24	0.24	0.65	0.63	0.65	0.72
R	0.63	0.66	0.69	–	0.43	0.42	0.42	–	0.28	0.29	0.29	–
	<i>1–4 km</i>											
Mean modeled, ppbv	65.07	62.05	59.59	60.16	164.04	143.34	144.46	135.66	5.00	3.58	3.34	2.38
Mean bias, ppbv	4.91	1.89	−0.58	–	28.37	7.67	8.79	–	2.62	1.20	0.96	–
S.D./mean modeled	0.34	0.27	0.26	0.23	0.38	0.28	0.28	0.34	0.86	0.71	0.73	0.88
R	0.57	0.63	0.65	–	0.52	0.66	0.66	–	0.64	0.64	0.65	–
	<i>4–6 km</i>											
Mean modeled, ppbv	50.61	59.74	58.97	66.80	92.08	101.37	101.77	106.66	0.82	1.34	1.29	1.13
Mean bias, ppbv	−16.19	−7.06	−7.83	–	−14.57	−5.28	−4.88	–	−0.31	0.21	0.16	–
S.D./mean modeled	0.14	0.20	0.19	0.16	0.13	0.24	0.24	0.33	0.67	0.62	0.64	0.35
R	0.15	0.44	0.46	–	0.11	0.36	0.36	–	0.24	0.40	0.39	–

^aModeled versus observed data, WP-3 platform. 0–6 km range. Frost LPS: NEI2001-FrostLPS. Frost LPS*: NEI2001-Frost LPS*. SD, standard deviation.

modeling performance, with R increasing from 0.15 to 0.46, and bias decreasing from −16.19 to −7.83. Similarly CO performance increased because of boundary conditions incorporating biomass burning (R increases from 0.11 to 0.36, and the negative bias is improved from −14.57 to −5.28). It is important to note that while the predicted biases in NO_y were reduced significantly in the postanalysis runs, they remain quite high (~1 ppbv overprediction when averaged over all altitudes for the DC-8 and WP-3 observations). The NO_y distributions and their comparison with various models used during ICARTT are discussed in detail by Singh *et al.* [2006]. In Figure 5 we plot the observed and predicted contribution of individual species to NO_y for the DC-8 observations and for the NEI2001-Frost LPS simulations. This plot shows that the predicted contributions are similar to those observed. Nitric acid is shown to compose the largest NO_y fraction below ~4 km, above which PAN contributes from 30 to 45% up to about 8 km; the relative contribution of NO in comparison to NO₂ increases with

altitude in the upper troposphere (up to 45% of total nitrogen) Within the boundary layer PAN and NO₂ each contribute ~20% to NO_y. The predicted distributions differ in comparison with the observations in that the relative contributions of HNO₃ are lower than those observed, possibly being linked to overprediction of rainout or wash-out processes upwind of the sampling. In addition the predicted contribution of NO increases with altitude at a slower rate than observed. This fact is probably related to the treatment of the lightning NO_x emissions. Lightning NO_x emissions at higher altitudes (4–6 km) improved the modeling of reactive nitrogen species, decreasing the negative bias of NO_y in the upper troposphere.

[13] Recurring positive bias of NO_y and its components suggest that NO_x emissions in the model are still higher than the actual emission in the summer of 2004. While the emissions used in this simulation have the large point source sector updated to 2004, emissions from the other sectors are based on 2001 values. The transportation sector is the major

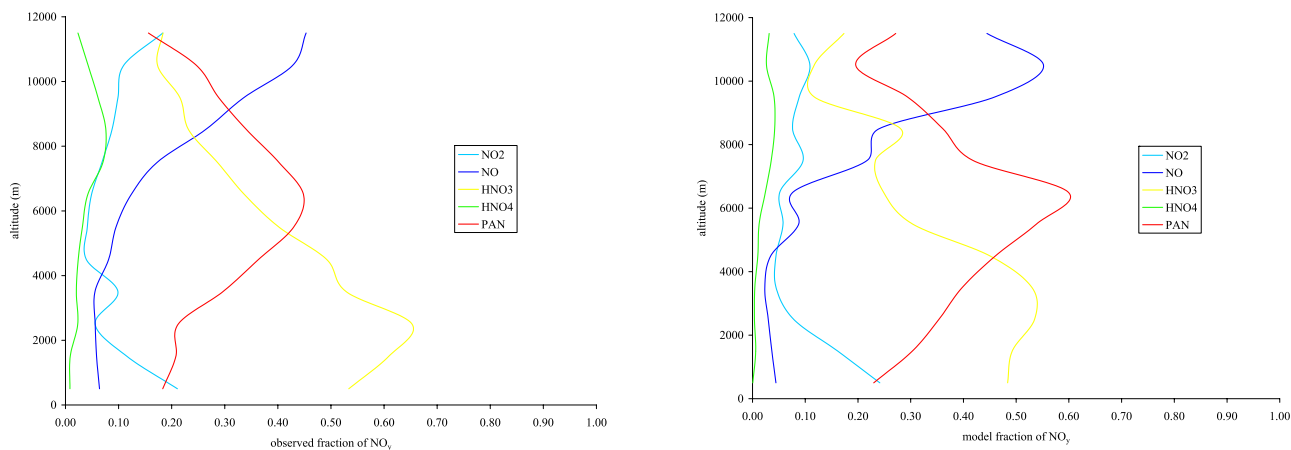


Figure 5. Comparison of mean contributions to NO_y along the DC-8 flight tracks as a function of altitude. (left) Observed values and (right) predicted for the NEI2001-Frost-LPS case. Values are plotted as fraction of total NO_y, defined as the sum of NO₂, NO, HNO₃, HNO₄, and PAN.

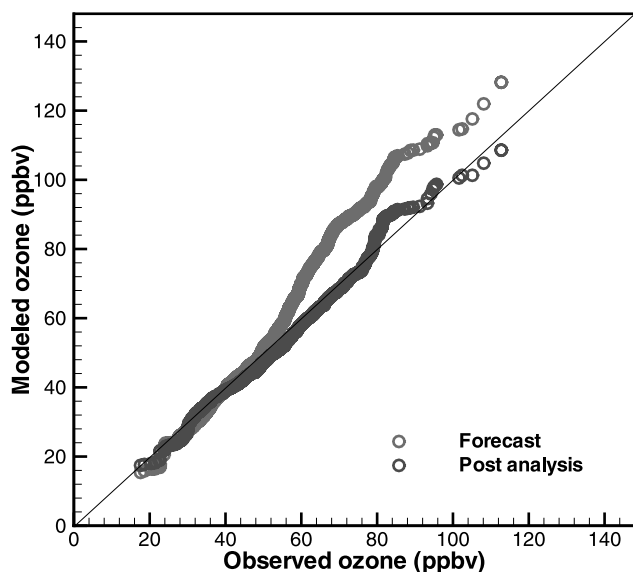


Figure 6. Quantile-quantile plot of modeled ozone with observed ozone for DC-8 platform, data points collected at altitude less than 4000 m, STEM-2K3. Forecast is NEI 1999, and postanalysis is NEI2001-Frost LPS*. MOZART-NCAR boundary conditions.

emission sector for NO_x and transport emissions have shown a downward trend [U.S. Department of Transportation, 2006]. Continuously monitored emissions of NO_x from large point sources have also decreased [Frost *et al.*, 2006]. So it is most likely that NO_x emissions in 2004 are actually lower than those in 2001 [Kim *et al.*, 2006]. To reflect this case we performed an additional simulation (NEI2001-Frost LPS* case) where the NO_x emissions were reduced by an additional 12% with respect to total NO_x on a national level (but by 60% for area NO_x emissions for selected states, as is discussed in Table 1). The results of this case are also presented in Tables 3 and 4, and Figure 4. The effect of this reduction in NO_x emissions is to further reduce by $\sim 20\%$ the bias in NO_y . Since ozone production in the ICARTT region is largely NO_x limited, this reduction in NO_x emissions also reduced the mean ozone levels by ~ 1 ppbv, and further reduced the bias in the lowest layers by 50% (to 1.45 ppbv for the case of the DC-8), compared to the NEI2001-Frost LPS case.

[14] The modeling of volatile organic compounds (VOCs) has always been a challenge because of the uncertainty of volatile organic compound emissions (VOC) emissions inventories in the United States [Parrish, 2006]. At altitudes above 2 km, all model scenarios showed a negative bias for the prediction for ethane, ethene, and propane, largely due to the global model boundary conditions with contributions from errors from imprecise treatment of convective events. In general R values decrease with altitude reflecting the fact that model performance is highly dependent on boundary conditions at higher altitudes.

[15] Figure 6 shows a quantile-quantile plot of observed and modeled O_3 for the DC-8 for measurements under 4000 m altitude range. The forecast values show a systematic over-prediction across the whole range, while the NEI2001-Frost LPS* case shows great improvement for values over

55 ppbv, which represented the majority of the points sampled. Figure 7 shows that the frequency for which ozone bias (modeled-observed bias divided by average observed value $\times 100$) is between -20 to 20% has increased from 51 to 73% of the time with respect to the forecast, and model error between -5 and 5% has increased from 14 to 23% with respect to forecast. Postanalysis work therefore resulted in a nearly unbiased distribution of ozone errors with a significant reduction in the standard deviation of ozone errors.

3.2. Case Studies

[16] The results above provide a mission wide perspective. NEI2001-Frost LPS* ozone bias is shown on a flight by flight basis in Figure 8. Generally the bias in the lowest layers is less than 5 ppbv, while the bias in the 8–12 km range is large, and particularly high in flights 3, 11, 15, 16, and 17. This is a reflection of the boundary conditions from the global models. Tang *et al.* [2007] shows that boundary conditions varied greatly among global models, and depending on the global models used the ozone bias in the upper troposphere varied from large negative to large positive values, reflecting differences in the global models in placing stratospheric intrusion of ozone. To show how the model predictions changed from the forecast to postanalysis details for specific flights were also analyzed. Figure 9 shows how model performance improved during DC-8 flights 12 and 14 (25 and 31 July 2004). The plots show that at low altitudes (0–1 km) the model bias is reduced from 8 to 2 ppb for flight 14, and from 22 to 12 ppbv for flight 12, when using improved emissions. For the upper troposphere the model showed a large biases (positive for flight 14, and negative for flight 12), which were decreased significantly by using updated boundary conditions from NCAR MOZART. This improvement is reflected in an increase in the correlation coefficients from 0.65 to 0.84 for flight 14, and from 0.01 to 0.78 for flight 12.

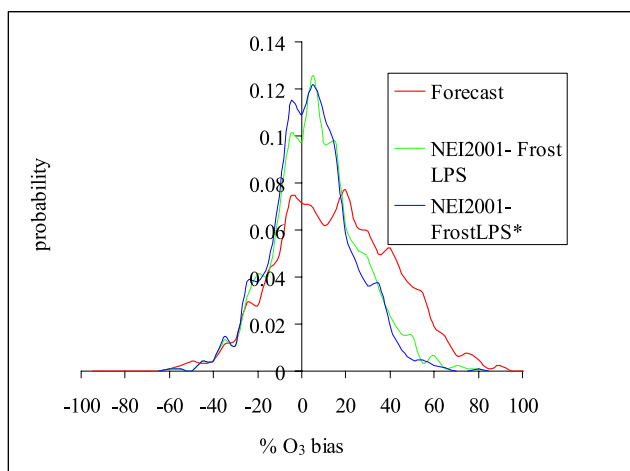


Figure 7. Probability distribution of % ozone bias for forecast (NEI 1999) and postanalysis runs (NEI2001-FrostLPS and NEI2001-FrostLPS*) for DC-8 measurements under 4000 m.

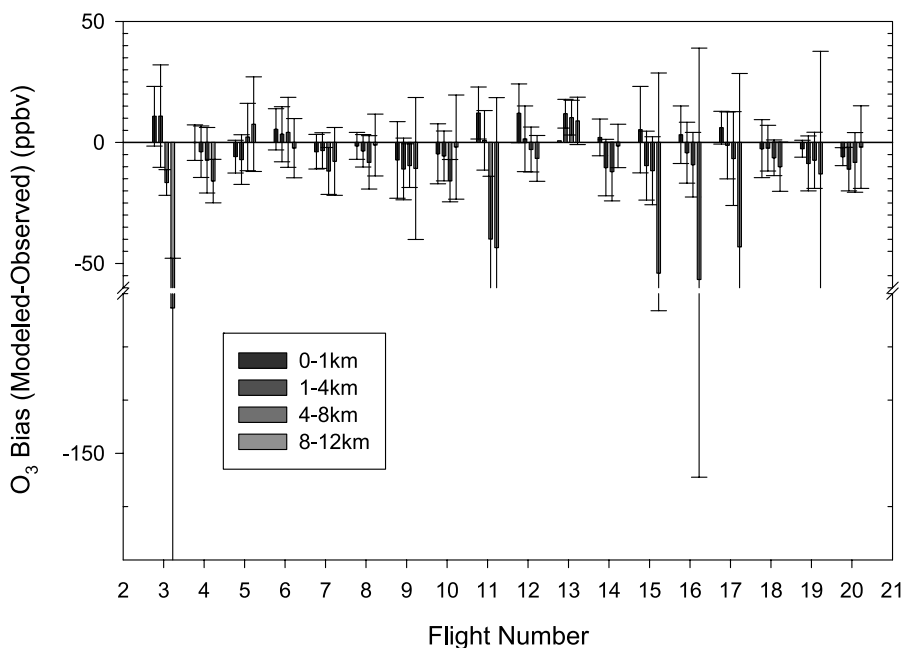


Figure 8. Mean ozone bias for DC-8 flights 3–20, separated by altitude range, for the NEI2001-FrostLPS* case. Error bars represent standard deviation.

4. Analysis of Model Error

[17] The relationship between model errors is a key step in understanding model behavior and identifying model deficiencies. This information is also becoming increasingly important as estimates of error covariance are an important aspect of chemical data assimilation [Chai *et al.*, 2007]. The

ICARTT experiment produced observations for a large spectrum of species that are involved in the photochemical oxidant cycle. Thus it is possible to use these data to explore the relationships between the calculated and observed species concentrations with respect to ozone. In this section we analyze two relationships: the correlation of observed

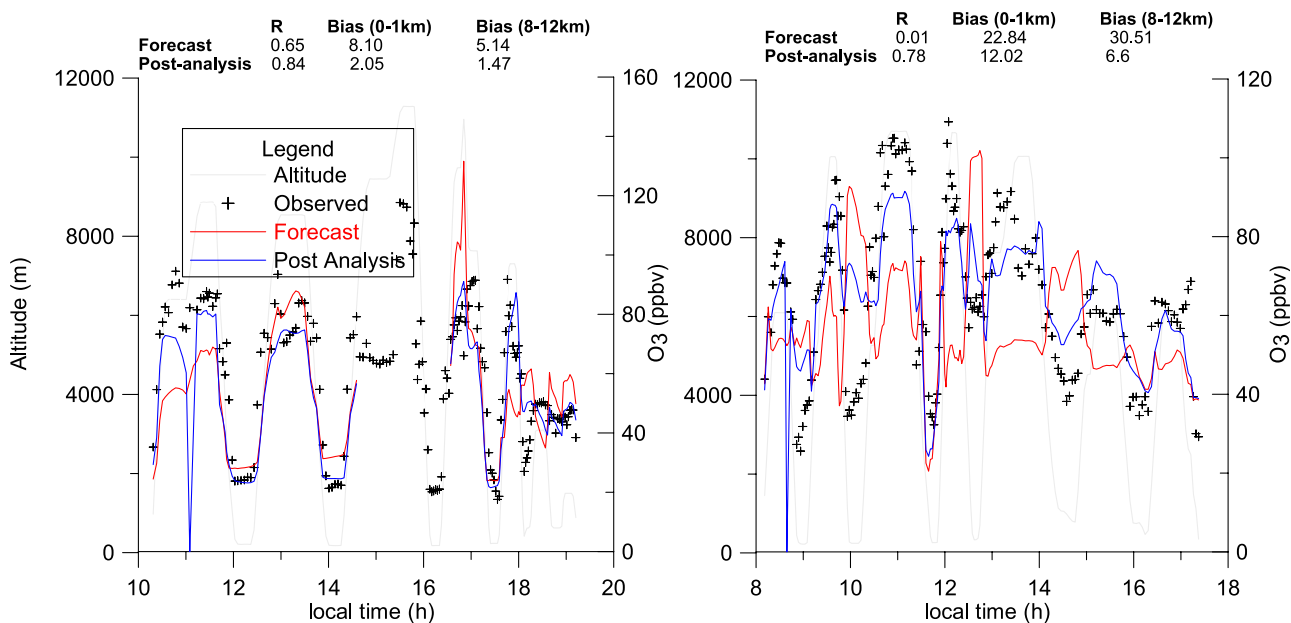


Figure 9. (left) Time series of observed and modeled ozone along DC-8 flight 14 flight track (31 July 2004). (right) Time series of observed and modeled ozone along DC-8 flight 12 track (25 July 2004). Absence of modeled data denotes that flight went beyond model boundaries. Forecast run is NEI 1999, MOZART boundary conditions. Postanalysis is NEI 2001-Frost LPS*, RAQMS boundary conditions.

Table 5. Correlation Coefficients of Observed Parameters Versus Observed Ozone and Bias (Modeled-Observed) of Parameters to Ozone Bias, at Different Altitude Ranges for DC-8 Platform^a

	Observation Correlation Coefficients				Bias Correlation Coefficients.			
	0–1 km	1–4 km	4–8 km	8–12 km	0–1 km	1–4 km	4–8 km	8–12 km
Acetone	0.68	0.21	0.11	−0.12	0.42	0.15	0.10	−0.14
ARO2	−0.02	0.11	0.31	–	0.04	0.04	0.89 (n = 6)	−0.95
ARO1	0.47	0.40	0.01	0.07	0.09	0.07	0.13	0.02
Acetaldehyde	0.38	−0.07	0.04	0.10	0.24	0.20	−0.07	0.12
CO	0.70	0.44	0.08	−0.24	0.51	0.44	0.19	−0.32
Ethene	0.24	0.06	−0.04	−0.07	0.04	−0.05	0.02	−0.05
Ethyne	0.75	0.48	0.14	0.00	0.18	0.25	0.26	−0.07
H ₂ O ₂	0.51	0.24	−0.29	−0.28	0.28	−0.01	−0.27	−0.29
Formaldehyde	0.53	0.07	−0.09	−0.12	0.20	0.11	0.18	−0.04
HNO ₃	0.86	0.50	0.19	0.67	0.47	0.45	0.44	0.36
HO ₂	0.48	0.23	−0.14	−0.28	0.15	0.09	0.14	−0.10
Isoprene	−0.03	−0.03	−0.19	–	−0.48	−0.38	−0.34	–
MEK	0.65	0.17	0.08	−0.14	0.48	0.44	0.11	−0.03
NO	−0.03	−0.06	0.19	−0.04	−0.17	0.07	0.10	0.04
NO ₂	0.15	0.07	0.48	0.14	−0.07	0.20	0.20	0.10
NO _y	0.63	0.30	0.48	0.27	0.34	0.55	0.39	0.10
NO _z	0.77	0.37	0.47	0.42	0.45	0.57	0.50	0.28
OH	0.55	0.24	0.23	0.12	0.24	0.23	−0.09	−0.01
PAN	0.64	0.46	0.39	0.03	0.60	0.64	0.43	−0.05
Propane	0.40	0.16	0.06	−0.10	0.14	0.14	0.01	−0.10
Propene	−0.06	0.05	0.06	–	−0.25	−0.12	0.13	–
RH	−0.21	−0.38	−0.28	−0.35	−0.14	−0.05	−0.03	−0.08
SO ₂	0.28	0.12	0.09	0.04	0.07	0.35	0.17	−0.01
Temperature	0.33	−0.09	−0.35	−0.19	0.27	0.15	−0.08	−0.41
Wind speed	−0.16	−0.06	0.13	0.14	0.04	0.10	0.09	0.06

^aBias calculated with respect to NEI2001-FrostLPS. Ranges 0–1 km, 1–4 km, 4–8 km, and 8–12 km.

ozone concentrations with respect to other measured parameters, and the correlation of ozone bias (or error), with respect to other species. Simultaneous analysis of these relationships provides valuable insight on ozone chemistry, and sources of model error.

4.1. Correlation Between Model Biases

[18] Comprehensive field experiments such as ICARTT provide an opportunity to examine the complex relationships between the processes that govern ozone distributions in the troposphere. Toward these ends we examined the correlations between the observed ozone, and ozone bias to various species using the DC-8 data and the results are presented in Table 5. Shown are results for the postanalysis simulation NEI2001-Frost LPS.

[19] We first examine the relationships between ozone and other parameters using all the DC-8 flight data binned by altitude. Table 5 shows that the observed ozone concentrations at low altitudes (0–1 km) are most strongly correlated to HNO₃ (R = 0.86), ethyne (R = 0.75), CO (R = 0.70), acetone (R = 0.68), MEK (R = 0.65), NO_z (R = 0.77), PAN (R = 0.64) and NO_y (R = 0.63). These species represent the general features important in photochemical production, i.e., precursor emissions of CO, NO_x and nonmethane hydrocarbons. The correlations are small for short-lived species such as NO, propene and isoprene (−0.03, −0.06 and −0.03, respectively), reflecting that the DC8 observations when averaged over 3 min map out large spatial scales (i.e., 60 km) and thus represent air masses that have been photochemically aged for many hours. These relationships change with altitude, and the correlations decrease in value. For the 4–8 km range, where only the nitrogenous species

concentrations show the highest correlation to ozone, with NO₂ (R = 0.48), NO_y (R = 0.48), NO_z (R = 0.47), and PAN (R = 0.39). At higher altitudes (8–12 km) correlations HNO₃ (R = 0.67), NO_z (R = 0.42), NO_y (R = 0.27) and with RH (R = −0.35) and CO (R = −0.24) are the highest.

[20] The decrease in correlations with altitude in the aggregate data reflects many factors, including the fact that the ozone relationships are driven by many processes with distinct correlations. For example stratospheric exchange and convection events result in very different and strong correlations, which show up when the data is first sorted by process, but are lost when all data representing all processes are analyzed together. The high correlations near the surface represent the dominance of the sources and photochemical production processes.

[21] The correlations of the ozone bias with respect to the biases in other species are also shown in Table 5. In general the model biases show similar relationships between species as discussed above for the ozone observations. For altitudes <1 km PAN, CO, MEK, and HNO₃ are among the most highly correlated biases with respect to the ozone bias (0.6, 0.51, 0.48, and 0.47, respectively). The correlation relationships for the biases tend to extend up to ~4 km, above which the values tend to decrease. In the upper troposphere the correlation structure is very different than that at the surface, reflecting the different processes and the strong contribution of the boundary conditions to the biases.

[22] The simple correlation analysis discussed above shows the complex relationships between ozone and the ozone biases, with those relationships in the lower troposphere reflecting the emissions and the photochemical production processes. To better understand these relations,

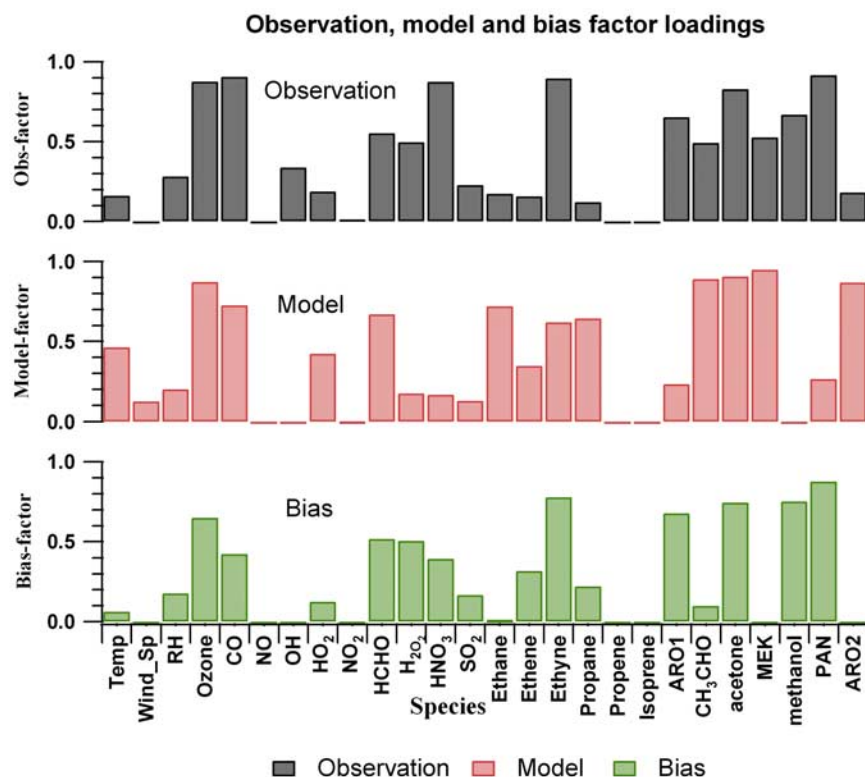


Figure 10. Factor analyses for DC-8, points in 0–1 km range. (top) Observations. (middle) Model. (bottom) Bias of model with respect to observations. Factor criteria are Eigen value one, varimax rotation.

factor analysis was performed on the observational data, on the model predicted values, and on the biases. Factor Analysis (FA) is essentially a variable reduction of data sets consisting of large number of intercorrelated variables into a small number of factors, which account for most of the variance in the original variables [Kulkarni, 2004]. This technique is frequently used in air quality studies to unravel the hidden source information from a rich ambient data set. It is particularly very useful for source apportionment studies when there is no prior information available about the nature of the major aerosol sources affecting a particular receptor station [Seinfeld and Pandis, 1998]. Factor Analysis is extremely useful in identifying the relationships among variables that are driven by common processes such as sources, transport, and chemistry. In addition, the Factor Analysis approach is unaffected by errors in modeled emissions, chemistry, or transport [Millet *et al.*, 2006]. In this section, the results obtained by the application of the Factor Analysis to the observed values, the corresponding model predictions and the model errors for the DC-8 at altitudes below 1 km are presented. The main objective of this analysis is to identify the underlying relationships between ozone and its other precursor species. All the analyses presented in this section were performed using SAS 9.0 software [SAS Institute, 2004]. All rows with missing values were deleted prior to performing the analysis, so all analysis uses identical sampling. The factors were extracted using Eigen value one criterion and the extracted factors were subject to the varimax rotation.

[23] Figure 10 depicts the output obtained from FA analysis. We only show the factor that accounts for the highest variance among the retained factors in each analysis here. Figure 10 (top) in gray shows the factor for the observed species. This factor contains a spectrum of species related to ozone and its precursors, and clustering together those photochemical factors identified in Table 5. Figure 10 (middle) in red shows the output from the same analysis conducted using the predicted values (NEI2001-Frost LPS*). In general the factors identified by the model predictions show many similarities to those based on observations suggesting that the modeled processes are capturing many of the ozone relationships in the real atmosphere. Figure 10 (bottom) shows the output of the factor analysis performed for the model errors. The clustering of errors shows a structure similar to that for the species dependencies. Highest factor loadings appear for primary species such as CO, ethyne and ARO1, as well as for secondary species (the largest factor loading is for PAN). These underlying error structures provide guidance into further model improvements, such as continued improvements in emissions, and improvements in the chemical and physical processes controlling key species such as PAN and HCHO. Furthermore this error structure may also be useful in better definition in the error covariance estimates needed by modern data assimilation techniques as discussed by Chai *et al.* [2007].

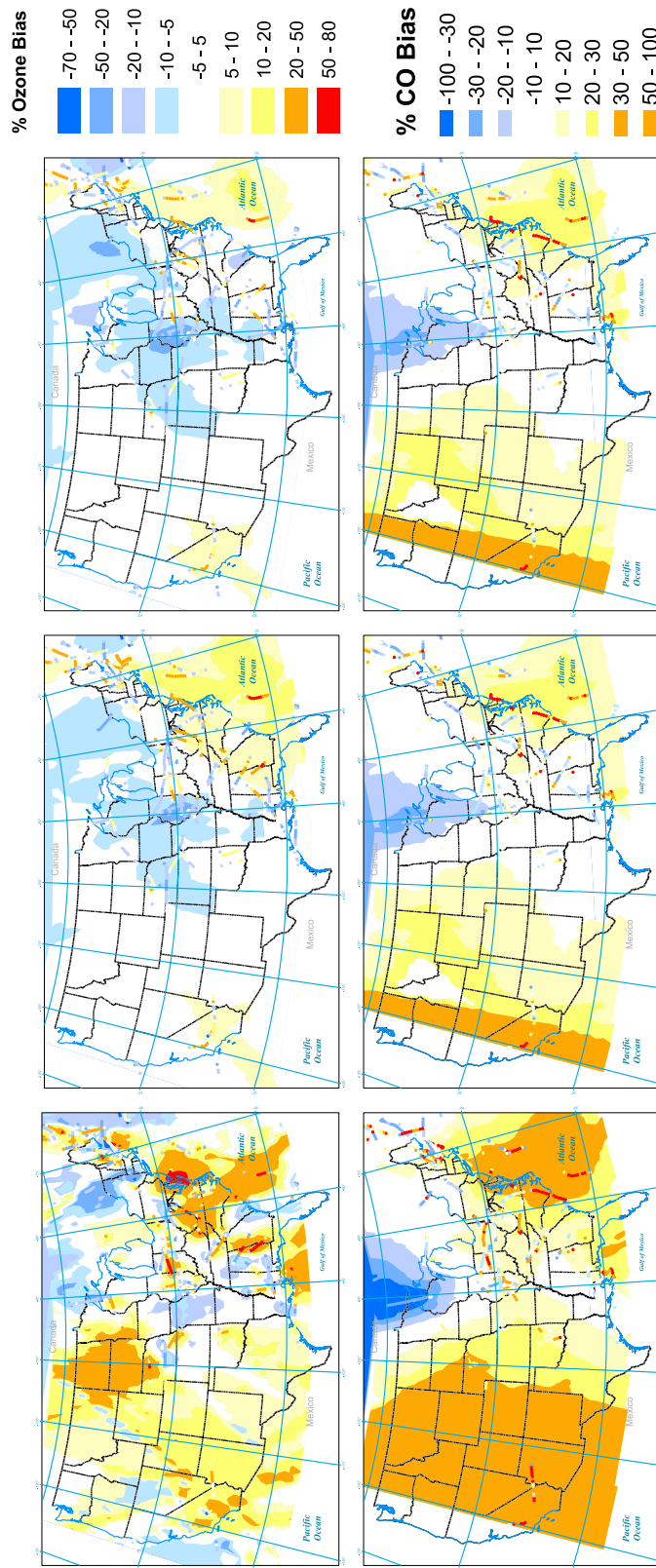


Figure 11. Kriged ozone (n = 1208) and CO (n = 1001) percent bias (modeled-observed) for alt <4000 m, DC-8 platform. (top) Ozone. (bottom) CO. (left) Forecast, NEI 2001-Frost LPS. (middle) NEI 2001-Frost LPS*. (right) NEI 2001-Frost LPS*.

4.2. Bias in a Regional Context

[24] Geographical context is given to the point bias estimations (modeled-observed) by interpolating them through kriging, generating a continuous surface. Data was restricted to the 0–4 km range, for all DC-8 flights. The previous section suggested which variables need to be improved to lower ozone bias. The interpolated bias surface gives guidance toward where model bias needs to be improved, and qualitatively how changes in modeled inputs affect the bias. The surface of ozone bias is shown in Figure 11. For the forecast (Figure 11, left) we observe that during the forecast there was a positive bias in the central and eastern United States with bias in the range of 10–50%. The biases in CO, NO_y, NO_x, HNO₃, and PAN, which showed strong correlation with ozone bias as discussed previously, were also analyzed. Wherever ozone presented a positive bias in the forecast CO (Figure 11, left), NO_x, NO_y, and HNO₃ (Figure 12, left) also presented positive biases. This is particularly clear for NO_y (Figure 12, middle left) where the bias in some regions of Ohio, North Carolina and Virginia are as large as 300–400%. When the emission inventories were modified, updating to NEI 2001 and the large point sources (Frost LPS), the ozone bias decreased to ~5–10% across the domain (Figure 11, top middle). The bias in CO was reduced through the continental United States, along with the bias of NO_y. However, a positive systematic bias persisted for ozone and NO_y, and its components, in portions of the domain.

[25] The effect of changes in the dry deposition velocities is also apparent for the bias plots for ozone and HNO₃. In the regions dominated by agricultural crops (i.e., the central United States) the biases of ozone and HNO₃ are significantly reduced. For those species with low dry deposition rates the effect on bias reductions are small (CO and NO_x). For example, North and South Dakota, regions with negligible changes in NO_x emissions between NEI 1999 and 2001 where ozone bias decreased from 20–50% to –5 to 5%.

[26] As a sensitivity study aimed to further reduce the bias, an additional simulation was conducted with a 60% decrease of NO_x area emissions (12% reduction of total NO_x emissions) for Alabama, Mississippi, Georgia, North Carolina, South Carolina, Tennessee, West Virginia, Indiana, and Ohio, chosen since these states presented the highest bias in NO_y, NO_x, and HNO₃ according to the kriging results. The results of this run showed enhancement of correlation factors for nitrogen species (Table 3 and 4), particularly for HNO₃, and NO_y, while decreasing their positive bias. As shown in Figure 11 (top right), the ozone bias decreased to a range of –10 to 10% for most of the continent, with large portions showing bias in the –5 to 5% range. The offshore ozone positive bias persists, but to lesser geographical extent, and lower magnitude than the NEI 2001 scenario, and the forecast scenario. NO_y for this same scenario decreases its regional bias to less than 100% over large portions of the domain (Figure 12, middle). Figure 12 (bottom right) shows that HNO₃ bias also decreases significantly with the updated and modified emissions, with some areas presenting a negative bias. Figure 12 (top right) shows that NO_x bias decreased in South Carolina, North Carolina, and Virginia from 200–

400% to –30 to 50%, in accordance to locations where the ozone bias was decreased.

[27] Special notice must be taken of Figure 11 (bottom), which shows that forecast CO, using NEI 1999 presented positive biases of 30–50% for the western portion of the United States, which decreased to 20–30% using the NEI 2001. Negative CO bias over Michigan decreased from –30 to –20% to –20 to –10% as a result of improved boundary conditions. The offshore Atlantic positive CO bias decreased from 30–50% to the 10–30% range and from 10–20% to –10 to 10% for the southeastern United States. These changes are due largely to changes in emissions as this region is largely under outflow conditions for the period of study.

5. Discussion

5.1. Sensitivity Analysis

[28] As illustrated in Figure 2 a variety of sensitivity studies were performed to investigate the sensitivity of the model to other important parameters. The most significant parameters studied in terms of reduction of model bias, dry deposition, emissions, and boundary conditions, have been discussed previously. In terms of near surface ozone the largest impacts are due to dry deposition velocities and emissions, with each contributing equally to the bias reduction. In the mid to upper troposphere, the boundary conditions dominated the improvements.

[29] Simulations were also performed for the eastern United States using a 12 km horizontal resolution. The biggest impact was found near the surface for the 12 km resolution. For the 0–1 km range the mean for the WP-3 increased by 3 ppb with a slight increase in correlation to (R = 0.71). For CO the 12 km resolution increased the mean value by 5 ppb and also increased the correlation coefficient. Biogenic emissions represent an additional source of uncertainty. We repeated a simulation using the BIES3 biogenic emissions algorithm, which led to higher biogenic emissions. Under these conditions the near surface ozone increased by 1 ppbv throughout the eastern United States and CO by 10 ppbv.

5.2. Ozone Production Efficiency

[30] Up to now we have shown that the ozone bias was strongly correlated to NO_z and NO_y bias. We have also discussed that there is geographical concordance of ozone bias with NO_z bias. Previous work has related ozone and NO_z [Kleinman, 2005; Trainer *et al.*, 1993], in a relationship for ozone production efficiency (OPE), which for the purposes of this analysis is the slope of the plot of odd oxygen (NO₂ + O₃) to NO_z. In Figure 13 the ozone production efficiency for the observed and predicted values for data points less than 4000 m is plotted. For the DC-8 data the observed OPE is 7.8, while the forecast, NEI2001-Frost LPS, and NEI2001-Frost LPS* cases have OPEs of 6.7, and 7.8, respectively. The observed ozone production efficiency for the DC-8 data suggests efficient formation of ozone, typical of NO_x limited conditions. The ozone production efficiency in the area sampled by WP-3 is lower than the DC-8 (Observed OPE = 3.49, Modeled OPE = 5.28), which reflects the fact that the area sampled was closer to emission source regions in the northeast.

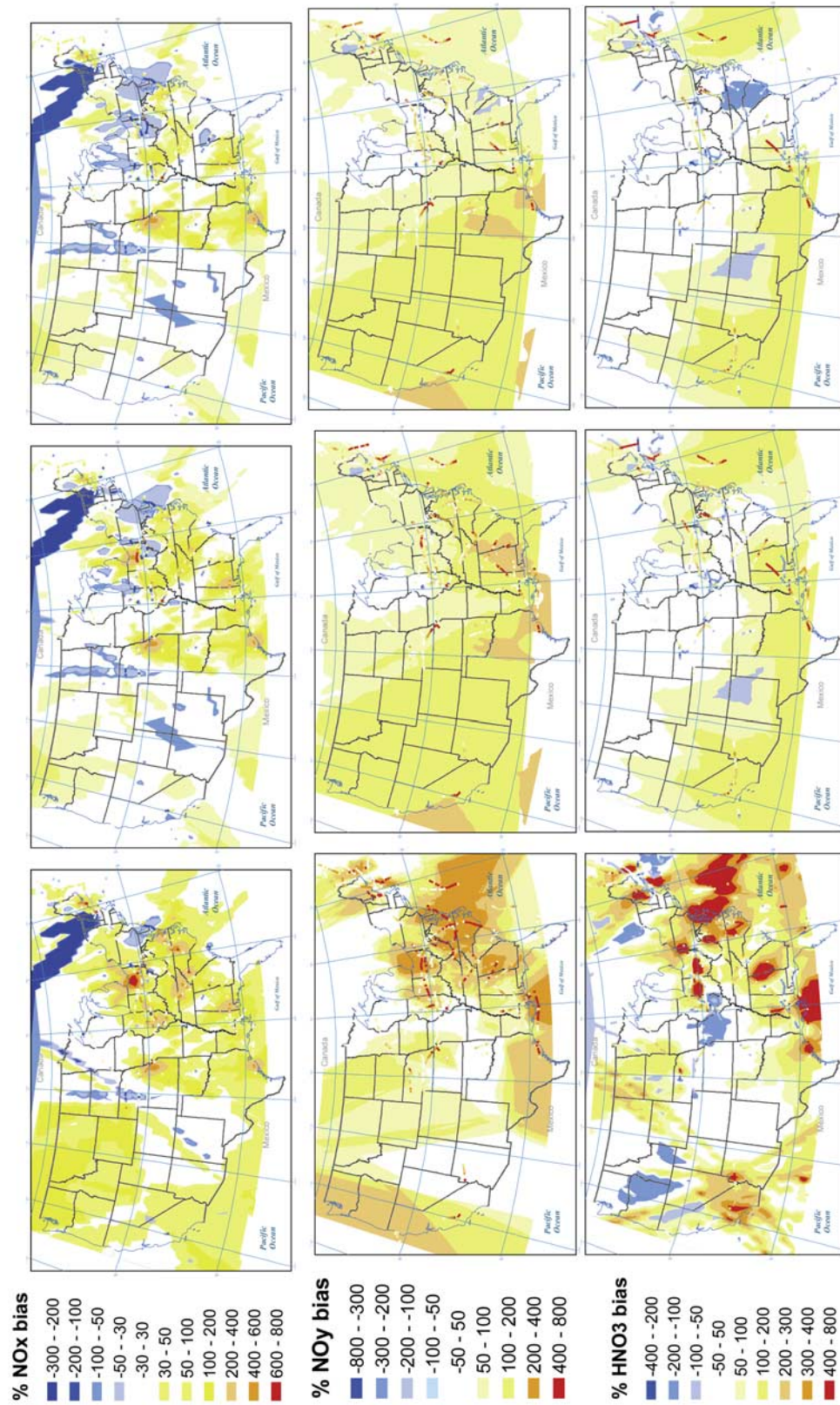


Figure 12. Kriged NO_x (n = 1028) and NO_y (n = 902) and HNO₃ (n = 1157) percent bias (modeled-observed) for alt <4000 m, DC-8 platform. (top) NO_x. (middle) NO_y. (bottom) HNO₃. (left) Forecast, NEI 1999. (middle) NEI 2001-Frost LPS. (right) NEI 2001-Frost LPS*.

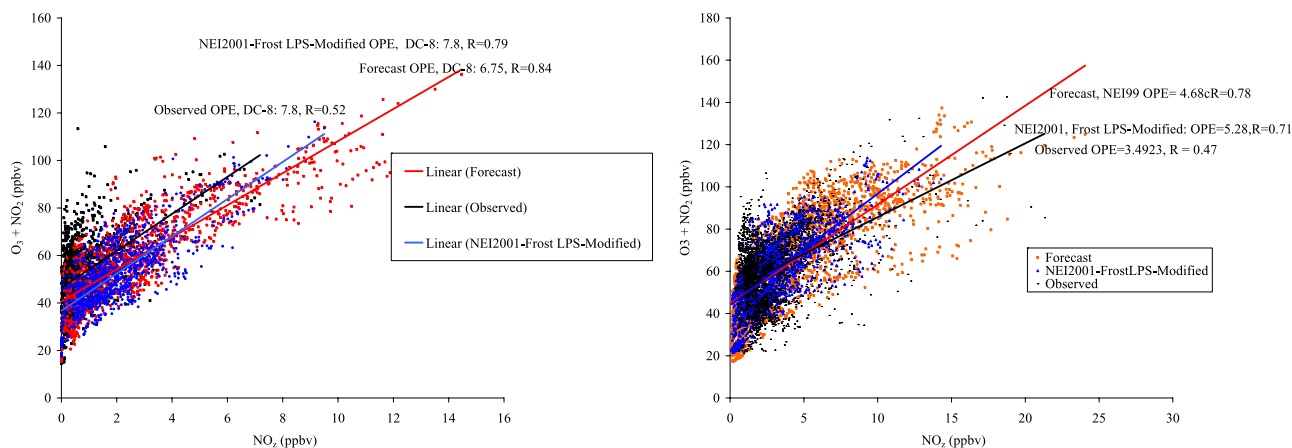


Figure 13. Observed and modeled ozone production efficiency (OPE). (left) DC-8. (right) WP-3 for data points with altitude <4000 m, all flights. Red indicates forecast, NEI 1999, and blue indicates NEI 2001-Frost LPS*.

5.3. Model Sensitivity to Emissions

[31] The fact that the modeled and observed production efficiencies are similar suggests that the underlying ozone relationships of the model are reasonably well represented, and therefore that the results of sensitivity studies may be meaningful. Further insights into ozone production can be seen by comparing the change in ozone to changes in emissions. The mean predicted near surface ozone (0–1 km) values for the NEI 2001 Frost LPS* is shown in Figure 14, in which higher values are found in California, Arizona, and the Atlantic states. The sensitivity of O_3 to emissions of precursors is also shown in Figure 15. These simulations were done for the period of 21 July to 18 August 2004, in the absence of anthropogenic VOCs, CO, and reduced NO_x . The contribution of each precursor is calculated as the difference between the average daytime surface ozone concentration in the presence of a precursor minus the concentration in the absence of the precursor, normalized by the total yearly emissions of the precursor (ppb ozone formed/(Tg precursor/year)). Ozone formation is most sensitive to changes in NO_x (as is also shown in Figure 14), especially in the Midwest, where ozone per Tg of NO_x /year increases by 10–20 ppbv (note that NO_x sensitivity was calculated on the basis of 30% reduction of total NO_x). For reference, 1 Tg of NO_x /year is roughly equivalent to the emissions of 28 coal fired power plants of 2500 MW (36,000 tonnes NO_x /year each) [Miller and Van Atten, 2004]. In the northeast United States, ozone is equally sensitive to NO_x and anthropogenic VOCs, while in large parts of the western United States, ozone is more sensitive to VOCs. CO effects are similar to VOCs, but smaller on a per Tg/year basis. Since CO emissions are larger than VOCs, total CO effects on ozone can be in the same order of magnitude. Figure 15 shows that VOCs and CO contribute to a large portion of ozone formation in portions of the northeast United States.

[32] As pointed out above, NO_x plays an important role in ozone production. Furthermore an appreciable fraction of NO_y is composed of PAN (representing $\sim 20\%$ near the surface and $\sim 50\%$ at 6–8 km altitudes, Figure 5), and ozone levels and errors were shown to be significantly

correlated with those for PAN. PAN plays important roles as both a key photochemical product and as a reservoir for NO_x . To assess the role of PAN on ozone production we conducted a simulation where PAN levels within the regional domain (but not in the boundary conditions) were continuously set to zero. In this way the formation of PAN was allowed, but the thermal decomposition source of NO_x was blocked, by setting PAN concentrations consistently to zero. The impact of PAN on predicted mean surface ozone for the month of July is shown in Figure 16 (right). This indirect ozone production pathway of PAN via production of NO_x is estimated to be over 20% throughout the continental United States with large regions with values between 30 and 50%. This impact extends to all altitudes with values exceeding 8% throughout the domain at an altitude of 5.6 km. Differences become small above this height as the PAN levels are dominated by the boundary conditions values, which were not changed, and the low rates of PAN thermal decomposition due to cool temperatures. The results point out the importance of accurately predicting PAN levels, which requires the close coupling between the regional and global models, as PAN sources and sinks reflect process occurring throughout the vertical extent of the atmosphere and over large geographic scales. Figure 16 (left) shows results for a sensitivity analysis where the formation of PAN itself is blocked. As PAN is a reservoir for NO_x under these conditions, with its formation pathway is blocked, a greater fraction of NO_x is oxidized into nitric acid, with the net effect of more ozone being formed. Mean surface ozone concentration increases up to ~ 7 ppb.

6. Conclusion

[33] The comprehensive ICARTT aircraft observations were used to evaluate and improve ozone prediction for the STEM model by simultaneously analyzing model performance with conventional statistics, mean vertical profiles, and geospatial interpolation to provide a comprehensive three dimensional context on model performance, laying the groundwork for subsequent model improvement.

[34] The STEM model forecasts of ozone were found to have a significant positive high (overprediction) bias near

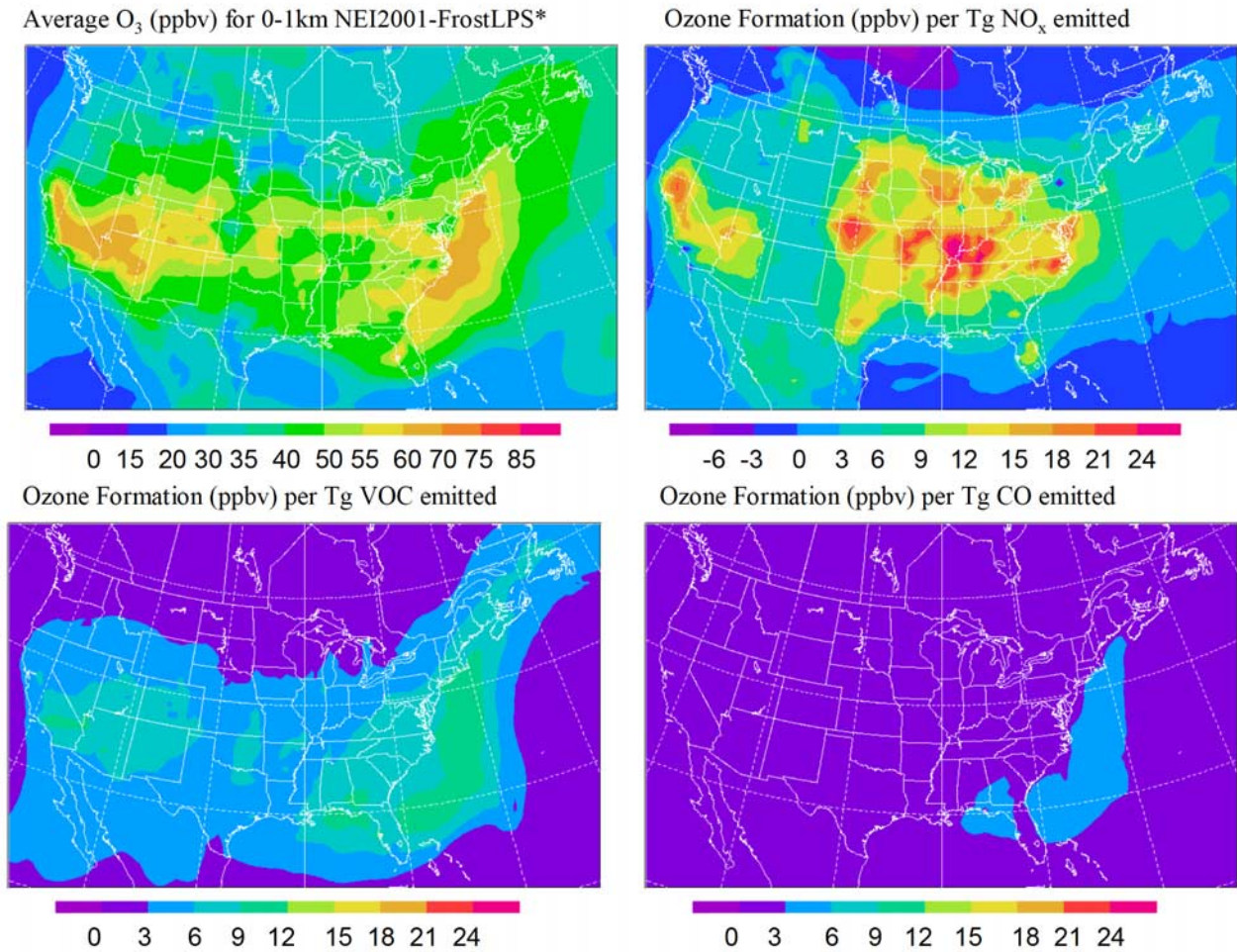


Figure 14. (top left) Modeled near surface ozone (0–1 km average) for NEI-2001-FrostLPS* case. Modeled net ozone formation (ppbv/(Tg of precursor/year)): (top right) NO_x and (bottom left) anthropogenic VOCs emissions. (bottom right) Anthropogenic CO. Study period is 21 July to 18 August 2004.

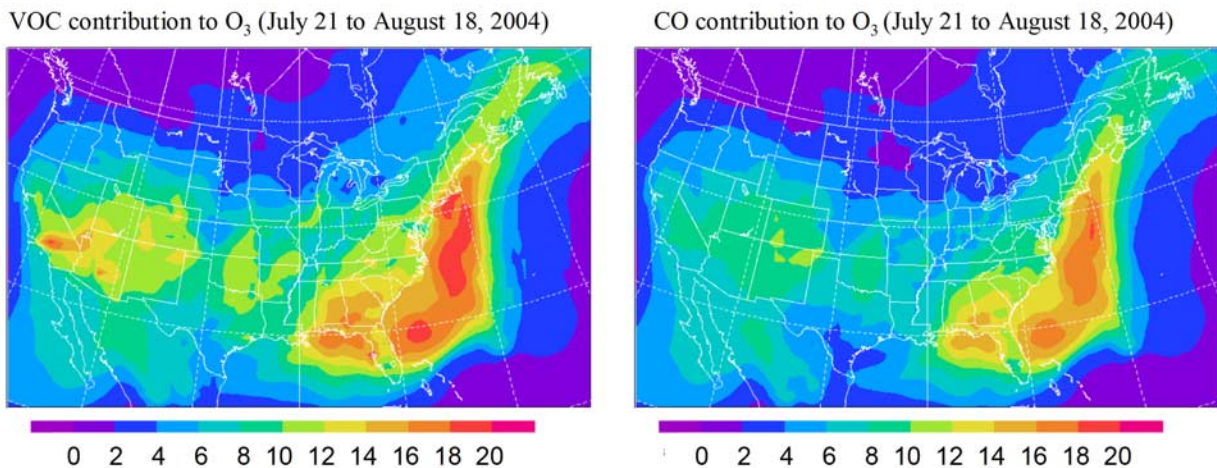


Figure 15. Average surface ozone contribution (ppbv) due to (left) anthropogenic VOCs and (right) anthropogenic CO, calculated as the difference between average 0–1 km ozone for NEI 2001-FrostLPS*, and the same in scenario in the absence of VOCs and CO, respectively. Study period is 21 July to 18 August 2004.

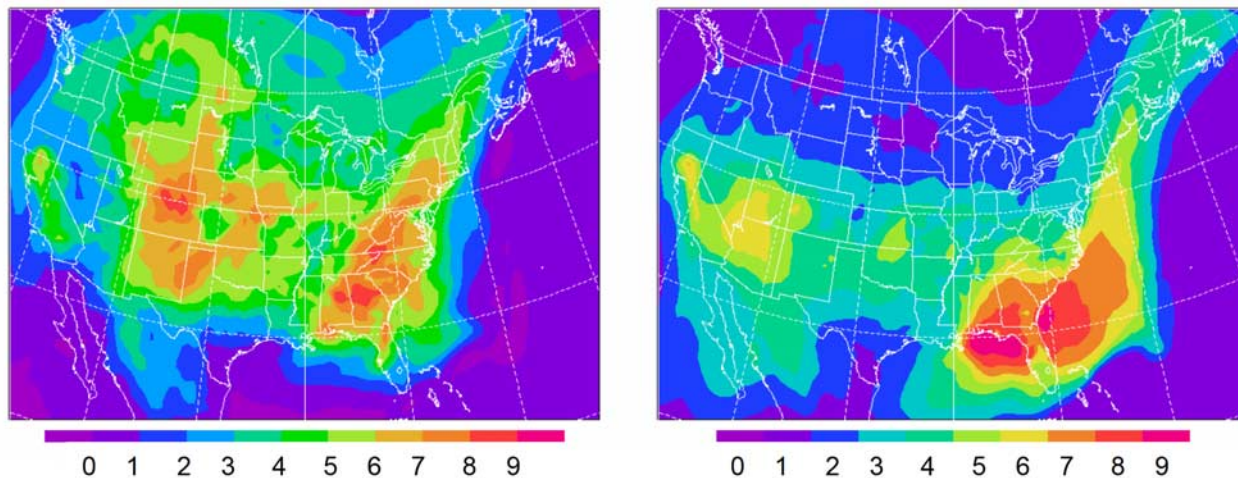


Figure 16. Calculated impact of the thermal decomposition of PAN on ozone. (left) Average difference between surface ozone with the formation of PAN and with the formation of PAN blocked. (right) Average difference between surface ozone concentration with unconstrained PAN and with PAN concentrations set to zero. Study period is 21 to 28 July 2004.

the surface and a large negative bias in the upper troposphere. These biases were due in part to model errors, which included the use of outdated emissions in the forecasts (i.e., the use of NEI 1999 emissions), out of season dry deposition velocities for agricultural crops, and the use of global model boundary conditions (BCs) that were based on climatological biomass burning emissions. Postanalysis simulations were conducted using corrected dry deposition velocities, NEI 2001 inventory with point source emissions updated to 2004, and BCs from global models using biomass burning emissions reflective of 2004 fire activity. These changes resulted in a decrease in low altitude (0–1 km) mean ozone bias from 11.21 to 1.45 ppbv in comparison to DC-8 observations and from 8.26 to -0.34 ppbv for the WP-3 data. The upper troposphere ozone negative bias persisted, but was reduced in magnitude. In addition, the postanalysis simulations resulted in a nearly unbiased distribution of ozone errors with a significant reduction in the standard deviation of ozone errors.

[35] A series of analyses were performed to study the structure of the ozone errors. These included the correlation of ozone errors with errors in other species. In general the model ozone errors show high correlation for altitudes <1 km with the errors in PAN, CO, MEK, and HNO_3 . These correlation relationships for the biases tended to extend up to ~ 4 km, above which the values decreased. To better understand the relationships among these errors, factor analysis was applied. The clustering of errors shows a structure similar to that for the species dependencies, and included contributions from meteorological features of temperature and RH, precursor species such as CO, ethyne and aromatics, as well as photochemical products such as HNO_3 , PAN, H_2O_2 and HCHO.

[36] The spatial distribution of the model errors were examined using Kriging. The geospatial distribution of forecast errors identified distinct features, which when compared to the errors in the postanalysis simulations,

helped document model improvements. For example, changes in the dry deposition to crops was shown to reduce substantially the high bias in the forecasts in the Midwest, while the updated emissions accounted for the decrease in bias in the eastern United States. In addition, improvements in boundary conditions from global models, which accounted for biomass burning emissions, improved model performance for CO in the upper troposphere, in comparison to the forecast stage.

[37] Nitrogen species, namely NO_y , NO_z , and HNO_3 showed positive bias during forecast stage, which decreased during postanalysis. Reductions in these biases due largely to emission changes resulted in reduction of ozone bias, especially in the 0–4 km range. However, a persistent high NO_y bias suggests that the NEI 2001 NO_x emissions are still overestimated. Kriging was shown to provide a geospatial analysis of these biases, and to provide information that can help guide further regional modification of emissions.

[38] Predicted ozone production efficiency (OPE) was studied and shown to be similar to the observed OPE (calculated values of OPE of 7.83 in comparison to observed OPE of 7.79 for the DC-8). The OPE for the WP-3 (observed 3.49, modeled 5.28) was lower than for the DC-8, suggesting that the area sampled by the WP-3 was closer to the source regions of NO_x .

[39] Simulations with perturbed emissions found ozone formation to be most sensitive to changes in NO_x , especially in the Midwest, where ozone per Tg of NO_x /year increases by 10–20 ppbv. However, in the northeast United States, ozone was found to be equally sensitive to NO_x and anthropogenic VOCs, while in large parts of the western United States, ozone was more sensitive to VOCs. CO effects were found to be similar to those for VOCs, but smaller on a per Tg/year basis. Since CO emissions are larger than VOCs, total CO effects on ozone can be in the same order of magnitude.

[40] The results presented here help demonstrate how more detailed analysis of errors can improve model performance. These results also point out that the underlying error structure is complicated, but that the underlying error structure can provide guidance into further model improvements, such as continued improvements in emissions, and improvements in the chemical and physical processes controlling key species such as PAN and HCHO. Furthermore this error structure may also be useful in better definition in the error covariance estimates needed by modern data assimilation techniques.

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References

- Bates, T. S., B. J. Huebert, J. L. Gras, F. B. Griffiths, and P. A. Durkee (1998), International Global Atmospheric Chemistry (IGAC) Project's First Aerosol Characterization Experiment (ACE 1): Overview, *J. Geophys. Res.*, *103*(D13), 16,297–16,318.
- Blond, N., L. Bel, and R. Vautard (2003), Three-dimensional ozone data analysis with an air quality model over the Paris area, *J. Geophys. Res.*, *108*(D23), 4744, doi:10.1029/2003JD003679.
- Carmichael, G. R., L. K. Peters, and R. D. Saylor (1991), The STEM-II regional scale acid deposition and photochemical oxidant model—I. An overview of model development and applications, *Atmos. Environ.*, *25*(10), 2077–2090.
- Carmichael, G. R., et al. (2003), Evaluating regional emission estimates using the TRACE-P observations, *J. Geophys. Res.*, *108*(D21), 8810, doi:10.1029/2002JD003116.
- Carter, W. P. L. (2000), Documentation of the SAPRC-99 chemical mechanism for VOC reactivity assessment, final report to California Air Resources Board, Contract 92-329, Univ. of Calif., Riverside, 8 May.
- Chai, T., G. R. Carmichael, Y. Tang, A. Sandu, and D. G. Streets (2007), Four-dimensional data assimilation experiments with International Consortium for Atmospheric Research on Transport and Transformation ozone measurements, *J. Geophys. Res.*, doi:10.1029/2006JD007763, in press.
- Frost, G. J., et al. (2006), Effects of changing power plant NO_x emissions on ozone in the eastern United States: Proof of concept, *J. Geophys. Res.*, *111*, D12306, doi:10.1029/2005JD006354.
- Geron, C., A. Guenther, and T. Pierce (1994), An improved model for estimating emissions of volatile organic compounds from forests in the eastern United States, *J. Geophys. Res.*, *99*, 12,773–12,792.
- Grell, G., J. Dudhia, and D. R. Stauffer (1995), A description of the Fifth Generation Penn State/NCAR Mesoscale Model (MM5), *NCAR Tech. Note, NCAR/TN-398+STR*, Natl. Cent. for Atmos. Res., Boulder, Colo. (Available at <http://cng.ateneo.net/cng/wyu/resources/mm5/desc/cover.pdf>)
- Hong, S.-Y., and H.-L. Pan (1996), Nonlocal boundary layer vertical diffusion in a medium-range forecast model, *Mon. Weather Rev.*, *124*, 2322–2339.
- Horowitz, L. W., et al. (2003), A global simulation of tropospheric ozone and related tracers: Description and evaluation of MOZART, version 2, *J. Geophys. Res.*, *108*(D24), 4784, doi:10.1029/2002JD002853.
- Huang, H.-M. H., S.-Y. Hong, and M. Kanamitsu (1997), The NCEP Regional Spectral Model: Update, *Bull. Am. Meteorol. Soc.*, *78*(10), 2125–2143.
- Jacob, D. J., J. H. Crawford, M. M. Kleb, V. S. Connors, R. J. Bendura, J. L. Raper, G. W. Sachse, J. C. Gille, L. Emmons, and C. L. Heald (2003), Transport and Chemical Evolution over the Pacific (TRACE-P) aircraft mission: Design, execution, and first results, *J. Geophys. Res.*, *108*(D20), 9000, doi:10.1029/2002JD003276.
- Kiley, C. M., et al. (2003), An intercomparison and evaluation of aircraft-derived and simulated CO from seven chemical transport models during the TRACE-P experiment, *J. Geophys. Res.*, *108*(D21), 8819, doi:10.1029/2002JD003089.
- Kim, Y. P., and J. H. Seinfeld (1995), Atmospheric gas-aerosol equilibrium III: Thermodynamics of crustal elements Ca²⁺, K⁺, Mg²⁺, *Aerosol Sci. Technol.*, *22*, 93–110.
- Kim, Y. P., J. H. Seinfeld, and P. Saxena (1993a), Atmospheric gas-aerosol equilibrium I: Thermodynamic model, *Aerosol Sci. Technol.*, *19*, 151–181.
- Kim, Y. P., J. H. Seinfeld, and P. Saxena (1993b), Atmospheric gas-aerosol equilibrium II: Analysis of common approximations and activity coefficient calculation methods, *Aerosol Sci. Technol.*, *19*, 182–198.
- Kim, S.-W., A. Heckel, S. A. McKeen, G. J. Frost, E.-Y. Hsie, M. K. Trainer, A. Richter, J. P. Burrows, S. E. Peckham, and G. A. Grell (2006), Satellite-observed U. S. power plant NO_x emission reductions and their impact on air quality, *Geophys. Res. Lett.*, *33*, L22812, doi:10.1029/2006GL027749.
- Kleinman, L. I. (2005), The dependence of tropospheric ozone production rate on ozone precursors, *Atmos. Environ.*, *39*, 575–586.
- Kulkarni, S. (2004), Surface elemental composition of aerosols at Beijing (China), Gosan (Korea), and Tango (Japan) during ACE-Asia, M. S. Thesis, 183 leaves, Chem. and Biochem. Eng., Univ. of Iowa, Iowa City.
- Liao, D., D. Peuquet, Y. Duan, E. Whitsel, J. Dou, R. Smith, H.-M. Lin, J.-C. Chen, and G. Heiss (2006), GIS approaches for the estimation of residential level ambient PM concentrations, *Environ. Health Perspect.*, *114*(9), 1374–1380.
- McKeen, S., J. Wilczak, G. Grell, S. Peckham, and M. Pagowski (2006), Real-time ozone forecasts over eastern North America during the summer of 2004: An assessment of several models and their ensemble, *Geophys. Res. Abstr.*, *7*, 1–2.
- Menu, L., et al. (2000), Atmospheric pollution over the Paris area: The ESQUIF project, *Ann. Geophys.*, *18*, 1467–1481.
- Miller, P. J., and C. Van Atten (2004), North American power plant emissions, Comm. for the Environ. Coop. of N. Am., Montreal, Que., Canada.
- Millet, D. B., et al. (2006), Chemical characteristics of North American surface layer outflow: Insights from Chebogue Point, Nova Scotia, *J. Geophys. Res.*, *111*, D23S53, doi:10.1029/2006JD007287.
- Oliver, M. A., and R. Webster (1990), Kriging: A method of interpolation for geographical information systems, *Int. J. Geogr. Inf. Syst.*, *4*(3), 313–332.
- Parrish, D. D. (2006), Critical evaluation of US on-road vehicle emission inventories, *Atmos. Environ.*, *40*, 2288–2300.
- Pfister, G., P. G. Hess, L. K. Emmons, J.-F. Lamarque, C. Wiedinmyer, D. P. Edwards, G. Pétron, J. C. Gille, and G. W. Sachse (2005), Quantifying CO emissions from the 2004 Alaskan wildfires using MOPITT CO data, *Geophys. Res. Lett.*, *32*, L11809, doi:10.1029/2005GL022995.
- Ramana, M. V., V. Ramanathan, I. A. Podgorny, B. B. Pradhan, and B. Shrestha (2004), The direct observations of large aerosol radiative forcing in the Himalayan region, *Geophys. Res. Lett.*, *31*, L05111, doi:10.1029/2003GL018824.
- SAS Institute (2004), SAS/STAT 9.1 user's guide, Cary, N. C.
- Seinfeld, J. H., and S. N. Pandis (1998), *Atmospheric Chemistry and Physics: From Air Pollution to Climate Change*, John Wiley, New York.
- Singh, H. B., W. H. Brune, J. H. Crawford, D. J. Jacob, and P. B. Russell (2006), Overview of the summer 2004 Intercontinental Chemical Transport Experiment—North America (INTEX-A), *J. Geophys. Res.*, *111*, D24S01, doi:10.1029/2006JD007905.
- Tang, Y., et al. (2003), Impacts of aerosols and clouds on photolysis frequencies and photochemistry during TRACE-P: 2. Three-dimensional study using a regional chemical transport model, *J. Geophys. Res.*, *108*(D21), 8822, doi:10.1029/2002JD003100.
- Tang, Y., et al. (2004), Three-dimensional simulations of inorganic aerosol distributions in east Asia during spring 2001, *J. Geophys. Res.*, *109*, D19S23, doi:10.1029/2003JD004201.
- Tang, Y., et al. (2007), Influence of lateral and top boundary conditions on regional air quality prediction: A multiscale study coupling regional and global chemical transport models, *J. Geophys. Res.*, *112*, D10S18, doi:10.1029/2006JD007515.
- Trainer, M., et al. (1993), Correlation of ozone within NO_y in photochemically aged air, *J. Geophys. Res.*, *98*(D2), 2917–2925.
- U.S. Department of Transportation (2006), Transportation air quality: Selected facts and figures, *Fed. Highway Admin. Off. of Nat. and Human Environ. Publ. FHWA-HEP-05-045 HEP/12-05 (8M)E*, Washington, D. C.
- U.S. Environmental Protection Agency (1999), Report of the December 15, 1999 EPA Satellite Forum on Ozone Monitoring, Mapping, and Public Outreach, Cincinnati, Ohio. (Available at <http://epa.gov/empact/techtransfer/satellite.htm>)
- U.S. Environmental Protection Agency (2006), 1999 National Emission Inventory, Research Triangle Park, N. C. (Available at <http://www.epa.gov/ttn/chief/net/1999inventory.html#final3crit>)
- Vautard, R., et al. (2003), A synthesis of the Air Pollution Over the Paris Region (ESQUIF) field campaign, *J. Geophys. Res.*, *108*(D17), 8558, doi:10.1029/2003JD003380.

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