Characterization of NO$_x$, SO$_2$, ethene, and propene from industrial emission sources in Houston, Texas


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[1] The Houston-Galveston-Brazoria urban area contains industrial petrochemical sources that emit volatile organic compounds and nitrogen oxides, resulting in rapid and efficient ozone production downwind. During September to October 2006, the NOAA WP-3D aircraft conducted research flights as part of the second Texas Air Quality Study (TexAQS II). We use measurements of NO$_x$, SO$_2$, and speciated hydrocarbons from industrial sources in Houston to derive source emission ratios and compare these to emission inventories and the first Texas Air Quality Study (TexAQS) in 2000. Between 2000 and 2006, NO$_x$/CO$_2$ emission ratios changed by an average of $-29\%\pm20\%$, while a significant trend in SO$_2$/CO$_2$ emission ratios was not observed. We find that high hydrocarbon emissions are routine for the isolated petrochemical facilities. Ethene (C$_2$H$_4$) and propene (C$_3$H$_6$) are the major contributors to ozone formation based on calculations of OH reactivity for organic species including C$_2$–C$_{10}$ alkanes, C$_2$–C$_5$ alkenes, ethyne, and C$_2$–C$_5$ aldehydes and ketones. Measured ratios of C$_2$H$_4$/NO$_x$ and C$_3$H$_6$/NO$_x$ exceed emission inventory values by factors of 1.4–20 and 1–24, respectively. We examine trends in C$_2$H$_4$/NO$_x$ and C$_3$H$_6$/NO$_x$ ratios between 2000 and 2006 for the isolated petrochemical sources and estimate a change of $-30\%\pm30\%$, with significant day-to-day and within-plume variability. Median ambient mixing ratios of ethene and propene in Houston show decreases of $-52\%$ and $-48\%$, respectively, between 2000 and 2006. The formaldehyde, acetaldehyde, and peroxyacetyl nitrate products produced by alkene oxidation are observed downwind, and their time evolution is consistent with the rapid photochemistry that also produces ozone.


1. Introduction

[2] Ozone (O$_3$) is the most abundant oxidant in the troposphere and can be considered a principal product of tropospheric chemistry. It is formed by photochemical reactions involving nitrogen oxides (NO$_x$) and reactive volatile organic compounds (VOCs) [Haagen-Smit, 1952; Crutzen, 1979]. Reduction of NO$_x$ and VOC emissions under the U. S. Clean Air Act of 1970 and its subsequent amendments has been partially successful in reducing ozone concentrations within the United States, but understanding and predicting exceedances remain a challenge.

[3] During daytime conditions, VOCs react primarily with hydroxyl radicals (OH). The resulting peroxy radicals can react with nitric oxide (NO) to regenerate OH and form nitrogen dioxide (NO$_2$). The rate and efficiency of ozone production depends on the absolute concentrations of NO$_x$ and VOCs, as well as the ratio of these species. Ambient measurements have confirmed that the rate and magnitude of ozone production differs for plumes downwind of different anthropogenic source types characterized by different NO$_x$ and VOC emission ratios [e.g., Gillani et al., 1998; Ryerson et al., 1998; Daum et al., 2000; Ryerson et al., 2001, 2003]. The fastest rate of ozone formation and highest yields per NO$_x$ molecule emitted are predicted for conditions where elevated concentrations of NO$_x$ and reactive VOCs are simultaneously present.

[4] The city of Houston, Texas, is the fourth largest in the United States, and the greater Houston urban area is home to
more than five million people [U. S. Census Bureau, 2006]. Since 2004, Houston-Galveston-Brazoria has been designated as a nonattainment area under the 1997 National Ambient Air Quality Standards (NAAQS) for ozone. The stricter 2008 NAAQS rules require that the 3 year average of the fourth highest daily maximum 8 h average ozone concentrations measured at each monitoring site within an area over each year must not exceed 0.075 ppm. The high ozone levels in Houston are attributed to anthropogenic NOₓ and VOC sources that combine to form ozone during periods of intense solar radiation and stagnant meteorological conditions. In addition to the typical urban emission sources, such as motor vehicles, the greater Houston area is home to the largest grouping of petrochemical industrial sources in the United States. However, the problems of petrochemical VOC emissions are not unique to Houston. Many cities contain industrial petrochemical sources, albeit on a smaller scale, and Houston serves as an important case study.

In order to evaluate and predict air quality in the Houston area, it is important to understand the magnitude and variability of these industrial sources and how these sources are evolving over time. Previous studies in Houston reported high concentrations of VOCs that were not correctly included in emission inventories [Kleinman et al., 2002; Karl et al., 2003; Ryerson et al., 2003; Wert et al., 2003; Jobson et al., 2004; de Gouw et al., 2009; Mellqvist et al., 2010]. Although concentrations of all measured VOCs exceeded values in emission inventories, the reactivity was dominated by a limited subset of highly reactive VOCs that included ethene and propene [Ryerson et al., 2003; Wert et al., 2003]. The high ozone concentrations observed in Houston were attributed to the presence of these highly reactive VOCs in a NOₓ-rich atmosphere. Biogenic sources of VOCs also exist in and around Houston, although detailed vegetation surveys [Wiedinmyer et al., 2001], aircraft measurements [Wert et al., 2003], and ship-based measurements [Gilman et al., 2009] indicate that these emissions are less important for ozone formation than those from anthropogenic sources near the Houston Ship Channel.

One of the primary goals of the second Texas Air Quality Study (TexAQS II) in 2006 was to continue the work begun during the TexAQS 2000 study to improve understanding of the chemical and meteorological causes of ozone exceedance episodes in the Houston-Galveston-Brazoria and Dallas metropolitan areas. A second important goal was to evaluate potential changes in ozone precursor emissions between 2000 and 2006. During September to October 2006, measurements of gas phase species and particles were made by the NOAA WP-3D aircraft in plumes from power plant, petrochemical, and urban sources in Texas and neighboring states. Further description of the scientific goals and logistics of the TexAQS 2006 field campaign can be found in the work of Parrish et al. [2009].

[7] This work focuses on the emissions of NOₓ, sulfur dioxide (SO₂), and VOCs from industrial sources in Houston, Texas. We examine trends in NOₓ and SO₂ emissions between 2000 and 2006 for industrial sources. Using relative OH reactivity, we identify the highly reactive VOCs that are likely to contribute to ozone formation. We then present emission trends for ethene and propene between 2000 and 2006. Finally, we examine the photochemical oxidation products measured in downwind transects from industrial sources.

2. Experimental

2.1. Overview of TexAQS Field Measurements

During the TexAQS 2006 study, the NOAA WP-3D was based at Ellington Field in Houston, Texas. The aircraft conducted 18 research flights between 31 August and 13 October 2006 in eastern Texas. Each research flight was approximately 6 h in duration. The flights focused on measurements of anthropogenic pollution during active photochemical periods, including downwind of urban areas, power plants, and petrochemical industrial facilities in the greater Houston area. Most data were acquired within the planetary boundary layer during daytime, with typical flight altitudes of 500 m above sea level for the plume transect measurements presented here. Plume transects were sampled by flying perpendicular to the prevailing wind direction under conditions when emissions from industrial complexes could be individually distinguished.

For the measurements used in this analysis, Table 1 summarizes the instrumental techniques, uncertainty, frequency, and references containing further details. The stated uncertainty in Table 1 includes both precision and accuracy. Instruments onboard the NOAA WP-3D included 1 Hz measurements of O₃, NO, NO₂, peroxyacetyl nitrates (PANs), total reactive nitrogen (NOₓ), carbon monoxide (CO), SO₂, and carbon dioxide (CO₂) [Ryerson et al., 1998, 1999;
Holloway et al., 2000; Ryerson et al., 2000; Slusher et al., 2004; Peischl et al., 2010]. In addition, several instruments measured VOCs. Formaldehyde (CH2O) was measured at 1 Hz by difference frequency generation absorption spectroscopy [Weibring et al., 2007]. Ethene (C2H4) was measured at 5 s resolution with laser photoacoustic spectroscopy (LPAS) [de Gouw et al., 2009]. Speciated VOCs were measured by both proton transfer reaction mass spectrometry (PTR-MS) [de Gouw and Warneke, 2007] and gas chromatography (GC) of 80 whole air samples acquired during each flight [Schauffler et al., 1999, 2003]. The PTR-MS measured a select set of VOCs, including acetaldehyde (CH3CHO), acetic acid, acetone, acetonitrile, benzene, toluene, C8 and C9 aromatics, methyl ethyl ketone, methanol, isoprene and its oxidation products (methyl vinyl ketone and methacrolein), and the sum of monoterpenes. The whole air sample measurements were less frequent but included a much larger suite of speciated VOCs, including C2–C10 alkanes, C2–C4 alkenes, ethyne, and C2–C5 aldehydes and ketones (see Ryerson et al. [2003] for a complete list), along with CO and methane (CH4).

[10] During the TexAQS 2000 study, the National Center for Atmospheric Research L-188C Electra aircraft was used. This aircraft was similarly based at Ellington Field in Houston, Texas, and conducted research flights during 18 August to 13 September 2000. A NOAA WP-3D transit flight south of Houston in 2002 provided additional data to construct trends. Both of these data sets have been described in detail previously [Ryerson et al., 2003]. The 2000, 2002, and 2006 aircraft campaigns used many of the same instruments, including O3, NO, NO2, NOx, CO, SO2, and whole air sampler, which minimizes measurement bias in the observed temporal trends.

2.2. Meteorological and Ancillary Data

[11] During the 2000, 2002, and 2006 field campaigns, aircraft instruments recorded wind speed, wind direction, ambient temperature, and other meteorological parameters. Meteorological and chemical measurements acquired during vertical profiles were used to determine information about boundary layer height and vertical mixing within and above the boundary layer. In addition, the Radar Wind Profiler Network measured vertical profiles of boundary layer winds at 6 Texas sites in 2000 [Nielsen-Gammon et al., 2008] and 10 Texas sites in 2006 [Parrish et al., 2009]. Vertical mixing and boundary layer height were determined for the sites closest to Houston, which were Ellington Field in 2000 and La Porte in 2006. Uncertainties in wind speed of ±1 m s−1 and in boundary layer height of ±10% were previously estimated by comparing derived values from the various aircraft and ground-based data sets [Ryerson et al., 2003].

2.3. Emission Inventories

[12] Emission estimates derived from measurements are compared here to Point Source Emissions Inventories (PSEI) for 1999 and 2006. The 1999 inventory was previously provided by the Texas Natural Resource Conservation Commission (as described in the work of Ryerson et al. [2003]), and the 2006 inventory was provided by the Texas Commission on Environmental Quality (TCEQ) (John Jolly, personal communication). These are more detailed versions of the standard inventories that are reported by TCEQ to the Environmental Protection Agency (EPA) for inclusion in the National Emissions Inventory (NEI) data base. In particular, these inventories contain speciated VOC emissions data. In 2006, the Houston-Galveston-Brazoria area contained 506 accounts, which represent individual facilities and sites. In order to obtain the most complete and accurate bottom-up emission inventory, the 2006 PSEI inventory used in this work has been updated to include more detailed information when possible. The updates are described below.

[13] 1. During 15 August to 15 September 2006, 86 accounts were asked to provide hourly emissions data at the level of a “path,” which is a uniquely defined combination of account, process unit, and emission point. Hourly path emissions were reported for 6% of the total paths for these 86 facilities, representing about 1% of the total paths in the Houston-Galveston-Brazoria area. This information has been incorporated into the PSEI.

[14] 2. Continuous Emissions Monitoring System (CEMS) instruments are installed on 17 of the largest sources in the Houston-Galveston-Brazoria area to monitor NOx, SO2, and CO2 emissions. We have updated the PSEI to include the average CEMS measurements reported by these sources during September to October 2006.

[15] The TCEQ PSEI includes NOx, SO2, CO, total VOC, PM2.5, and PM10 for sources in Texas. Emissions of hundreds of individual VOC compounds were also reported in the 1999 and 2006 TCEQ PSEI. When examining specific time periods in our analysis, we have considered hourly averaged emissions data from CEMS and estimates provided by facility operators. The timing and nature of nonroutine emission events at many facilities were also reported to TCEQ and are considered in this study.

3. Results and Discussion

3.1. Emission Inventory and Predicted Trends

[16] Figure 1 shows NOx and total VOC emissions from the TCEQ 2006 PSEI for the Houston-Galveston-Brazoria area, as described in section 2.3. The map shows a 160 km × 160 km area and indicates the major petrochemical industrial complexes, including the Houston Ship Channel, Mont Belvieu, Texas City, Chocolate Bayou, Freeport, and Sweeny. In the southeastern United States, ozone production downwind of power plants is typically most efficient in the presence of reactive biogenic hydrocarbons, such as isoprene [Ryerson et al., 2001]. The colocation of major NOx and VOC sources at the petrochemical industrial complexes shown in Figure 1 has been shown to lead to rapid downwind ozone formation [Kleinman et al., 2002; Daum et al., 2003; Ryerson et al., 2003]. In contrast, coal- and gas-fired power plants, such as W. A. Parish, emit relatively minor amounts of VOCs, as shown for the emissions inventory in Figure 1 and confirmed by downwind measurements during the TexAQS 2006 field campaign. The inventory values and trends for the six industrial complexes are given in Table 2. The TCEQ PSEI indicates reductions in NOx and SO2 emissions between 1999 and 2006.

[17] Reported CEMS monitoring has been found to be consistent with both NOx and SO2 measurements by the
NOAA WP-3D aircraft [Peischl et al., 2010]. Previous work has shown that NO\textsubscript{x} controls on large power plants, such as W. A. Parish, have resulted in significant, measurable decreases in NO\textsubscript{x} emissions. These reductions have been observed by the NOAA WP-3D [Peischl et al., 2010] and satellite measurements [Kim et al., 2006]. Some industrial sources in Houston have implemented NO\textsubscript{x} and SO\textsubscript{2} controls, similar to the large power plants. These reductions may have led to the decreasing values for NO\textsubscript{x} and SO\textsubscript{2} emissions in the TCEQ PSEI between 1999 and 2006, shown in Table 2.

3.2. Trends in NO\textsubscript{x} and SO\textsubscript{2} Emissions for Industrial Sources

Two different techniques are typically used to compare aircraft measurements to emission inventories. The first technique is the calculation of absolute fluxes [White et al., 1976; Trainer et al., 1995], which requires accurate knowledge of wind speed, wind direction, and aircraft speed. In addition, the boundary layer must be well defined and well mixed, and the emissions must have had sufficient time to mix vertically throughout the boundary layer. These are potential sources of error during meteorological conditions that do not have vigorous mixing or a well-defined boundary layer height or for aircraft transects that are close enough to the source that plumes have not had sufficient time to mix vertically. Poorly defined boundary layer heights and weak vertical mixing were observed during a number of the WP-3D flights in 2006. For this analysis, we have chosen to calculate emission ratios instead. This second method does not require knowledge of the boundary layer height or mixing and is not subject to the same assumptions and errors described above. Because emission ratios are determined from the enhancement of two species in a discrete plume, they are not sensitive to seasonal or meteorological changes that may affect the background concentration. Attribution of observed enhancement ratios to a single known source becomes complicated when multiple plumes coalesce. In this analysis, we have selected discrete plumes that exhibit enhanced mixing ratios for anthropogenic species and can be unambiguously matched to a source directly upwind.

Figure 2a shows a time series of CO\textsubscript{2}, NO\textsubscript{x}, and SO\textsubscript{2} mixing ratios for one transect measured downwind of the Sweeny industrial complex. This particular transect was measured on 21 September 2006 10:37 local standard time at an altitude of 340 m. The plume was intersected 11.4 km from the source, after a transit time of approximately 0.3 h. Plume transit time is determined from the average wind speed at the time of the plume intersection and the distance from the source. Figures 2b and 2c show the NO\textsubscript{x}/CO\textsubscript{2} and SO\textsubscript{2}/CO\textsubscript{2} correlation plots for these data. The large measured ratio of NO\textsubscript{x}/NO\textsubscript{y} (97% ± 2%) shows that oxidation of NO\textsubscript{y} was minimal, as expected from the short time between emission and measurement. Thus, losses of NO\textsubscript{y} were minimal and NO\textsubscript{x} measured downwind is a good proxy for...
the direct emissions of NO\textsubscript{x}. The fit in each case is determined from the linear orthogonal distance regression, weighted by the measurement imprecision (as given in Table 1). The industrial complexes contain multiple discrete units whose emissions typically cannot be separated in the WP-3D data. The slopes of the correlation plots for downwind transects are equal to the net emission ratios for the entire complex.

[20] Emission ratios of NO\textsubscript{x}/CO\textsubscript{2} and SO\textsubscript{2}/CO\textsubscript{2} were calculated for all WP-3D and Electra transects of Texas City, Sweeny, and Chocolate Bayou industrial plumes during 2000, 2002, and 2006 following the example shown in Figures 2b and 2c. Plumes included in this analysis were intersected 0.3–2.5 h after original emission and had \( R^2 \) correlation coefficients greater than or equal to 0.75. Figure 2d shows the geometric mean and range of NO\textsubscript{x}/CO\textsubscript{2} emission ratios observed at each industrial complex for each field campaign. Despite the day-to-day variability, the emission ratios show a decreasing trend in NO\textsubscript{x} emissions. A linear fit to the ratios, weighted by an imprecision of 30\% for each data point, indicates a change and 1-sigma uncer-

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\[ ^{a} \text{Values are reported in short tons d}^{-1}, \text{with NO}\textsubscript{x} \text{reported as NO}_2. \]

Figure 2. (a) Time series of NO\textsubscript{x}, SO\textsubscript{2}, and CO\textsubscript{2} measured downwind of the Sweeny industrial complex on 21 September 2006. (b) Correlation plot showing the relationship between NO\textsubscript{x} and CO\textsubscript{2} for the Sweeny plume transect. A linear fit to the data, weighted by the imprecision, gives an \( R^2 \) correlation coefficient of 0.91. (c) Similar correlation plot showing the relationship between SO\textsubscript{2} and CO\textsubscript{2}. A linear fit to the data, weighted by the imprecision, gives an \( R^2 \) correlation coefficient of 0.81. (d) Emission ratios were calculated for all industrial plumes intersected by the NOAA WP-3D aircraft during 2000, 2002, and 2006, following the method shown in Figures 2a, 2b, and 2c. Plumes included in this analysis were intersected 0.3–2.5 h after original emission and had an \( R^2 \) correlation coefficient greater than or equal to 0.75. The panel shows the geometric mean and annual range of emission ratios observed at each industrial complex. A linear fit to the ratios, weighted by an imprecision of 30\% for each data point, indicates a change and 1-sigma uncer-
tainty in \( \text{NO}_x/\text{CO}_2 \) of \(-29\% \pm 20\% \) between August 2000 and September 2006. \CO_2 emissions scale quantitatively with the mass of fuel consumed, so this result shows that the emissions of \text{NO}_x per unit of \text{CO}_2 has decreased. Figure 2e indicates no significant trend in \text{SO}_2/\text{CO}_2 emission ratios.

[21] The error bars in Figures 2d and 2e show the range of values observed during each year and indicate the variability between repeated measurements. Since each industrial complex contains numerous individual processes, which may run discontinuously, it is likely that this variability is real. Recent analysis of aircraft measurements found that \NO_x and \text{SO}_2 to \text{CO}_2 emission ratios for large power plants agree with available CEMS measurements of stack exhaust within \( \pm 14\% \) \[Peischl et al., 2010\]. This measurement uncertainty represents the uncertainty of emission ratios calculated for each individual plume transect.

3.3. Major Contributors to Ozone Formation

[22] Effective air quality regulations require quantification of the relative contribution of \NO_x and different VOC species to ozone formation. The importance of an individual hydrocarbon species to ozone formation depends both on its concentration, reactivity, and the reactivity of subsequent reaction products. The relative impacts of different hydrocarbons on urban ozone production can be determined to first order from their OH reactivity, because the initial peroxy radical formation is the rate-limiting step in ozone formation. This comparison is approximate, because it considers only the OH reactivity of the initial VOC species and not the subsequent chemistry of its products. OH reactivity is calculated as the product of the hydrocarbon concentration and its OH reaction rate coefficient at the measured ambient pressure and temperature, according to the expression,

\[
R_{\text{OH-VOC}} = k_{\text{OH-VOC}}[\text{VOC}].
\]

[23] The actual amount of ozone produced during the oxidation of a particular hydrocarbon species depends on the subsequent reaction products, meteorology, and the concentrations of other hydrocarbons and \text{NO}_x \[Carter, 1994\]. However, the relative OH reactivity of different hydrocarbons provides a simple method for comparing their impact on ozone production. In particular, this method allows the instantaneous reactivity of hydrocarbons measured in a complex mixture to be directly compared.

[24] Mixing ratios of an extensive suite of VOC compounds were acquired during summer 2000 at a ground-based site southeast of the Houston Ship Channel \[Jobson et al., 2004\]. These included \text{CO}, \text{C}_2–\text{C}_{10} alkanes, \text{C}_2–\text{C}_5 alkenes, ethyne, \text{C}_6–\text{C}_9 aromatics, and \text{C}_2–\text{C}_5 aldehydes and ketones. The authors found that reactivity was dominated by \text{C}_1–\text{C}_3 aldehydes, with ethene and propene occasionally dominating in highly concentrated plumes. During 2006, a similar suite of VOC measurements acquired onboard the NOAA R/V Ronald H. Brown in the Gulf of Mexico, Galveston Bay, and the Houston Ship Channel \[Gilman et al., 2009\] showed that \text{CO} and \text{CH}_4 dominated OH reactivity in the Gulf of Mexico, while VOCs from anthropogenic sources played the dominant role in Galveston Bay and the Houston Ship Channel. Ethene, propene, and formaldehyde were the largest contributors to OH reactivity, despite the higher mixing ratios of \text{C}_2–\text{C}_4 alkanes. By examining diurnal profiles, the authors determined that formaldehyde was produced predominantly by secondary chemistry rather than from primary emissions. This contradicts an analysis that correlates formaldehyde with other species and concludes that the formaldehyde budget is dominated by primary sources \[Rappengluck et al., 2010\].

[25] Aircraft measurements of VOCs provide important information that extends what is available from ground- and ship-based data sets by sampling VOC emissions directly downwind of selected sources. During 2000, 2002, and 2006, whole air samples acquired in the isolated industrial plumes showed elevated mixing ratios for many hydrocarbons, including alkanes, alkenes, aromatics, and ethyne. The most abundant VOCs observed in whole air samples acquired in industrial plumes were ethane (\text{C}_2\text{H}_6), ethene (\text{C}_2\text{H}_4), propene (\text{C}_3\text{H}_6), and isomers of butane (\text{C}_4\text{H}_{10}) and pentane (\text{C}_5\text{H}_{12}). In some cases, 1,3-butadiene (\text{C}_4\text{H}_6) and isomers of butene (\text{C}_4\text{H}_8) andpentene (\text{C}_5\text{H}_{10}) were also abundant. OH reactivities for the measured VOCs and \text{NO}_2 are presented for the industrial sources in Houston in the following subsections. The OH reaction rate coefficients used here to calculate OH reactivity are taken from several references \[Atkinson, 1986, 1994, 1997; DeMore et al., 1997; Atkinson et al., 2005; Sander et al., 2006\].

3.3.1. Houston Ship Channel

[26] The Houston Ship Channel is the largest complex of industrial sources of \text{NO}_x and reactive VOCs in the Houston-Galveston-Brazoria area, so the relative contribution of different VOCs to ozone formation is particularly important for this conglomeration of sources. Figure 3 presents measurements from a representative transect downwind of the Houston Ship Channel on 26 September 2006. Emission sources in the Houston Ship Channel are shown in Figure 3a. Figure 3b shows mixing ratios of \text{NO}_x, \text{SO}_2, and \text{CO}_2 observed during the transect. It is not always possible to attribute specific plumes to individual sources, although some geographical groupings can be made. For example, the large \text{SO}_2 source at the west end of the Houston Ship Channel is attributed to industrial sulfuric acid manufacturing.

[27] Figure 3c shows OH reactivity in the Houston Ship Channel plume for the 1 Hz \text{NO}_x, \text{CH}_4, and semicontinuous \text{CH}_3\text{CHO} data. The discrete bars represent the seven second sampling period of the whole air sampler, with formaldehyde and PTR-MS measurements (including acetaldehyde and isoprene) interpolated onto this time base. \text{CO} and \text{CH}_4 are excluded. Figure 3c shows that anthropogenic VOCs in the Houston Ship Channel are a major contributor to ozone formation. During this transect, total OH reactivity when formaldehyde measurements were available varied between 6.5 and 9.3 \text{ s}^{-1}, consistent with daytime observations made onboard the R/V Brown in the Houston Ship Channel \[Gilman et al., 2009\]. Although anthropogenic alkane emissions contribute to the OH reactivity, the total is dominated by ethene, propene, formaldehyde, and acetaldehyde. Formaldehyde and acetaldehyde are each formed primarily as secondary products from the oxidation of ethene and propene \[Wert et al., 2003\]. In contrast, the OH reactivity attributed to biogenic isoprene and other biogenic VOCs is comparatively small in the Houston Ship Channel.
This is consistent with other measurements in the ship channel that showed the importance of anthropogenic alkene emissions [Kleinman et al., 2002; Ryerson et al., 2003; Wert et al., 2003; Jobson et al., 2004; Gilman et al., 2009].

### 3.3.2. Other Industrial Sources

High VOC emissions are typical not only for the Houston Ship Channel but also for other industrial facilities in the Houston area. Ozone precursors at the other industrial complexes with fewer individual sources than the Houston Ship Channel can similarly be examined to give insight into the reactivity and relative importance of VOC species. Figure 4 shows representative transects downwind of Texas City, Chocolate Bayou, Freeport, and Sweeny acquired on 19 and 29 September 2006. Each of the industrial complexes shown in Figure 4 has comparable or greater OH reactivity with VOCs than with NO₂. Similar to the Houston Ship Channel, TCEQ 2006 PSEI values for NOₓ (solid red circles) and VOC (open black circles) are shown. The flight track of the NOAA WP-3D is shown as a black line. Wind direction is from the northeast.

**Figure 3.** Measurements conducted by the NOAA WP-3D aircraft downwind of the Houston Ship Channel on 26 September 2006. (a) Map of the Houston Ship Channel. TCEQ 2006 PSEI values for NOₓ (solid red circles) and VOC (open black circles) are shown. The flight track of the NOAA WP-3D is shown as a black line. Wind direction is from the northeast. (b) Time series of NOₓ, SO₂, and CO₂ measured along the flight track. (c) Solid bars showing OH reactivity for isoprene (green), alkanes (pink), ethene (orange), propene (blue), acetaldehyde (purple), formaldehyde (brown), and other measured VOC species (gray). CO and CH₄ are excluded from the reactivity calculation. OH reactivity with NO₂ (red), formaldehyde (brown), and acetaldehyde (purple) are shown as semicontinuous traces.
Channel, ethene, propene, formaldehyde, and acetaldehyde are major contributors to the OH reactivity. For transects downwind of Texas City, Sweeny, and Freeport, these VOCs account for more than half of the OH reactivity. At Chocolate Bayou, the reactivity includes an additional contribution from 1-butene and 1,3-butadiene. These highly reactive compounds may be emitted by other facilities as well but are rapidly removed in the atmosphere by oxidation. Both Sweeny and Chocolate Bayou indicate a larger OH reactivity due to isoprene than is observed in the Houston Ship Channel. This is expected from biogenic emission inventories for the region [Wiedinmyer et al., 2001; Warneke et al., 2010]. At Freeport, the fraction of OH reactivity attributed to VOCs is smaller than at the other complexes.

Elevated ethene, propene, formaldehyde, and acetaldehyde concentrations are consistently observed downwind of industrial sources in the aircraft measurements. Previous analysis of ethene measurements concluded that emissions from mobile sources were small downwind of the Houston urban core [de Gouw et al., 2009]. Similarly, we observe here that ethene, propene, formaldehyde, and acetaldehyde are predominantly associated with industrial point sources and not mobile sources.

3.4. Temporal Trends in Ethene and Propene Emissions for Industrial Sources

According to the calculations of OH reactivity presented in Figures 3 and 4, ethene and propene are two of the most significant VOC precursors to ozone formation from industrial sources in Houston. The aircraft data acquired in 2000, 2002, and 2006 can be used to determine emission trends and variability for ethene and propene. Previous studies in 2000 showed substantial discrepancies between emission inventories and measurements for these highly reactive VOCs [Ryerson et al., 2003; Wert et al., 2003]. We can compare the ethene and propene emissions measured in 2006 to the TCEQ PSEI to quantify the agreement and examine changes.

3.4.1. C2H4/NOx and C3H6/NOx Ratios

Emission ratios are the best choice for comparing the aircraft measurements of ethene and propene emissions to the TCEQ Point Source Emissions Inventory and examining temporal trends in their emissions. For the sparse samples acquired by the whole air sampler, it is not possible to calculate absolute fluxes. Continuous ethene measurements were acquired by the LPAS instrument, and these data were used to determine emission fluxes for ethene from the Mont Belvieu chemical complex [de Gouw et al., 2009]. However, ethene concentrations in some plume transects analyzed here were at or near the detection limit of the LPAS. When conserved species are coemitted from a single large source, their enhancement ratio should remain constant as the plume is transported downwind and diluted. However, their ratio will change if chemical or physical processes preferentially remove one of the species. The location of NOx and VOC emissions within a facility do not have to coincide, as long as the plume is well mixed at the time of downwind sampling. For the Texas City, Chocolate Bayou, Freeport, and Sweeny industrial complexes, we use aircraft measurements acquired during plume transects in 2000, 2002, and 2006 to calculate emission ratios of C2H4/NOx.
Measurements of C$_2$H$_4$/NO$_x$ measured for plume transects during mean, median, range, and emission inventory values for the 2006 data at each of the four industrial complexes. The measured C$_2$H$_4$/NO$_x$ ratios exceed the emission inventory values by factors of 1.4–20. Independent flux measurements acquired in 2006 by solar occultation and DOAS [Mellqvist et al., 2010; Rivera et al., 2010] are shown in Figure 5, and these agree well with the aircraft determinations. The consistent disagreement between the measurements and emissions inventory over multiple sampling dates and multiple years makes it unlikely that the high ethene emissions are due to upsets or nonroutine emission events. The discrepancies between the emissions inventory and the measurements are consistently large.

[34] Identifying temporal trends in the C$_2$H$_4$/NO$_x$ ratio from Figure 5 is difficult. For the NO$_x$/CO$_2$ and SO$_2$/CO$_2$ ratios presented in Figure 2, the uncertainty in the individual ratio determinations is comparable to the variability between repeated measurements. For the C$_2$H$_4$/NO$_x$ and C$_2$H$_6$/NO$_x$ ratios, the variability is higher than the measurement uncertainty. The day-to-day and within-plume variability is large (approximately −50% to +100%), as indicated by the range bars. This is consistent with previous studies that reported variability in ethene emissions [Murphy and Allen, 2005; de Gouw et al., 2009; Mellqvist et al., 2010]. It appears that decreases in ethene emissions relative to NO$_x$ may have occurred at the Sweeny industrial complex. For the other industrial complexes, any interannual trend is hidden by variability within the plume and between different days.

[35] Figure 6 shows the observed ratio of C$_2$H$_6$/NO$_x$ for the same plume transects as displayed in Figure 5. As for ethene, the solid black circles indicate the average emission ratio and range. As described above, the measured C$_2$H$_4$/NO$_x$ ratio represents a lower limit of the actual emission ratio due to the higher reactivity of C$_2$H$_4$ relative to NO$_x$. Like the C$_2$H$_4$/NO$_x$ ratios in Figure 5, the measured C$_2$H$_6$/NO$_x$ ratios in Figure 6 exceed the emission inventory values by factors of 1–24. This indicates that the actual discrepancy between measurements and the TCEQ Point Source Emissions...

![Figure 5](image-url)

**Table 3.** VOC/NO$_x$ Ratios Measured for Plume Transects During 2006 Compared to TCEQ Point Source Emissions Inventory Values$^a$

<table>
<thead>
<tr>
<th></th>
<th>C$_2$H$_4$/NO$_x$</th>
<th>C$_2$H$_6$/NO$_x$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n Mean Median Range E.I.</td>
<td>n Mean Median Range E.I.</td>
</tr>
<tr>
<td>Texas City</td>
<td>17 0.32 0.39 0.01–1.5 0.03</td>
<td>18 0.11 0.16 0.002–0.75 0.02</td>
</tr>
<tr>
<td>Chocolate Bayou</td>
<td>3 0.40 0.38 0.38–0.42 0.28</td>
<td>3 0.12 0.13 0.08–0.15 0.13</td>
</tr>
<tr>
<td>Freeport</td>
<td>4 0.36 0.61 0.04–1.1 0.08</td>
<td>4 0.05 0.07 0.02–0.13 0.01</td>
</tr>
<tr>
<td>Sweeny</td>
<td>7 0.93 1.1 0.49–1.2 0.06</td>
<td>7 0.68 0.65 0.48–1.1 0.03</td>
</tr>
</tbody>
</table>

$^a$Geometric mean is given.

and C$_2$H$_4$/NO$_x$. The daytime plumes examined in this work were intersected 0.2–2.7 h (2–61 km) downwind of the industrial complexes, as calculated from the average wind speed at the time of plume intersection and the distance from the source. Reaction with OH is the primary loss mechanism for C$_2$H$_4$, C$_2$H$_6$, and NO$_x$. The respective OH reaction rates of these molecules at standard temperature and pressure (298 K; 1013 mb) are $9.1 \times 10^{-12}$, $30.2 \times 10^{-12}$, and $10.6 \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ [Atkinson et al., 2005; Sander et al., 2006]. This indicates that the C$_2$H$_4$/NO$_x$ enhancement ratio will be approximately conserved downwind. In contrast, the measured C$_2$H$_6$/NO$_x$ ratio can only be considered a lower limit of the original C$_2$H$_4$/NO$_x$ emission ratio because of the greater OH reactivity of C$_2$H$_6$.

[33] Figure 5 shows the observed ratio of C$_2$H$_4$/NO$_x$ for plume transects acquired during 2000, 2002, and 2006 downwind of the four industrial complexes. The solid black circles indicate the geometric mean emission ratio and range. The four panels are plotted on a log scale to better compare with the C$_2$H$_4$/NO$_x$ ratio calculated from the TCEQ Point Source Emissions Inventory. Table 3 gives the number of samples, mean, median, range, and emission inventory values for the 2006 data at each of the four industrial complexes. The measured C$_2$H$_4$/NO$_x$ ratios exceed the emission inventory values by factors of 1.4–20. Independent flux measurements acquired in 2006 by solar occultation and DOAS [Mellqvist et al., 2010; Rivera et al., 2010] are shown in Figure 5, and these agree well with the aircraft determinations. The consistent disagreement between the measurements and emissions inventory over multiple sampling dates and multiple years makes it unlikely that the high ethene emissions are due to upsets or nonroutine emission events. The discrepancies between the emissions inventory and the measurements are consistently large.

[34] Identifying temporal trends in the C$_2$H$_4$/NO$_x$ ratio from Figure 5 is difficult. For the NO$_x$/CO$_2$ and SO$_2$/CO$_2$ ratios presented in Figure 2, the uncertainty in the individual ratio determinations is comparable to the variability between repeated measurements. For the C$_2$H$_4$/NO$_x$ and C$_2$H$_6$/NO$_x$ ratios, the variability is higher than the measurement uncertainty. The day-to-day and within-plume variability is large (approximately −50% to +100%), as indicated by the range bars. This is consistent with previous studies that reported variability in ethene emissions [Murphy and Allen, 2005; de Gouw et al., 2009; Mellqvist et al., 2010]. It appears that decreases in ethene emissions relative to NO$_x$ may have occurred at the Sweeny industrial complex. For the other industrial complexes, any interannual trend is hidden by variability within the plume and between different days.

[35] Figure 6 shows the observed ratio of C$_2$H$_6$/NO$_x$ for the same plume transects as displayed in Figure 5. As for ethene, the solid black circles indicate the average emission ratio and range. As described above, the measured C$_2$H$_4$/NO$_x$ ratio represents a lower limit of the actual emission ratio due to the higher reactivity of C$_2$H$_4$ relative to NO$_x$. Like the C$_2$H$_4$/NO$_x$ ratios in Figure 5, the measured C$_2$H$_6$/NO$_x$ ratios in Figure 6 exceed the emission inventory values by factors of 1–24. This indicates that the actual discrepancy between measurements and the TCEQ Point Source Emis...
Measurements of $C_2H_6$/$NO_x$ ratio by the NOAA is even greater than it is for $C_2H_4$. Independent flux measurements acquired in 2006 by solar occultation flux and DOAS [Mellqvist et al., 2010; Rivera et al., 2010] are shown as open blue circles with their calculated error bars. Again, the discrepancy between the emissions inventory and the measurements appears to be real. The day-to-day and within-plume variabilities are large, as indicated by the range bars, obscuring temporal trends in the $C_2H_6$/$NO_x$ ratio. It appears that $C_2H_6$ emission reductions relative to $NO_x$ may have occurred at the Chocolate Bayou industrial complex. For the other industrial complexes, any interannual trend is hidden by variability within the plume and between different days.

The large discrepancies shown in Figures 5 and 6 between $C_2H_4$/$NO_x$ and $C_2H_6$/$NO_x$ ratios reported in the TCEQ PSEI and measured by the WP-3D could be due to inaccuracy in the alkene emissions, $NO_x$ emissions, or both. Previous work has shown that $NO_x$ is better represented than VOCs in available point source emission inventories [Ryerson et al., 2003]. $NO_x$ emissions in the TCEQ PSEI are derived from CEMS data for many of the large $NO_x$ sources at each industrial complex. As discussed in section 3.2, the CEMS emission ratios have been verified within ±14% [Peischl et al., 2010] and the EPA requires the absolute accuracy to be within ±10%. Additionally, previous calculations of $NO_x$ mass flux for industrial sources during the TexAQS 2000 campaign showed agreement with emission inventories within ±50% [Ryerson et al., 2003]. There is no evidence that $NO_x$ inventory values are inaccurate by more than this factor in Houston. Thus, the large inventory discrepancies in $C_2H_4$/$NO_x$ and $C_2H_6$/$NO_x$ ratios shown in Figures 5 and 6 are unlikely due to errors in inventory $NO_x$ values. These discrepancies are more likely attributed to $C_2H_4$ and $C_2H_6$ emissions that are consistently much higher than represented in the TCEQ PSEI. $C_2H_4$ and $C_2H_6$ emissions from industrial sources that are much higher than values in the inventory have been observed during every flight in Houston during 2000, 2002, and 2006. These higher emissions were not reported as flares or upset events, but instead represent typical operating conditions in the Houston urban area.

Examination of the temporal trends for $C_2H_4$/$NO_x$ and $C_2H_6$/$NO_x$ ratios further requires consideration of how both VOC and $NO_x$ concentrations change through time. Trends in $NO_x$ emissions, as observed in $NO_x$/$CO_2$ ratios for the industrial complexes, were discussed in detail in section 3.2. The TCEQ PSEI indicates $NO_x$ reductions of 40%–84% (see Table 2), and the WP-3D measurements show reductions of 29% ± 20% (see Figure 2d). We estimate the total change in emissions of $C_2H_4$ and $C_2H_6$ for the industrial complexes over the 2000–2006 period to be −30% ± 30%.

### 3.4.2. Ambient $C_2H_4$ and $C_2H_6$ Concentrations

The previous section presented emission ratios that can be linked to known industrial sources. It is also possible to examine trends in ambient $C_2H_4$ and $C_2H_6$ concentrations observed during 2000 and 2006 onboard the NOAA aircraft. Figure 7a shows a geographical box that includes the Houston Ship Channel, Mont Belvieu industrial complex, and the Houston urban core. During 2000 and 2006, 117 and 273 whole air samples, respectively, were acquired within this sampling area at an aircraft altitude of 200–1000 m. Figures 7b and 7c show the cumulative probability distributions for ethene and propene mixing ratios measured during each of the two years. The median observed value changed by −52% and −48% for ethene and propene, respectively. Although the mean concentrations may be biased high values, the mean has a well-defined uncertainty that is useful for evaluating whether the trend is significant. The mean concentration changes for ethene and propene were −52% ± 8% and −52% ± 21%, respectively. Surface measurements acquired during 2000 and 2006 have also indicated a decrease in ambient concentrations of $C_2H_4$ and $C_2H_6$ [Gilman et al., 2009]. Reductions in measured VOC concentrations may be due either to reductions in emissions or to differences in meteorology that result in different dilution or oxidation of the emissions. Fully quantifying the potential differences in boundary layer heights, solar radiation, and mixing during the two years is difficult. However, all emissions are subject to the same meteorology.

Ethene, shown in Figure 7d, is emitted in vehicle exhaust and is typically used as an urban tracer. The cumulative probability plot for ethene shows a more modest change in the median value between 2000 and 2006 of −10%. The trend in the mean concentration of ethene was −37% ± 10%.
Analysis of CO, another urban tracer, shows a change in the median value of $-18.0\%$ during this same time period, with a trend in the mean concentration of $-17\% \pm 3\%$. Figures 7e and 7f show the calculated cumulative probability for ambient temperature and wind speed observed during the 2 years. Average wind speeds were higher in 2006 than in 2000, which would lead to a greater dilution of ethene emissions and lower mixing ratios in 2006. Conversely, average temperatures were lower in 2006 than in 2000, consistent with observations of shallower boundary layers and less vigorous vertical mixing, which would lead to higher mixing ratios in 2006. A similar cumulative analysis of formaldehyde, the main photooxidation product of ethene, for 2000 and 2006 in the same geographical area shows that its median concentration decreased by more than 40%. The changes in the ambient concentrations of all of these species are consistent with the conclusion that the decreases in $C_2H_4$ concentration are due to decreases in emissions, rather than to differences in meteorological conditions between the two data sets.

### 3.5. Secondary Photochemical Products Formed Downwind

The large suite of measurements onboard the NOAA WP-3D aircraft makes it possible to examine chemical processing downwind of the industrial complexes. Oxidation of each of the alkene species results in different photochemical products. Initially, alkenes react with OH, O$_3$, or NO$_x$ to form ketones and aldehydes. The carbon-carbon double bond is broken, forming two carbonyl groups. The aldehydes proceed to react with OH or photolyze. In particular, terminal alkenes react with OH to form formaldehyde, which may subsequently react with OH to yield a hydroperoxy radical and CO, or photolyze through one of two pathways. 2-Alkenes react with OH to form acetaldehyde, which may then react with OH in the presence of NO$_2$ to form peroxyacetyl nitrate (PAN). Using the NOAA WP-3D measurements, we can identify the ketone, aldehyde, and PAN reaction products downwind.

A series of three WP-3D transects acquired downwind of the Texas City industrial complex on 19 September 2006 is shown in Figure 8a. The portion of each flight track crossing the plume is shown in black, with measured NO$_x$ plotted in red to indicate the plume location. The plume was intersected at distances of 19.6, 41.3, and 56.0 km downwind, corresponding to plume ages of 0.6, 1.5, and 2.3 h. As discussed previously and shown in Figure 4a, the Texas City industrial complex emits high concentrations of reactive VOCs and NO$_x$. Figures 8b and 8c show the mixing ratios of primary emissions (ethene, propene, and NO$_x$) and secondary photochemical products (formaldehyde, acetaldehyde, PAN, and O$_3$) during the first and third transects. The initially high ethene concentrations observed within the first transect are greatly decreased by the third transect, as seen in the continuous LPAS ethene data and single WAS sample, demonstrating the high reactivity of ethene. Similarly, while NO$_x$ accounts for more than half of the NO$_y$ observed in the first transect, it has been largely oxidized by the third transect.

The loss rates of the primary emissions and production rates of the secondary photochemical products can be calculated by assuming that NO$_x$ is a conserved plume tracer over the transport time of these three transects. Figure 8d shows ln(VOC/NO$_y$) ratios for ethene and propene as a function of downwind transport time with linear fits weighted by the measurement imprecision. The corresponding expo-
Figure 8. (a) Map of three transects measured downwind of the Texas City complex on 19 September 2006. Measured NO\textsubscript{x} is plotted on the map, indicating the plume location. (b and c) Primary emissions and secondary photochemical products measured during the first and third downwind transects. The x axis indicates the cross-plume distance, with 0 km at the center of the plume. Primary emissions include ethene (orange bar) and propene (blue bar) measured by the whole air sampler, ethene measured by LPAS (solid orange circles) and NO\textsubscript{x} (red line). Secondary photochemical products include formaldehyde (brown line), acetaldehyde (solid purple circles), NO\textsubscript{y} (green line), PAN (solid gray circles), and O\textsubscript{3} (black line). (d) Natural log of the ratio of VOC species to NO\textsubscript{y}, as a function of transport time downwind. NO\textsubscript{y} is used here as an approximately conserved plume tracer. Slopes for ethene measured by the LPAS and whole air sampler are $-0.4 \pm 0.2$ and $-0.6 \pm 0.2$ h\textsuperscript{-1}, respectively. The slope for propene is $-1.1 \pm 0.2$. (e) Ratios of VOC photochemical products to NO\textsubscript{y}. The slopes for formaldehyde, acetaldehyde, and PAN are $0.27 \pm 0.02$, $0.35 \pm 0.24$, and $0.112 \pm 0.004$ ppbv ppbv\textsuperscript{-1} h\textsuperscript{-1}. The slope of $1.6 \times$ ethene + $1.0 \times$ propene is $-0.22 \pm 0.08$ ppbv ppbv\textsuperscript{-1} h\textsuperscript{-1}.
ential decay rate of ethene for the LPAS instrument and whole air sampler are $-0.4 \pm 0.2$ and $-0.6 \pm 0.2$ h$^{-1}$, respectively. The observed propene loss is $-1.1 \pm 0.2$ h$^{-1}$. Figure 8e shows the production rates of formaldehyde, acetaldehyde and PAN as $0.27 \pm 0.02$, $0.35 \pm 0.24$, and $0.112 \pm 0.004$ ppbv ppbv$^{-1}$ h$^{-1}$. The sum of $1.6 \times C_2H_4/NO_x + 1.0 \times C_2H_4/NO_x$ represents a formaldehyde yield of 1.6 for ethene [Niki et al., 1981] and 1.0 for propene [Niki et al., 1978]. This sum has a fitted slope of $-0.22 \pm 0.08$ ppbv ppbv$^{-1}$ h$^{-1}$, indicating that the loss of these two alkenes accounts for more than 80% of the observed formaldehyde production, neglecting photochemical losses for formaldehyde. The propene losses are insufficient to account for the production of acetaldehyde and PAN, indicating that other 2-alkenes may be important in the carbon budget for these two secondary products.

4. Conclusions

[45] Observed reductions of 29% $\pm$ 20% in NO$_2$ emissions at the industrial complexes between August 2000 and September 2006 are consistent with observations of reduced NO$_2$ emissions at larger power plants throughout the southeastern United States that have implemented emission abatement controls. Previous analysis of industrial emissions measured during the TexAQS 2000 field campaign showed that the principal VOC precursors leading to the rapid ozone formation characteristic of the Houston-Galveston-Brazoria area were petrochemical emissions of ethene and propene. Our examination of temporal trends in C$_2$H$_4$/NO$_x$ and C$_3$H$_6$/NO$_x$ ratios from isolated petrochemical sources between 2000 and 2006 leads to the conclusion that emissions of these alkenes have decreased by 30% $\pm$ 30%. Additionally, we observe that median ambient concentrations of ethene and propene within the Houston urban area have changed by $-52\%$ and $-48\%$, respectively, over this same time period. One principal result from the 2000 field study was the identification of large underestimates of reactive light alkenes in VOC emission inventories for petrochemical sources [Kleinman et al., 2002; Karl et al., 2003; Ryerson et al., 2003; Wert et al., 2003; Jobson et al., 2004]. The analysis presented here for 2000, 2002, and 2006 measurements in the Houston-Galveston-Brazoria area indicates that emission inventory inaccuracies persist. Measured ratios of C$_2$H$_4$/NO$_x$ and C$_3$H$_6$/NO$_x$ exceed emission inventory values by factors of 1.4–20 and 1–24, respectively. De Gouw et al. [2009] reported ethene emissions of 520 $\pm$ 140 kg h$^{-1}$ from the Mont Belvieu industrial complex. Comparing those measurements to the updated 2006 emission inventory shown in Table 2 gives a factor of 6 discrepancy, which is similar to the results reported here for other industrial complexes. Because of the inaccurate accounting of ethene, propene, and other reactive VOC emissions in the 2006 TCEQ PSEI, chemical transport models will continue to have difficulty in accurately predicting ozone levels in the greater Houston area. Although the current analysis has focused on the isolated petrochemical industrial areas, including Texas City, Chocolate Bayou, Sweeny, and Freeport, the largest source in this area remains the Houston Ship Channel. The high concentration of colocated NO$_x$ and VOC sources in the Houston Ship Channel make it difficult to analyze, but it is certainly a major contributor to ozone formation.

[45] Acknowledgments. We thank the management, staff, and flight crew from the NOAA Aircraft Operations Center for their support during the field mission. We thank Bryan Lambeth and TCEQ for the La Porte wind profiler data. We thank John Jolly for TCEQ emissions inventory data. We thank Johan Mellqvist, Claudia Rivera, and Jerker Samuelsson for sharing their solar occultation flux and DOAS measurements. We thank Dr. Sacco te Lintel Hekkert from Sensor Sense for LPAS measurements. R.A.W. thanks Dr. Chuck Brock for helpful discussions. We acknowledge financial support for the field measurements from the NOAA Air Quality and NOAA Climate Research and Modeling programs. We acknowledge financial support for the analysis from Texas Commission on Environmental Quality contract 582-8-86246-FY09. R.A.W. acknowledges a National Research Council Postdoctoral Fellowship.

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