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Biomass burning and urban air pollution over the Central Mexican Plateau

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Abstract. Observations during the 2006 dry season of highly elevated concentrations of cyanides in the atmosphere above Mexico City (MC) and the surrounding plains demonstrate that biomass burning (BB) significantly impacted air quality in the region. We find that during the period of our measurements, fires contribute more than half of the organic aerosol mass and submicron aerosol scattering, and one third of the enhancement in benzene, reactive nitrogen, and carbon monoxide in the outflow from the plateau. The combination of biomass burning and anthropogenic emissions will affect ozone chemistry in the MC outflow.

1 Introduction

The 20 million (2005) inhabitants of Mexico City experience some of the worst air quality in the world. The high population density coupled with the topography of the city (2200 m, surrounded on three sides by mountains) leads to the daily buildup of pollutants in the region (Molina and Molina, 2002). Despite growing population and greatly increased automobile use, air quality has improved measurably in the last decade as the federal and city governments implemented a series of air quality regulations broadly similar to those that have been effective in, for example, Los Angeles (Lloyd, 1992). Nevertheless, ozone and particulate matter (PM) in the city often exceed international standards (WHO, 2008) and the city is consistently enveloped in a pall by the large amount of aerosol present.

The pollution from the city has impacts beyond the basin. Aerosols and ozone produce important forcing on regional climate through their interaction with both thermal infrared and visible radiation (Solomon et al., 2007). Indeed, the effluents from megacities, such as Mexico City, are now seen as globally important sources of pollution.

In the last decade, there have been several intensive studies of the air quality in the Mexico City basin. A major study undertaken in the Mexico City Metropolitan Area in spring of 2003 (MCMA-2003) included significant international cooperation (Molina et al., 2007). In the spring of 2006, a consortium of atmospheric scientists expanded significantly on this study, obtaining a large suite of measurements in and around Mexico City in an effort to understand both the controlling chemistry in the basin and the impacts of the outflow pollution on the broader region. Named Megacity Initiative: Local and Global Research Observations (MILAGRO), this campaign involved measurements at several ground sites along the most common outflow trajectory, and from several
Fig. 1. MODIS-Aqua images of the Mexico City basin on (a) 5 March 2006 at 13:35 CST and (b) 10 March 2006 at 13:55 CST, illustrate how large fires in the hillsides surrounding the city can impact visibility. Red boxes are thermal anomalies detected by MODIS. Black lines represent Mexican state boundaries. Images courtesy of MODIS Rapid Response Project at NASA/GSFC.

Biomass burning can be a significant contributor to poor air quality in many regions of the world, including southern California (Muhle et al., 2007). Several previous studies have suggested that fires in and around the Mexico City basin can impact air quality (Molina et al., 2007; Bravo et al., 2002; Salcedo et al., 2006). During the springtime (March–May), many fires occur in the pine forests on the mountains surrounding the city, both inside and outside the basin. These fires are virtually all of human origin. Primarily they originate from accidental means (escaped agricultural/land maintenance fires, escaped campfires, smoking, fireworks, vehicles, etc.), with a smaller number originating from intentional ignition (E. Alvarado, Univ. of Washington, personal communication, 2009). Typically, the biomass burning season intensifies in late March, reaching a maximum in May (Fast et al., 2007; Bravo et al., 2002). The heat from these fires is observable from space by the infrared channels of the moderate resolution imaging spectroradiometer (MODIS) instruments operated from NASA’s Aqua and Terra platforms (Giglio et al., 2003). Figure 1, for example, shows two visible images from MODIS taken on 5 March (panel a) and 10 March (panel b) 2006. The locations of the detected thermal anomalies are shown as red boxes. The aerosol haze from the fires can be seen covering large areas of land around and above MC, particularly on 5 March. MODIS imagery suggests that the total biomass burning around MC in March 2006 was greater than climatological amounts, and closer to what is normally observed during the month of April.

Using tracers of pollution from biomass burning and urban emissions, we show that fires significantly impacted air quality above and downwind of Mexico City in March 2006. We use aircraft measurements of hydrogen cyanide (HCN) to estimate the contribution of biomass burning to the regional air quality. HCN is produced in the pyrolysis of amino acids (Ratcliff et al., 1974) and has been widely used as an atmospheric tracer of biomass burning emissions (e.g., Li et al., 2003). We use simultaneous observations of acetylene ($C_2H_2$) to characterize the contribution of urban
emissions. We show that a simple two end-member mixing model (biomass burning and urban emissions) developed from these tracers can explain most of the observed variability in several pollutants including carbon monoxide, benzene, organic aerosol, reactive nitrogen oxides (NO$_y$), and the amount of submicron aerosol particles.

2 Observations

The NSF C-130 flew through the Mexico City region on eleven flights in March 2006 (Fig. 2). Of these flights, three had fewer than ten samples of C$_2$H$_2$ within our study area (3×3 degree box centered on MC, shown in Fig. 2 and termed Central Mexican Plateau) and one flight did not have HCN observations. These flights are excluded from the calculation of the overall fire impact (4, 12, 26, and 28 March). In Fig. 2a, the aircraft flight tracks are colored by HCN–HCN$_{background}$, and in Fig. 2b observed C$_2$H$_2$. The map has been divided into 0.2×0.2 degree pixels and all observations from the C-130 across the MILAGRO campaign within a given pixel have been averaged together. High concentration data have been rounded down to capped values of 800 and 1500 ppbv for HCN and C$_2$H$_2$, respectively. Color scales range from 0 ppbv (dark blue) to the capped value (dark red). The black star in each panel is the center of Mexico City, and the black box outlines the 3×3 degree box centered on Mexico City which is the study area considered for the fire impact analysis.

![Figure 2](https://example.com/figure2.png)

**Figure 2.** C-130 flight tracks during MILAGRO colored by tracers: (a) observed HCN–HCN$_{background}$, and (b) observed C$_2$H$_2$. The map has been divided into 0.2×0.2 degree pixels and all observations from the C-130 across the MILAGRO campaign within a given pixel have been averaged together. High concentration data have been rounded down to capped values of 800 and 1500 ppbv for HCN and C$_2$H$_2$, respectively. Color scales range from 0 ppbv (dark blue) to the capped value (dark red). The black star in each panel is the center of Mexico City, and the black box outlines the 3×3 degree box centered on Mexico City which is the study area considered for the fire impact analysis.

![Table 1](https://example.com/table1.png)

**Table 1.** Anthropogenic and biomass burning emission ratios derived here (TLS) and those measured directly in the Mexico City area.

<table>
<thead>
<tr>
<th>Species</th>
<th>Urban (Δ[xy]/Δ[C$_2$H$_2$])$^a$</th>
<th>Biomass Burning (Δ[xy]/Δ[HCN])$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>opt$^b$ 29 March</td>
<td>fire obs.$^d$</td>
</tr>
<tr>
<td>CO</td>
<td>96±3 92</td>
<td>104±8 117</td>
</tr>
<tr>
<td>C$_6$H$_6$</td>
<td>0.137±0.005 0.13</td>
<td>0.19±0.01 7.1$^f$</td>
</tr>
<tr>
<td>NO$_y$</td>
<td>4.3±0.3 3.7</td>
<td>5.4±1.0 78</td>
</tr>
<tr>
<td>OA</td>
<td>3.9±1.0 2.9</td>
<td>22±4 14</td>
</tr>
<tr>
<td>scattering$^e$</td>
<td>22±3 22</td>
<td>75±5 78</td>
</tr>
<tr>
<td>HCN</td>
<td>0.056$^c$ – 1– 1–</td>
<td>1– 1– 0.17</td>
</tr>
<tr>
<td>C$_2$H$_2$</td>
<td>– – –</td>
<td>– – – 1– 1</td>
</tr>
</tbody>
</table>

$^a$ Units are mol/mol, except for $d[OA]/d[y]$ and $d[scat]/d[y]$, which have units of µg sm$^{-3}$ ppbv$^{-1}$ and Mm$^{-1}$ ppbv$^{-1}$, respectively.

$^b$ Calculated emission ratios determined from TLS analysis.

$^c$ An upper limit for how much HCN comes from urban emissions in MC, derived from data collected from C-130 on 29 March 2007, a day with low BB influence. This factor is not optimized.

$^d$ Median values for fires sampled by the Twin Otter around Mexico City in March 2006 (Yokelson et al., 2007b).

$^e$ This refers to submicron scattering measured at 550 nm.

$^f$ This value is the NO$_x$/HCN emission ratio, not NO$_y$/HCN. For this comparison we assume that in the fresh smoke sampled by Yokelson et al. (2007b), NO$_y$≈NO$_x$.

![Figure 2b](https://example.com/figure2b.png)

Figure 2b shows the C-130 flight tracks colored by the mixing ratio of C$_2$H$_2$. C$_2$H$_2$ is produced in the combustion of both gasoline and diesel fuels. We chose C$_2$H$_2$ as our urban tracer because its atmospheric lifetime is quite long. We calculate that with respect to its major loss mechanism (reaction with the hydroxyl radical, OH), the atmospheric lifetime is ten days to two weeks in the Mexico City region. We did not use other tracers of city emissions such as toluene or methyl tert-butyl ether (MTBE) because their shorter atmospheric lifetimes complicate the regional analysis. In fresh city plumes (as determined by the ratio of toluene to C$_2$H$_2$), all the urban tracers (e.g. C$_2$H$_2$, MTBE, toluene) are highly correlated.

Although the regions of enhanced C$_2$H$_2$ and HCN appear geographically coincident in Fig. 2, the sources of these
Fig. 3. Timeline from the flight of 8 March 2006: (a) measured HCN, C\(_2\)H\(_2\), and radar altitude. The observed (dots) and reconstructed (bars) CO, benzene, and organic aerosol concentrations, are shown in panels (b–d), respectively. The bar widths represent the relative bottle sampling time (width has been expanded by 3 \times for clarity). The measurements in panels (b–d) are colored by [toluene]/[C\(_2\)H\(_2\)]\(^*\) with color scale ranging from red=1 to blue=0, and grey meaning data was unavailable for this calculation. Data points with black borders in panels (b–d) lie within the 3 \times 3 degree study area.

Fig. 3a. On 8 March 2006, the C-130 flew into the Mexico City basin and over a short period encountered air masses significantly enhanced in either HCN, C\(_2\)H\(_2\), or both (panel a). Fig. 4 shows the flight track corresponding with data shown in Fig. 3 on top of the MODIS – Aqua image for 8 March.

To quantify the contribution of both fire and urban emissions to the distribution of a trace gas (or aerosol), \(Y\), we implement a simple two end-member model using the measured excess HCN, \([HCN]^*\), and measured excess C\(_2\)H\(_2\), \([C_2H_2]^*\), as tracers:

\[
Y = F_Y (\text{fire}) \times [HCN]^* + F_Y (\text{urban}) \times [C_2H_2]^* \tag{1}
\]

where,

\[ [HCN]^* = [HCN] - S_{HCN}(\text{urban}) \times [C_2H_2]^* - [HCN]_{\text{background}} \tag{2} \]

\[ [C_2H_2]^* = [C_2H_2] - S_{C_2H_2}(\text{fire}) \times [HCN]^* - [C_2H_2]_{\text{background}} \tag{3} \]

\(F_Y\) are scalars that relate the emission of \(Y\) from fire and urban sources to the emissions of HCN and C\(_2\)H\(_2\), respectively (Table 1). \(S_{HCN}\) and \(S_{C_2H_2}\) are the emission ratios of HCN to C\(_2\)H\(_2\) and C\(_2\)H\(_2\) to HCN for urban and fire emissions, respectively. These cross terms account for the contribution of urban and fire emissions to the excess HCN and C\(_2\)H\(_2\), respectively. We also account for the amounts of these tracers advected into the region from afar (backgrounds).

We derive a set of emission ratios, \(F_Y\), for the pollutants using total least squares (TLS) analysis (Table 1). We weight \([HCN]^*\) and \([C_2H_2]^*\) by estimates of their error determined primarily from uncertainties in the backgrounds and in the variability of the emission ratios. We estimate the uncertainty in the derived emission ratios using a bootstrap method (Efron and Tibshirani, 1993). The bootstrap method creates \(x\) alternate data sets by picking \(n\) random samples with replacement from the original data set, where \(n\) equals the
number of samples in the original data set. The TLS analysis is then performed across all alternate data sets, and statistics are computed on the results of all analyses. For this analysis we used $x=1000$. In Table 1, we also summarize the emission factors determined independently from measurements made directly in biomass burning plumes measured in the basin during MILAGRO.

Gasoline and diesel engine exhaust contain HCN, though previous measurements of the emissions vary by orders of magnitude (Baum et al., 2007). Automobiles lacking catalytic converters can produce 100 times more HCN than automobiles with functional catalysts (Baum et al., 2007; Harvey et al., 1983). To estimate the appropriate emission ratio for Mexico City, $S_{HCN}$, we use observations made from the C-130 on 29 March when the C-130 sampled city emissions with what appears to be minimal fire influence (Fast et al., 2007). The measured slope of HCN to C$_2$H$_2$ in the city plumes encountered on this day is 0.056 (mol/mol). The ratio of CH$_3$CN to HCN in the city emissions is 0.5, similar to the ratio measured in both fire plumes and in the region as a whole. This is in contrast to observations from Asia where urban emissions had a much lower ratio CH$_3$CN to HCN (Li et al., 2003). Thus it is possible that even on the 29th, some of the HCN is from burning. Given our cross term corrections, and assuming urban fire sources such as garbage, coal, and biofuel burning was no different on 29 March than on other days, emissions from these urban fire sources are counted as urban emissions and not as fire emissions. Using the 0.056 (mol/mol) as an upper limit for the HCN/C$_2$H$_2$ emission ratio from urban emissions, we estimate that all urban emissions of HCN account for no more than 15% of the total emissions in the basin during March 2006. Other sources of HCN from, for example, coal burning and petrochemical industries are also estimated to be small (see Appendix A2).

We use measurements of C$_2$H$_2$ and HCN observed in forest fires in and around Mexico City to estimate $S_{C_2H_2}$ (Yokelson et al., 2007b). The emission of C$_2$H$_2$ from these forest fires was near the low end of the range typically observed for extratropical forest fires (Yokelson et al., 2007b; Andreae and Merlet, 2001). We estimate that the contribution of biomass burning to C$_2$H$_2$ accounts for less than 10% of the C$_2$H$_2$ in and around Mexico City.

To account for the background amounts of C$_2$H$_2$ and HCN advected into the region, we use our observations in air sampled aloft, away from the Mexico City basin. We use separate HCN and C$_2$H$_2$ backgrounds for the gas phase species and for organic aerosol/scattering, as aerosols have a more variable atmospheric lifetime. For the gas phase species (CO, benzene, and NO$_x$), we use a constant value of 140 pptv for the background values of HCN for flights before 21 March. Following a shift in weather on 21–22 March (Fast et al., 2007) (see also Figs. A5–A7, and supplementary material (http://www.atmos-chem-phys.net/9/4929/2009/acp-9-4929-2009-supplement.pdf) for higher resolution MODIS images), we find higher background concentrations for HCN (220 pptv). C$_2$H$_2$ backgrounds are quite small (0–30 pptv), so we have used a constant background value of 0 pptv for the analysis of the gas phase species. For organic aerosol and scattering, we have used a flight-by-flight analysis of the correlation of organic aerosol mass and scattering with HCN and C$_2$H$_2$ to define the background HCN and C$_2$H$_2$. For flights in early March, the implied background of HCN and C$_2$H$_2$ for organic aerosol (the abundance of these gases when the amount of organic aerosol is zero) is very close to the global backgrounds used in the analysis of the gases, but following the rainy period, 20–26 March, the apparent backgrounds for both HCN and C$_2$H$_2$ increased more drastically than for the gas phase pollutants; organic aerosol concentrations were near zero at significantly higher concentrations of HCN and C$_2$H$_2$ (300 and 150 pptv, respectively). This difference is likely due to removal of aerosol (but not insoluble gases such as HCN, CO, C$_2$H$_2$, etc.) in the region during the rain storms.

The variance in the abundance of our fire and urban tracers (HCN and C$_2$H$_2$) explains most of the variability in CO and other pollutants. For example, in panels (b), (c), and (d) of Fig. 3, we show the observations (circles) for CO, benzene, and organic aerosol mass along the C-130 flight track. The observations are averaged to the sampling time of the whole air samples used to determine C$_2$H$_2$. The bars show the contributions from fire and urban emissions, estimated from [HCN]$^*$ (orange) and acetylene, [C$_2$H$_2$]$^*$ (black) using the emission ratios described in Table 1 and the mixing model described by Eq. (1).

Figure 5a shows a scatter plot of the predictions from the two component model and all observations (from all flights)
Fig. 5. Scatter plots for observed vs. reconstructed CO (a) and benzene (d). Associated scatter plots for CO vs. each tracer, [C$_2$H$_2$]$^*$ (b) and [HCN]$^*$ (c) and for benzene vs. each tracer, [C$_2$H$_2$]$^*$ (e) and [HCN]$^*$ (f). Lines are best fit to all data using total least squares (TLS) regression. Points are colored by [toluene]/[C$_2$H$_2$]$^*$, where red=1 and blue=0. Points with black border lie within the 3×3 degree study area.

made from the C-130 during MILAGRO for CO. The observations made within the 3×3 degree box surrounding MC, during the 7 flights considered in this analysis are highlighted with a black border. CO has a relatively long lifetime in the atmosphere and there is a persistent northern hemispheric background of between 60 and 150 ppb that varies with season and latitude. To estimate the regional increase in [CO], we assume that the background [CO] is equal to the simulation of background CO taken from the Model for OZone And Related chemical Tracers (MOZART) chemical transport model (Horowitz et al., 2003), plus a constant 34 ppbv offset (see Fig. 3b, green bars). The offset was determined from the bias between MOZART simulations and the observed CO in the cleanest air encountered during MILAGRO – typically aloft and outside the Mexico City basin. Figure 5d shows the comparison for benzene. From the tracer analysis, we estimate that biomass burning accounts for (31±3)%, (36±3)%), and (34±7)% of the CO, benzene, and reactive nitrogen (NO$_x$). These estimates are the mean of the mass-weighted, daily-averaged fire/excess fractions for observations made within the 3×3 degree box centered on MC, for the seven flights considered in this analysis. The sensitivity of these ratios to the size of the box is described in Table A1. Consistent with expectation, the fraction of pollution from biomass burning is lower for a smaller box centered over the city.

Figure 6a shows a scatter plot of the predictions from the two component model for organic aerosol. The data are colored by the ratio of toluene to acetylene. High ratios (red colors) are indicative of very fresh emissions with little photochemical processing. A similar figure for submicron aerosol scattering is shown as Fig. A3. From the tracer analysis, we estimate that biomass burning accounts for (66±11)%, and (57±5)% of the organic aerosol mass and total submicron
scattering (which determines visibility). These estimates are the mean of the mass-weighted, daily-averaged fire/excess fractions for observations made within the 3×3 degree box centered on MC for the seven flights considered in this analysis.

3 Discussion

In Table 1, we compare the emission ratios estimated from the TLS analysis with observations made in the city plume encountered on 29 March (little BB influence) and in fresh fire plumes sampled within the study area by the Twin Otter (Yokelson et al., 2007b). The emission ratios derived here do not change if the data from the 29th are excluded from the least-squares analysis. Note, while Yokelson et al. (2007b) report emission factors for up to 5 fires (or groups of fires) as well as the geometric mean of the reported individual emission factors, here we report (Table 1) the median values of the individual emission ratios calculated from the emission factors given therein. Median values are reported here because of the high variability of the individual emission factors, and the non-uniform weighting across fires in the geometric mean calculation. In particular, the unusually high modified combustion efficiency (MCE) of the “17 March – Planned Fire” and the group of fires reported as “6 March – Fires 1–4” are given 25% of the weight as the remainder of the fires in calculating the geometric mean. While not as good, the comparison using the mean of the emission ratios derived from Yokelson et al. (2007b) does not alter the conclusions.

The urban ratio of CO to C2H2 measured on 29 March, 92 (mol/mol), is very close to the emission ratio estimate from the total least squares (TLS) analysis, 96 (mol/mol) (Table 1). This ratio is lower than typically observed in US (300–500 mol/mol), and Asian cities (220 mol/mol) (Xiao et al., 2007), but comparable to the value observed by Grosjean et al. (1998) in urban air in Brazil. The lower emission ratios in Mexico and Brazil may reflect differences in fuel composition or catalytic converter functionality (Sigsby et al., 1987). The relatively high concentration of C2H2 in traffic exhaust combined with the emissions of C2H2 from fires near the low end of the typical range makes C2H2 a good tracer for the Mexican urban emissions. Observations made during the same period of CO and C2H2 at a ground station (T1) in Mexico City and described by de Gouw et al. (2009) show a mean CO to C2H2 ratio of 150 mol/mol. It is not known at this time what accounts for this difference, but likely it is due to a difference in calibration factors. The analysis described here is not sensitive to an absolute error in C2H2 or CO as the emission ratios are derived internally to the data set. While a calibration error in either CO or C2H2 will affect the C2H2 emission ratios reported in Table 1, it will not change the calculated fire impact for the trace gasses reported in Table A1.

The median ratio of CO to HCN observed in biomass burning plumes within the study area, 117 mol/mol (Yokelson et al., 2007b), is close to that derived in the total least-squares analysis, 104 mol/mol. As noted by Yokelson et al. (2007b), this ratio is at the low end of the range typically observed for biomass burning. The relatively high emissions of HCN in biomass burning plumes combined with the low emission ratios from urban sources makes HCN a very good tracer for biomass burning in Mexico City.

Biomass burning is a significant global source of benzene and the impact of this source in and around the Mexico City...
basin is quite apparent in the correlations of benzene with the tracers (Fig. 5d–f). From the least squares analysis, we estimate that fires contributed 36% of this pollutant to the atmosphere above the central Mexican Plateau in March 2006. Relative to CO, the benzene emission ratio for biomass burning derived here is similar to those reported in other studies (Andreae and Merlet, 2001). As an aside, measurements of the ratio of benzene and toluene have been used in many previous studies to estimate the aging of an urban airmass as the atmospheric oxidation of toluene occurs much more rapidly than benzene (e.g., Cubison et al., 2006). Because the benzene/toluene emission ratio from fires (∼1–3) is much greater than the same emission ratio from urban emissions (0.2), this method is not appropriate for cities – such as Mexico City during the biomass burning season – that have significant contributions of benzene from fire emissions.

Organic aerosol generally accounts for more than half the mass of fine particulate matter (PM$_{2.5}$) in (Salcedo et al., 2006) and around (DeCarlo et al., 2008) Mexico City. As shown in Fig. 6b–c, the amount of organic aerosol is highly correlated with [HCN]$^*$ and [C$_2$H$_2$]$^*$, suggestive of large fire and urban emission influences. Using the emission ratios described in Table 1, we estimate that biomass burning contributes 66% of the organic aerosol to the study area in March 2006. These estimates are quite uncertain, however, due to complex aerosol chemistry.

Organic aerosol is both formed in and lost from the atmosphere on relatively fast timescales. Although direct (or primary) emissions of organic aerosol from automobiles are quite small (∼5–10 µg per standard ($T=273$ K, $P=1$ atm) cubic meter of air (sm$^3$)) per ppmv CO (de Gouw et al., 2005; Robinson et al., 2007)), subsequent atmospheric oxidation of co-emitted hydrocarbons such as toluene and other aromatics, as well as biogenic and biomass burning hydrocarbon emissions, can yield low vapor pressure compounds that condense on the existing particulate forming secondary organic aerosol (SOA) (Kroll and Seinfeld, 2008). The amount of SOA produced from these gas-phase sources over the period of a day substantially exceeds the primary emissions from urban sources (de Gouw et al., 2005; Kleinman et al., 2008; Robinson et al., 2007; Volkamer et al., 2006). The influence of this process is apparent in the aircraft data. In Fig. 6a, the organic aerosol data is colored by the ratio of toluene to acetylene. Toluene is co-emitted with acetylene in urban emissions, but is oxidized in the atmosphere with a lifetime of approximately 12 daylight hours. In samples containing very high toluene (less oxidized), the total least-squares analysis tends to over-predict the amount of organic aerosol. For example, the urban emission ratio derived from the least-squares analysis (∼39 µg per sm$^3$ per ppmv CO or ∼3.9 µg per sm$^3$ per ppbv of C$_2$H$_2$) is greater than the factor derived from the slope of the correlations in the fresh plumes encountered on 29 March.

In addition to aerosol growth, aerosol mass can be lost through several mechanisms. As aerosol is transported away from its source and diluted with clean air, semi-volatile compounds evaporate to the gas phase (Robinson et al., 2007). Dry and wet deposition also remove aerosol from the atmosphere. As mentioned earlier, the changing backgrounds for aerosol relative to insoluble gases observed during late March are likely explained by aerosol loss via wet deposition.

Despite the complexity of the aerosol chemistry, the simple two end-member mixing model does describe much of the variability of organic aerosol mass observed from the C-130. This result may be related to our sampling – most of the observations were made in the afternoon when the city and fire emissions had experienced some aging. The apparent organic aerosol emission ratio for urban emissions derived from the 29 March plume, 32 µg OA per sm$^3$ per ppmv CO, is not inconsistent with other estimates for organic aerosol from urban emissions aged 3–6 h (de Gouw et al., 2005; Volkamer et al., 2006). The aerosol burden continues to increase as the air masses are further oxidized in the MC outflow (Kleinman et al., 2008); similar to what has been observed in the New York City (de Gouw et al., 2005) and Atlanta plumes (Weber et al., 2007).

The complexity of the mixing of the urban-core aerosol emissions with the regional aerosol emissions (often dominated by fire emissions) as well as mixing with the clean free troposphere precludes strong statements on the aerosol dynamics of these different sources.

The average NO$_x$/VOC emission ratio for the fires surrounding Mexico City is similar to the urban NO$_x$/VOC emission ratio for Mexico City. This is due in part to the higher than expected NO$_x$ emission from the fires, which is significantly (2–4 times) larger than typical for fires (Yokelson et al., 2007). As the fire and urban NO$_x$/VOC emissions are similar, the fire emissions will significantly impact the ozone production in the Mexico City outflow. Modeling of the transport and aging of the MC plume using a 3-D chemical transport model with full chemistry and accurate urban and fire emissions will be required to quantify the full impact of the fire emissions on ozone concentrations.

4 Implications for air quality improvement

The implications of this study for air quality engineering in Mexico City are not straightforward. Although visibility within the city and the export of aerosol, ozone, and other trace gases is significantly impacted by biomass burning during the period of our observations, it is not possible to estimate from this data set the long-term impact of such burning on the urban dwellers of Mexico City. Biomass burning in March 2006 was significantly higher than typical for March, though not unlike the amount of burning usually observed in the height of the biomass burning in April/May. During most of the year (June–February), however, these sources are negligible. Thus, annually-averaged, the impact
of biomass burning will certainly be much smaller. In addition, because our sampling was from aircraft and primarily in the afternoon, and because the biomass burning was primarily in the forest above the city (Yokelson et al., 2007b), the impact of fire on the air breathed by people within Mexico City will be smaller than the regional impact estimated here. Indeed, we observe that even in the relatively well mixed afternoon planetary boundary layer, the impact of fire increases with altitude (Fig. A4). Consistent with this finding, estimates of the impact of fire on air quality at the ground stations within the city suggest a smaller fire influence (Molina et al., 2007; Bravo et al., 2002; Salcedo et al., 2006; Yokelson et al., 2007b; Stone et al., 2008; Moffet et al., 2008; Aiken et al., 2009). A possible method for estimating the impact of fires to people on the ground in Mexico City would be through the use of a high resolution 3-D chemical transport model constrained by accurate winds and meteorological conditions, accurate fire emissions, and coupled with population maps. For example, an extension of the Fast et al. (2009) study could provide such an estimate, however additional work would be required to properly model the nighttime and early morning boundary layer, as well as an adequate parameterization of SOA growth which considers both anthropogenic and biomass burning precursors. Finally, unlike reducing emissions from the urban sources, reducing fire emissions through fire suppression efforts may have environmental costs as well as benefits; although forest fire suppression in and around the basin would yield improvement in visibility, such fire suppression actions may be inconsistent with proper forest management practices.

### Table A1. Fire impact for pollutants considering several box sizes.

<table>
<thead>
<tr>
<th>Species</th>
<th>Box 1 (%)</th>
<th>Box 2 (%)</th>
<th>Box 3 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>21</td>
<td>28</td>
<td>31</td>
</tr>
<tr>
<td>C6H6</td>
<td>25</td>
<td>33</td>
<td>36</td>
</tr>
<tr>
<td>NO2</td>
<td>24</td>
<td>31</td>
<td>34</td>
</tr>
<tr>
<td>OA</td>
<td>52</td>
<td>61</td>
<td>66</td>
</tr>
<tr>
<td>scattering</td>
<td>43</td>
<td>52</td>
<td>57</td>
</tr>
</tbody>
</table>

*a* Box defined by latitude: [19.269 to 19.867] and longitude: [-99.300 to -98.867].

*b* Box defined by latitude: [19.067 to 20.033] and longitude: [-99.433 to -98.667].

*c* Box defined by latitude: [17.900 to 20.900] and longitude: [-100.400 to -97.400].

### Fig. A1. Scatter plot of the HCN and the CH3CN observations made aboard the C-130 during the MILAGRO experiment ($r^2=0.78$). The data (black x symbols) were fit using least squares (red: slope=0.45, intercept=0.039 ppbv), and robust total least squares (blue: slope=0.39, intercept=0.054) regression techniques. Robust least squares is an iterative algorithm using reweighted least squares and a bisquare weighting function, where data which lie far from the best fit are given less weight in successive iterations. The $r^2$ value is calculated via the standard method.

### Appendix A

#### Additional information

**A1 Data sources**

HCN was measured on the NCAR C-130 as discrete 0.5 s samples obtained every 5 s. The analysis was performed by chemical ionization mass spectrometry (CIMS). While the Caltech CIMS instrument has been described previously in detail (Crounse et al., 2006), the particular HCN method has not. In brief, HCN reacts rapidly with the CF3O− (and CF3O−·H2O) anion to form the cluster product ion, CF3O−·HCN, which is monitored at $m/z=112$. Instrumental backgrounds were measured once every 15 min by passing ambient air through a filter containing nylon wool coated with NaHCO3. In-flight calibrations were performed once per hour using HNO3 and H2O2 calibration standards and proxied to laboratory calibrations of HCN. Similar to H2O2, the sensitivity of the CIMS instrument toward HCN is a function of water vapor. This is corrected for using the aircraft water vapor measurement and a water vapor sensitivity curve for HCN determined in the laboratory. Absolute laboratory calibrations were conducted using HCN permeation.
Fig. A2. Scatter plot for observed vs. reconstructed NOy (a). Associated scatter plots for NOy vs. each tracer, [C2H2]* (b) and [HCN]* (c). Lines are best fit to all data using total least squares (TLS) regression. Points are colored by [toluene]/[C2H2]*, where red=1 and blue=0. Points with black border lie within the 3×3 degree study area.

Fig. A3. Scatter plot for observed vs. reconstructed submicron scattering (a). Associated scatter plots for submicron scattering vs. each tracer, [C2H2]* (b) and [HCN]* (c). Lines are best fit to all data using total least squares (TLS) regression. Points are colored by [toluene]/[C2H2]*, where red=1 and blue=0. Points with black border lie within the 3×3 degree study area.

CH3CN was measured by a cryotrap concentrator coupled to a gas chromatograph mass spectrometer (cryo-GCMS), an instrument similar to the one described by Apel et al. (2003). The cryo-GCMS instrument concentrated ambient air in the...
cryotrap for 45 s prior to a 125 s analysis, yielding one data point every 170 s. The accuracy for the cryo-GCMS CH$_3$CN determination is estimated to be ±20% with a precision of ±3%.

C$_2$H$_2$, benzene, and toluene were recovered from 2 L canister samples that were periodically filled (approximately 12 samples per hour for the flights into the city). Each canister is filled over a period of 30–120 s. These samples were analyzed by gas chromatography at UC-Irvine (Blake et al., 1997). The detection limit for each of these compounds is 3 pptv, and the accuracy is estimated to be better than ±5% for C$_2$H$_2$ and ±10% for benzene and toluene.

NO$_y$ was measured via catalytic reduction of reactive nitrogen species to NO on a gold surface in the presence of a small flow of CO, used as a reducing agent. The resulting NO was measured by the standard chemiluminescence technique. In order to keep the conversion efficiency constant the converter was maintained at a constant pressure using a heated teflon valve just upstream of the converter. The converter was housed in the inlet pylon, extending about 30 cm into the airstream, in order to minimize the length of inlet tubing upstream of the converter. The upstream plumbing (tubing and valve) were limited to a heated length of about 15 cm. The inlet tubing and converter were oriented perpendicular to the airstream with an aft-facing 45° cut on the end of the tubing. This configuration minimized particle amplification, and tended to exclude larger particles (1 micron or so). At 1 ppbv, the estimated NO$_y$ precision and accuracy are ±3% and ±15%, respectively.

Carbon monoxide was measured continuously with an ultraviolet (VUV) fluorescence instrument similar to the one developed by Gerbig et al. (1999) with a precision of 3 ppbv and a typical accuracy of ±10% at 100 ppbv CO. On several flights, the VUV fluorescence CO measurements were not available and CO concentrations were determined from the canister samples.

Aerosol composition (organic, sulfate, nitrate, ammonium, and chloride) and mass were determined with a high resolution aerosol mass spectrometer (HR-AMS) (DeCarlo et al., 2006, 2008). The HR-AMS detection limit for the organic aerosol is 0.35 µg sm$^{-3}$ for a 12 s integration period, and its accuracy is estimated to be better than ±25%. Aerosol scattering coefficients were measured at 450, 550, and 700 nm wavelengths using two TSI-3563 nephelometers. One nephelometer measured submicron scattering employing a 1 µm aerodynamic impactor, and the other measured total scattering (Anderson et al., 2003).

**A2 Alternative HCN sources**

In addition to biomass burning and gasoline/diesel combustion, other sources of HCN may contribute to the enhanced HCN in the Mexico City basin. For example, HCN has also been shown to be produced in the pyrolysis of coal from the breakdown of pyrrolic and pyridinic nitrogen (Leppalahti and Koljonen, 1995). Coal burning, however, is minimal in the basin. According to the 1999 Mexico National Emissions Inventory (NEI), 88% of the CO produced in Mexico City, and surrounding states (summing over Distrito-Federal, Mexico, and Morales) comes from mobile sources (NEI, 2008).

We did observe elevated HCN in the plumes from the power plants (fuel-oil fired) and petrochemical complex in Tula, north of Mexico City. The ratio of HCN to CO in the Tula plume is similar to that from fire. The Tula CO emissions are, however, significantly smaller than the CO emissions from fire in the MC basin suggesting these emissions have minimal influence on the regional HCN budget.

**A3 HCN and acetonitrile correlation**

A reasonable correlation ($r^2$=0.78, $n=835$) exists between HCN and CH$_3$CN observations, suggesting similar sources (Fig. A1). However, on multiple occasions, directly over Mexico City, enhanced CH$_3$CN was observed without accompanying enhancements in HCN (see Fig. A1, data excursions well above the best fit line). Explanations for these CH$_3$CN plumes could include industrial, non-combustion, sources, or possibly a different compound interfering with the GC-MS CH$_3$CN measurement. Overall, $\Delta$CH$_3$CN is 39% of $\Delta$HCN (Fig. A1). On 29 March, a day with little fire influence according to HCN levels relative to CO, the slope of CH$_3$CN to HCN is similar to the overall relationship, indicating that these compounds are emitted from urban sources.
in about the same ratio as from fire, or that fire is still the dominating source of these compounds even for days without large fires.

A4 NO$_y$ and submicron scattering

Analogous to Figs. 5 and 6, NO$_y$ (Fig. A2), and submicron scattering at 550 nm (Fig. A3) reconstructions are shown. The two component fit for total scattering yielded very similar results as the one for submicron scattering (e.g. fire fraction also equal to 57%), with similar correlation coefficients. Significant amounts of aerosol nitrate were measured in and around Mexico City. For this analysis total NO$_y$ was taken as the sum of measured NO$_y$ and aerosol nitrate. While the NO$_y$ instrument likely measures some of the aerosol nitrate as NO$_y$, at this time it is not known what fraction of aerosol nitrate was sampled by the NO$_y$ instrument. To the extent aerosol nitrate is sampled by this NO$_y$ instrument, we are double counting the aerosol nitrate. The correlation with the 2-component model is significantly better if the aerosol nitrate is added to the NO$_y$ measurement to give total NO$_y$.

A5 Altitude dependence of fire impact

As observed from the C-130, and inferred by comparing the C-130 data with observations on the ground in Mexico City, the impact of the fires surrounding Mexico City is not as severe on the ground, as it is above the City. This is due in part to the location of the fires, elevated above the city on the mountainsides surrounding the city. The impact of fire...
Fig. A6. MODIS-Aqua images around Mexico City for 14 March through 22 March 2006. Red boxes represent detected fires. Black lines are state boundaries. Blank areas represent missing data. Images courtesy of MODIS Rapid Response Project at NASA/GSFC.

increases with altitude above Mexico City (Fig. A4), suggesting that the smoke from the fires does not fully impact the ground.

A6 Box size for study area

The fire impact for the pollutants reported in the main text, considered a $3 \times 3$ degree rectangular study area, centered on Mexico City. As one decreases the box size, the urban emissions become relatively more important (Table A1). This makes sense considering the fires emissions originate from many diffuse sources (individual fires) scattered across the plateau, while the urban emissions are more centrally located in and around the MC basin. Table A1 compares the results for the 2-component model using three different box sizes. Box 1 is the smallest, encompassing the populated area of Mexico City and some adjacent terrain. Box 2 is somewhat larger, including the ring of mountains around MC. Box 3 is the largest box, equating to the $3 \times 3$ degree box used for the results presented in the main body of this work.

A7 MODIS Aqua satellite image time line

A daily timeline of true color satellite images of the Mexico City area, collected from the MODIS instrument aboard the Aqua satellite, for the Month of March 2006 are shown in Figs. A5–A7. The images were taken at approximately 1:30 p.m. local time each day. The red boxes on each satellite image represent detected thermal anomalies. One can observe from inspection of the images that many fires
are not detected due to a number of reasons, including cloud cover, smoke cover, low fire temperature, or simply lack of satellite coverage. Also, the number of detected fires does not necessarily correlate with the impact of fires on the visibility as observed from the satellite pictures. Higher resolution images are available as supplementary material (http://www.atmos-chem-phys.net/9/4929/2009/acp-9-4929-2009-supplement.pdf) for both Aqua and Terra images of the Mexico City region during March 2006.

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